handbook of
Clinical Drug Data
tenth edition

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NOTICE: The editors, Philip O. Anderson, PharmD, James E. Knoben, PharmD, and William G. Troutman, PharmD, and the contributors have written this book in our private capacities. No official support or endorsement by any university, hospital, federal agency, or pharmaceutical company is intended or should be inferred.
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Preface

The Tenth Edition of the *Handbook of Clinical Drug Data* continues a long tradition of providing clinically relevant, well-referenced drug information compiled by expert clinicians and presented in a compact format. The formats of all sections should be familiar to users of the ninth edition. As with recent editions, information in the *Handbook* is divided into three parts.

**Drug Monographs** in Part I have been updated to include numerous newly marketed and promising investigational drugs. Areas with extensive revisions include the Antivirals reflecting the many new agents for HIV infection, Immunosuppressants, Anticonvulsants, and the Hematologic Drugs. Three new subsections have been added to reflect the growing number of agents for rheumatoid arthritis, glaucoma and osteoporosis: Antiarthritis Drugs in the Analgesic and Anti-inflammatory Drugs section, Ophthalmic Drugs for Glaucoma in the Central Nervous System section, and Bisphosphonates in the Renal and Electrolytes section.

**Clinical Drug Information** in Part II continues to provide clinically useful information that helps the reader to decide which drug(s) are most likely to have caused adverse reactions or which are the best choices for patients in special populations. All drug-induced diseases sections have been extensively updated, as have the Cytochrome P450 Interactions, Pregnancy, Breastfeeding, Renal Disease, Immunization, and Cardiac Arrest sections. **Drug-Laboratory Test Interferences** in Part III has also been updated.

In this edition, we welcome several new authors: Dan Baker, Jess Benson, Toy Biederman, Juliana Chan, Paul Cuddy, Rob DiDomenico, Allison Einhorn, Ray Hammond, Patty Marshik, Gary Matzke (a returning author), Renée Mercier, and Anna Taddio (our first “international” author). We would also like to thank the previous authors whose work in most cases served as the basis for revisions of the chapters that appear in this edition by new authors: Andrea Anderson (Drugs and Pregnancy), Lisa Ashton (Respiratory Drugs), Arasb Ateshkadi (Renal and Electrolytes), Rosemary Berardi (Gastrointestinal Drugs), Larry Borgsdorf (anaphylaxis) Larry Davis (NSAIDs and Hematologic Drugs), John Flaherty (Aminoglycosides and β-Lactams), John Gambertoglio (Renal Disease), Millie Gottwald (Antimigraine Drugs and Neurodegenerative Diseases), Amy Guenette (Inotropic Drugs and Nitrates), Brian Kearney (Renal Disease), and Carolyn Zaleon (Gastrointestinal Drugs). We are saddened to report the deaths of Drs. Ateshkadi and Gambertoglio since our last edition. Both will be remembered for their professional dedication and the quality of their work. John Gambertoglio had long-time personal and professional ties to the editors and will be particularly missed.

This edition also marks another major change, being the first edition produced with our new publisher, McGraw-Hill and new editors, Stephen Zollo and...
Nicky Panton. We thank them for their efforts to maintain the high quality of the Handbook that we desire and our readers have come to expect.

Philip O. Anderson
James E. Knoben
William G. Troutman
August 2001
Part I of this book is organized around 10 major drug categories, which have been subdivided into common therapeutic groups. Within these therapeutic groups, drug information is alphabetically presented in three formats: Monographs, Minimonographs, and Comparison Charts. Monographs and Comparison Charts are grouped together to ensure that related drugs are easy to compare and contrast. Charts are located after the monographs to which they relate. Drug antagonists are grouped together with agonists to simplify organization and accessibility.

Monographs are used for drugs of major importance and prototype agents. Minimonographs are used for drugs similar to prototype drugs, those of lesser importance within a therapeutic class, and promising investigational agents. Minimonographs contain only selected subheadings of information rather than all subheadings contained in the full monographs.

Comparison Charts are used to present clinically useful information on members of the same pharmacologic class and different drugs with a similar therapeutic use, as well as to present clinically relevant information on certain other topics.

The preferred method to gain access to complete information on a particular brand or generic drug is to use the index at the end of the book. The index may also direct the user to other pertinent information on the drug.

MONOGRAPH FORMAT

CLASS INSTRUCTIONS
This is an optional heading at the beginning of each drug class. It consists of patient instructions that apply to more than one of the drug monographs in this subcategory. If all drugs are not identical in their instructions, only the common information is found here. The Patient Instructions section of each monograph that is affected states, “See Class Instructions” as the opening phrase.

GENERIC DRUG NAME
The nonproprietary (generic) name is listed on the left, followed by common brand names listed on the right. Brand-name products listed are not necessarily superior or preferable to other brand-name or generic products; “Various” indicates the availability of additional brand and/or generic products.

Pharmacology. A description of the chemistry, major mechanisms of action, and human pharmacology of the drug in clinical application.

Administration and Adult Dosage. Route of administration, indications, and usual adult dosage range are given for the most common labeled uses. Dosages correspond
HOW TO USE THIS BOOK

to those in the product labeling or in standard reference sources. “Dose” refers to a single administration and “dosage” to a cumulative amount (eg, daily dosage).

**Special Populations.** Dosages in patient populations other than the typical adult are listed:

*Pediatric Dosage* (given by age or weight range)

*Geriatric Dosage* (given by age range)

*Other Conditions* (renal failure, hepatic disease, obesity, etc.)

**Dosage Forms.** The most commonly used dosage forms and available strengths are listed, as well as popular combination product dosage forms. Prediluted IV piggyback or large-volume parenteral containers are not listed unless this is the only commercially available product.

**Patient Instructions.** Key information that should be provided to the patient when prescribing or dispensing medication is presented. When introductions apply to an entire drug category, see “Class Instructions” at the beginning of that subcategory.

**Missed Doses.** What the patient should do if one or more doses are missed.

**Pharmacokinetics.** Data are presented as the mean ± the standard deviation. Occasionally the standard error of the mean (SE) is the only information available on variability, and it is identified as such.

**Onset and Duration** (time course of the pharmacologic or therapeutic effect)

**Serum Levels** (therapeutic and toxic plasma concentrations are given)

**Fate** (The course of the drug in the body is traced. Pharmacokinetic parameters are generally provided as total body weight normalized values. The volume of distribution is either a $V_d$ in a one-compartment system or $V_c$ and $V_{d\beta}$ or $V_{dss}$ in a two-compartment system.)

$t_{1/2}$ (terminal half-life is presented)

**Adverse Reactions.** Reactions known to be dose related are usually given first, then other reactions in decreasing order of frequency. Reaction frequency is classified into three ranges. However, percentages of reactions may be provided for reactions that occur more frequently than 1%.

- frequent (>1/100 patients)
- occasional (1/100 to 1/10,000 patients)
- rare (<1/10,000 patients)

**Contraindications.** Those listed in product labeling are given. “Hypersensitivity” is not listed as a contraindication because it is understood that patients should usually not be given a drug to which they are allergic or hypersensitive—exceptions are noted.

**Precautions.** Warnings for use of the drug in certain disease states and/or patient populations, together with any cross-sensitivity with other drugs. Part II, Chapter 3, “Drug Use in Special Populations,” should be consulted for more information, particularly regarding pregnancy and breastfeeding.

**Drug Interactions.** The most important drug interactions are listed.
Parameters to Monitor. Important clinical signs and/or laboratory tests to monitor to ensure safe and effective use are presented. The frequency of monitoring may also be given; however, for many drugs the optimal frequency has not been determined.

Notes. Distinguishing characteristics, therapeutic usefulness, or relative efficacy of the drug are presented, as well as unique or noteworthy physicochemical properties, handling, storage, or relative cost.
PART I

Drug Monographs

Principal Editor: Philip O. Anderson, PharmD

- Analgesic and Anti-inflammatory Drugs
- Antimicrobial Drugs
- Antineoplastics, Chemoprotectants, and Immunosuppressants
- Cardiovascular Drugs
- Central Nervous System Drugs
- Gastrointestinal Drugs
- Hematologic Drugs
- Hormonal Drugs
- Renal and Electrolytes
- Respiratory Drugs
Antimigraine Drugs

PHARMACOLOGY. Dihydroergotamine (DHE) is a semisynthetic ergot alkaloid that is hypothesized to exert its antimigraine effect via its agonist activity at the serotonin 5-HT\textsubscript{1D} receptor, resulting in vasoconstriction of intracranial blood vessels and inhibition of inflammatory neuropeptide release.\textsuperscript{1} The drug also binds with high affinity to adrenergic and dopamine receptors; however, the antimigraine effect of these events is unknown. Compared with ergotamine, DHE is a weaker vasoconstrictor, is less active as an emetic, and is less oxytocic.

ADMINISTRATION AND ADULT DOSAGE. \textbf{IM} 1 mg initially, then 1 mg q 1 hr prn, to a maximum of 3 mg/day or 6 mg/week. \textbf{IV} (for rapid effect) 0.5–1 mg, may repeat in 1 hr to a maximum of 2 mg/day or 6 mg/week. Consider administering metoclopramide 10 mg IV before DHE to treat nausea due to migraine and prevent nausea due to the drug.\textsuperscript{2} \textbf{Intranasal} one spray (0.5 mg) into each nostril; may repeat in 15 min to a maximum of 2 mg over 24 hr.

SPECIAL POPULATIONS. \textbf{Pediatric Dosage}. Safety and efficacy not established.

\textbf{Geriatric Dosage}. Same as adult dosage.

\textbf{Dosage Forms}. \textbf{Inj} 1 mg/mL; \textbf{Nasal Spray} 4 mg/mL.

\textbf{Patient Instructions}. This drug can cause numbness and tingling in fingers, toes, or face. Notify your physician if you are pregnant or have heart disease or high blood pressure. Do not exceed the maximum dosage. The nasal spray can cause local irritation. Do not reuse the applicator; use the solution right after opening. Review training materials with your health care provider and report the use of all cold or allergy medications and all over-the-counter medications.

\textbf{Pharmacokinetics}. Onset and Duration. Onset under 5 min IV, within 15–30 min after IM or intranasal spray; duration 3–4 hr. Intranasal 50–70\% of patients respond in 4 hr.

\textbf{Fate}. The drug is absorbed directly into the systemic circulation when administered intranasally, but it undergoes extensive first-pass metabolism if given orally. Bioavailability of the nasal spray is 38 ± 16\%, variable depending on self-administration technique.\textsuperscript{3} Protein binding is 93\%. After administration of 1 mg, peak levels are 1 ± 0.4 µg/L (intranasal) and 4.4 µg/L (IM), occurring at 0.9 ± 0.6 hr (intranasal) and 0.4 ± 0.3 hr (IM).\textsuperscript{3} After IM administration, V\textsubscript{c} is 12 ± 4 L/kg, and V\textsubscript{d} is 33 ± 0.2 L/kg.
suggesting distribution into deep tissue compartments. Cl is $1.6 \pm 0.17 \text{ L/hr/kg}$. The drug is metabolized to at least 5 metabolites, 3 of which are active. The major route of excretion for DHE and its metabolites is in the feces via the bile.\textsuperscript{1,3}

$t_{1/2}$: α phase (intranasal) $1 \pm 0.5 \text{ hr}$, (IM) $0.9 \pm 0.3 \text{ hr}$; β phase (intranasal) $7.9 \pm 4 \text{ hr}$, (IM) $7.2 \pm 2.2 \text{ hr}$.\textsuperscript{3}

**Adverse Reactions.** The most frequently reported adverse events with intranasal administration are rhinitis, pharyngitis, altered sense of taste, application site reactions, nausea, vomiting, and dizziness. With all routes of administration, nausea, vomiting, diarrhea, and localized edema occur frequently.\textsuperscript{4,5} Numbness and tingling of fingers and toes, muscle pain in extremities, weakness in legs, pruritus, rash, and infection occur occasionally. Pleural and retroperitoneal fibrosis occur rarely with prolonged use.

**Contraindications.** Pregnancy and lactation; peripheral vascular disease; coronary artery disease; ischemic heart disease; hemiplegic or basilar migraine; sepsis; recent history of vascular surgery; severely impaired hepatic or renal function; hypersensitivity to ergot alkaloids.

**Precautions.** Use caution to avoid overuse by patients with chronic vascular headaches. Patients with risk factors for coronary artery disease should undergo periodic cardiovascular evaluation.

**Drug Interactions.** (See Ergotamine Tartrate.) DHE can antagonize the antianginal effects of nitrates. The risk of bleeding with warfarin (eg, wound hematoma, anemia, hematuria) is worsened with co-administration of DHE. Macrolides including erythromycin can increase the risk of ergot toxicity. Sumatriptan can exacerbate coronary artery vasospasm and should not be taken within 24 hr of DHE. SSRIs can cause weakness, hyperreflexia, or incoordination.

**Notes.** IV DHE is used when oral agents have failed to abort migraine and for terminating cluster or migraine headache in an emergency setting. It is not intended for prophylaxis or the management of hemiplegic or basilar migraine. The intranasal preparation is a noninvasive option for outpatients. Intranasal administration also results in improved bioavailability over the oral form because it does not undergo a first-pass effect in the liver. DHE does not cause physical dependence and is associated with a more favorable side effect profile than ergotamine, especially with regard to GI and peripheral vascular effects. In one study, subcutaneously administered sumatriptan appeared to be more effective than DHE nasal spray; however, DHE was better tolerated.\textsuperscript{6}

**ERGOTAMINE TARTRATE**

**Pharmacology.** Ergotamine is an ergot alkaloid that is hypothesized to exert its antimigraine effects via its agonist activity at the serotonin 5-HT\textsubscript{1D} receptor, resulting in vasoconstriction of intracranial blood vessels and inhibition of inflammatory neuropeptide release. The drug also binds with high affinity to adrenergic receptors; however, the antimigraine effect of this binding is unknown. The mechanism in migraine is thought to be vasoconstriction of cranial blood vessels, with a concomitant decrease in the amplitude of pulsations as well as depression of serotonergic neurons that mediate pain.
Administration and Adult Dosage. **PO for migraine** 2 mg initially, then 1 mg each 1/2 hr prn, to a maximum of 6 mg/day or 10 mg/week; **PR** 2 mg initially, may repeat in 1 hr prn, to a maximum of 4 mg/attack or 10 mg/week; **SL** 2 mg initially, then 2 mg q 30 min as needed, to a maximum of 6 mg/day or 10 mg/week. Titrate the dosage during several attacks gradually, then administer the minimum effective dosage with subsequent attacks. Patients who routinely require over 2 mg/attack can be given the total effective dosage at the onset of the headache.

**Special Populations. Pediatric Dosage.** Safety and efficacy not established. (>12 yr) 1 mg initially, then 1 mg q 30 min prn, to a maximum of 3 mg/attack.

**Geriatric Dosage.** No specific data are available.

**Other Conditions.** Decrease dosage by 50% in patients receiving methysergide as prophylaxis.

**Dosage Forms.** **SL Tab** 2 mg; **Tab** 1 mg with caffeine 100 mg (Cafergot, Ercaf, various); **Supp** 2 mg with caffeine 100 mg (Cafergot, Wigraine).

**Patient Instructions.** Initiate therapy at the first signs of an attack. Take only as directed and do not exceed recommended dosages. Report tingling or pain in extremities immediately.

**Pharmacokinetics. Onset and Duration.** Onset (oral) 5 hr; (rectal) 1–3 hr.

**Serum Levels.** 200 ng/L (176 pmol/L) or greater appears to be therapeutic; a high frequency of adverse reactions has been associated with levels >1.8 µg/L (1.5 nmol/L).

**Fate.** Bioavailability 1–2% orally, 5% rectally; relative bioavailability decreases in the following order: PR > PO > SL. Peak serum level after 2 mg rectally is 454 ± 407 ng/L (390 ± 350 pmol/L), 50 ± 43 min after the dose. Peak serum level after 2 mg with caffeine 100 mg orally is 21 ± 12 ng/L (18 ± 11 pmol/L), 69 ± 191 min after the dose. Vd is 1.9 ± 0.8 L/kg; Cl is 0.68 ± 0.24 L/hr/kg. The drug is extensively metabolized in the liver, with 90% of metabolites excreted in the bile. t½ 1.9 ± 0.3 hr; apparent half-life is 3.4 ± 1.9 hr after rectal administration because of slow absorption.

**Adverse Reactions.** Nausea and vomiting occur frequently. Signs and symptoms of ergotamine intoxication include weakness in legs, coldness and muscle pain in extremities, numbness or tingling of fingers and toes, precordial pain, transient tachycardia or bradycardia, and localized edema; these rarely develop with recommended dosages. Frequent or worsening headaches can occur with frequent, long-term, or excessive dosages. Ergotamine dependence can result in withdrawal symptoms occurring within 24–48 hr after drug discontinuation. Rectal or anal ulceration can occur with suppository use.

**Contraindications.** Pregnancy; peripheral vascular disease; coronary artery disease; hypertension; hepatic or renal impairment; sepsis; severe pruritus.

**Precautions.** Lactation; avoid excessive dosage or prolonged administration because of the potential for ergotism and gangrene.

**Drug Interactions.** β-Blockers, dopamine, and epinephrine can cause increased vasoconstriction and increased risk of peripheral ischemia or hypertension. The
macrolides (especially erythromycin and troleandomycin) can inhibit the metabolism of ergot alkaloids.

**Notes.** The stimulant action of preparations containing caffeine can keep patients from the beneficial effects of sleep. Caffeine, however, can improve dissolution of the oral formulation. Ergotamine is commonly used for abortive therapy of migraine and provides relief in 50–90% of patients.\textsuperscript{7} **Aspirin** (650 mg) or **naproxen** (750–1250 mg/day) might be effective in aborting migraine headache in mild cases or in patients who cannot take ergotamine. OTC products containing aspirin, **acetaminophen**, and caffeine (Excedrin Migraine) or **ibuprofen** (Advil Migraine, Motrin Migraine) have FDA approval for mild to moderate migraine. Prescription combination products such as **Midrin** and **Fiorinal** might be useful, but overuse of any antimigraine combination product can lead to rebound headache. **NSAIDs** are useful for prophylaxis against menstrual-related migraines when taken during the perimenstrual period. **Butorphanol** spray might be beneficial for patients with infrequent, severe headaches who cannot tolerate ergot products or triptans, but frequent use can cause dependency. The \( \beta \)-blockers **propranolol** and **timolol** are approved by the FDA for migraine prophylaxis, but other \( \beta \)-blockers without intrinsic sympathomimetic activity (eg, **atenolol**, **nadolol**) are also useful. **Verapamil** can prevent migraines in some patients but can take several months to reach maximum effectiveness. **Tricyclic antidepressants** (eg, **amitriptyline**, **nortripyline**) have been more successful in migraine prophylaxis than SSRIs. **Divalproex** has been used successfully for prophylaxis. Consider frequency of attacks (more than 2/month), co-morbid conditions, and side effects when choosing prophylactic therapy. Effective doses for migraine prophylaxis drugs are usually lower than those used for other indications.\textsuperscript{11}

**Methysergide Maleate**

**Pharmacology.** Methysergide is a semisynthetic ergot alkaloid, thought to act centrally as a serotonin agonist and to inhibit blood vessel permeability to humoral factors that affect pain threshold. Unlike other ergots, methysergide does not inhibit reuptake of norepinephrine and has minimal oxytocic, vasoconstrictor, and \( \alpha \)-adrenergic blocking effects. Because of its toxicity, methysergide is usually used only after other prophylactic measures have failed.

**Adult Dosage.** PO for migraine or cluster headache prophylaxis 4–8 mg/day with food. A drug-free interval of 3–4 weeks must follow each 6-month course; however, reduce the dosage gradually to avoid rebound headache.

**Dosage Forms.** Tab 2 mg.

**Pharmacokinetics.** Methysergide undergoes extensive liver metabolism to methylergonovine, a compound with greater activity and a longer elimination half-life than the parent drug (3.5 hr vs 1 hr). About 56% of an oral dose is eliminated in the urine as unchanged drug and metabolites.

**Adverse Reactions.** Insomnia, postural hypotension, nausea, vomiting, diarrhea, and peripheral ischemia occur frequently. Occasionally, heartburn, peripheral edema, rash, or arrhythmias occur. Rarely, mental depression occurs. Long-term (>6 months) therapy can cause retroperitoneal and pleuropulmonary fibrosis and
thickening of cardiac valves. The drug is contraindicated in peripheral vascular, cardiovascular, or pulmonary disease; phlebitis; pregnancy; and impaired liver or kidney function. Precautions and drug interactions are similar to those of ergotamine.12

**SUMATRIPTAN SUCCINATE**

**Pharmacology.** Sumatriptan is a serotonin (5-HT) analogue and a selective agonist at 5-HT1D receptors in cerebral vascular smooth muscle. Receptor activation results in migraine relief by vasoconstriction of intracranial blood vessels and attenuation of the release of vasoactive peptides responsible for inflammation of sensory nerves.13,14 (See also Selective Serotonin Agonists Comparison Chart.)

**Administration and Adult Dosage.** PO for migraine 25–100 mg; a second dose of up to 100 mg may be administered in 2 hr if response is unsatisfactory. A 100 mg dose might not provide any greater effect than a 50 mg dose. If headache returns, additional doses may be given q 2 hr, up to 200 mg in a 24-hr period. SC for migraine 6 mg; a second 6 mg injection may be administered 1 hr after the initial dose, but limited to no more than 2 injections within a 24-hr period. Controlled studies have not verified a beneficial effect of a second dose. Intranasal 5–20 mg in one nostril or 5 mg in each nostril; may repeat in 2 hr to a maximum of 40 mg/day.

**Special Populations.** Pediatric Dosage. (<18 yr) safety and efficacy not established.

**Geriatric Dosage.** Same as adult dosage. Consider the possibility of undiagnosed heart disease in the elderly.

**Dosage Forms.** Tab 25, 50 mg; Inj 6 mg/0.5 mL; Nasal spray 5, 20 mg.

**Patient Instructions.** Sumatriptan is used for relief of migraine and not for the prevention of a migraine attack. Do not take this drug if you are pregnant without consulting with your physician. Inform your physician if you have high blood pressure, diabetes, seizures, or heart, liver, or kidney disease. Report pain or tightness in chest, shortness of breath, wheezing, or rash immediately.

**Oral.** Do not take more than 200 mg within 24 hours and allow at least 2 hours after the first tablet. SC injection. Do not take more than 2 injections within 24 hours and allow at least 1 hour between injections. Pain or redness at injection site lasts less than 1 hour. Nasal spray. If 1 dose does not provide adequate relief, you may take another dose after 2 hours. Do not take more than 40 mg in 1 day.

**Pharmacokinetics.** Onset and Duration. PO 50% of patients respond in 2 hr; peak 1.5 hr. SC 70% of patients respond within 1 hr and 90% within 2 hr;15 peak 10–15 min.16 Intranasal 50–60% of patients respond in 2 hr.

**Fate.** Oral bioavailability is 14 ± 3% owing to presystemic metabolism and erratic absorption. Absorption is delayed by about 0.5 hr if taken with food. After a 100 mg oral dose, a peak of 54 µg/L (180 nmol/L) occurs in about 1.5 hr. SC bioavailability is 97 ± 16%; a peak of 74 µg/L (250 nmol/L) occurs in 12 min after a 6 mg SC dose. After a 20 mg intranasal dose, the mean peak is 16 µg/L (54 nmol/L). Plasma protein binding is 14–21%. Vd is 2.7 L/kg; Cl is 0.96 ± 0.12 L/hr/kg. Hepatic metabolism is by MAO-A to an indole acetic acid, followed
by glucuronidation and renal elimination. About 40% is found in the feces and 60% is excreted renally, 22% unchanged, and 40% as the active indole acetic acid metabolite.  

$\frac{1}{2}$t $\approx 1.9 \pm 0.3$ hr.

**Adverse Reactions.** Frequent side effects are pain and redness at SC injection site, tingling, hot flushes, dizziness, and chest tightness or heaviness. With the nasal spray, throat discomfort and unusual taste occur frequently. With all routes of administration, occasional weakness, myalgia, burning sensation, tightness in chest, transient hypertension, drowsiness, headache, numbness, neck pain, abdominal discomfort, mouth/jaw discomfort, and sweating occur. Rarely, cardiac arrhythmias, myocardial ischemia, polydipsia, dehydration, dyspnea, skin rashes, dysuria, and dysmenorrhea occur. The drug can accumulate in melanin-rich tissues such as the eye with long-term use. Several cases of ischemic colitis have been reported after sumatriptan use.

**Contraindications.** IV administration; ischemic heart disease; Prinzmetal’s angina; uncontrolled hypertension; concurrent administration of MAO inhibitors or within 2 weeks of discontinuation; within 24 hr of an ergotamine-containing drug or ergot derivative such as methysergide or dihydroergotamine; hemiplegic or basilar migraine.

**Precautions.** Pregnancy. Use with caution in those with impaired hepatic function, seizure disorder, neurologic lesion, or cardiovascular disease; postmenopausal women; or men >40 yr.

**Drug Interactions.** Nonselective MAO inhibitors or MAO-A inhibitors can increase the systemic availability of sumatriptan (especially after oral administration). Theoretically, ergot alkaloids and sumatriptan can cause prolonged vasospastic reactions if used together. (See Contraindications.) SSRIs can cause weakness, hyperreflexia, and incoordination when given with sumatriptan and other triptans.

**Parameters to Monitor.** Renal, hepatic, and cardiovascular status initially and q 6 months.

**Notes.** Subcutaneous sumatriptan is much more expensive than alternatives. It is effective in the treatment of cluster headache and appears to be more effective than ergotamine/caffeine in aborting migraine. (See Selective Serotonin Agonists Comparison Chart.)
<table>
<thead>
<tr>
<th>DRUG</th>
<th>DOSAGE FORMS</th>
<th>ADULT DOSAGE</th>
<th>ONSET (HR)</th>
<th>HALF-LIFE (HR)</th>
<th>COMMENTS</th>
</tr>
</thead>
<tbody>
<tr>
<td>Almotriptan</td>
<td>Tab 6.25, 12.5 mg.</td>
<td>PO 12.5 mg.</td>
<td>1–2</td>
<td>3–3.7</td>
<td>Low headache recurrence rate. Similar efficacy to oral sumatriptan, but better tolerated. No propranolol interaction.</td>
</tr>
<tr>
<td>Axert</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>An 80 mg dose is superior to 100 mg of oral sumatriptan. Does not induce CYP3A4.</td>
</tr>
<tr>
<td>Eletriptan</td>
<td>—</td>
<td>PO 20–80 mg.</td>
<td>1–2</td>
<td>4–7</td>
<td>An 80 mg dose is superior to 100 mg of oral sumatriptan. Does not induce CYP3A4.</td>
</tr>
<tr>
<td>(Investigational—Pfizer)</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>(Investigational—Pfizer)</td>
</tr>
<tr>
<td>Frovatriptan</td>
<td>—</td>
<td>PO 2.5 mg.</td>
<td>2–4</td>
<td>25</td>
<td>Lowest recurrence rate.</td>
</tr>
<tr>
<td>(Investigational—Elan)</td>
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<tr>
<td>Naratriptan</td>
<td>Tab 1, 2.5 mg.</td>
<td>PO 1–2.5 mg, may repeat in 2 hr; may repeat sequence once in 4 hr, to a maximum of 5 mg/day.</td>
<td>1–3</td>
<td>6</td>
<td>Low headache recurrence rate. More specific than other agents for 5HT1B. Smoking increases and oral contraceptives decrease clearance.</td>
</tr>
<tr>
<td>Amerge</td>
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<tr>
<td>Rizatriptan</td>
<td>Tab 5, 10 mg (conventional and rapidly dissolving).</td>
<td>PO 5–10 mg, may repeat in 2 hr, to a maximum of 30 mg/day.</td>
<td>0.5–2</td>
<td>2–3</td>
<td>Onset of rapidly dissolving tablet is slightly slower than conventional. Reduce dose when used with propranolol.</td>
</tr>
<tr>
<td>Maxalt</td>
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<tr>
<td>Maxalt-MLT</td>
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<tr>
<td>Sumatriptan</td>
<td>Inj 6 mg, Tab 25, 50 mg, Nasal Spray 5, 20 mg/spray.</td>
<td>SC 6 mg, may repeat once in 1 hr, to a maximum of 12 mg/day. PO 25–100 mg, may repeat q 2 hr to a maximum of 200 mg/day. Nasal 5–20 mg, may repeat once in 2 hr to a maximum of 40 mg/day.</td>
<td>0.2 (SC) 0.5–1 (PO) &lt;1 (Nasal)</td>
<td>2.5</td>
<td>Headache recurrence rate of 40%; relatively high (5%) frequency of chest pain and tightness.</td>
</tr>
<tr>
<td>Imitrex</td>
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<tr>
<td>Zolmitriptan</td>
<td>Tab 2.5, 5 mg. Tab (rapidly dissolving) 2.5 mg.</td>
<td>PO 2.5–5 mg, may repeat once in 2 hr to a maximum of 10 mg/day.</td>
<td>0.5–2</td>
<td>3</td>
<td>Cimetidine or oral contraceptives increase AUC by 50%.</td>
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<tr>
<td>Zomig</td>
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<tr>
<td>Zomig-ZMT</td>
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</table>

From references 13, 14, and 21–29 and product information.
Antirheumatic Drugs

ETANERCEPT

**Pharmacology.** Etanercept is a dimeric fusion protein that binds to tumor necrosis factor (TNFα and β) and blocks its interaction with TNF receptors on the cell surface. This reduces the signs and symptoms of rheumatoid arthritis and delays joint damage in adults with moderate to severe rheumatoid arthritis. It is indicated for patients with inadequate response to one or more disease-modifying drugs.

**Administration and Adult Dosage.** SC for **rheumatoid arthritis** 25 mg twice weekly (72–96 hr apart).

**Special Populations.** **Pediatric Dosage.** (<4 yr) safety and efficacy not established; (4–17 yr) SC for **juvenile rheumatoid arthritis** 0.4 mg/kg twice weekly, not to exceed 25 mg per dose.

**Geriatric Dosage.** Same as adult dosage.

**Dosage Forms.** Inj 25 mg.

**Patient Instructions.** This drug may be self-administered. Instruct patient on proper injection preparation and subcutaneous injection technique along with appropriate syringe and needle disposal methods. Rotate the injection sites and give injections at least 1 inch from an old site; avoid tender, bruised, red, or hard areas. Inform your physician immediately of any persistent fever, bruising, bleeding, or pallor. (See also Notes.)

**Missed Doses.** Injections should be given 72–96 hours apart. Give a missed dose as soon as possible and resume usual schedule.

**Pharmacokinetics.** Fate. Bioavailability with SC injection is 58%, with peak plasma concentrations achieved within 48–96 hr. Median clearance is 52 mL/hr/m². $t_{1/2}$ 115 hr.

**Adverse Reactions.** Injection site reactions that involve mild to moderate erythema, itching, pain, or swelling occur in about 37% of patients. Upper respiratory infections, headache, rhinitis, dizziness, pharyngitis, and cough occur frequently. Etanercept is well tolerated in children with juvenile rheumatoid arthritis, with adverse reactions similar to those experienced by adults. Rare cases of CNS demyelinating disorders and pancytopenia have been reported.

**Contraindications.** Sepsis.

**Precautions.** Do not administer to patients with active infections or children with significant exposure to varicella virus. In patients with juvenile rheumatoid arthritis exposed to varicella zoster, temporarily discontinue etanercept and give varicella zoster immune globulin. Update vaccinations before initiating etanercept therapy. Do not give live vaccines during etanercept therapy. The needle cover provided with the diluent syringe contains latex and should not be handled by those with latex allergy. Administer with caution to patients with recent history of CNS demyelinating disorders.
Parameters to Monitor. Monitor patients closely for infection and hematologic abnormalities during therapy. Discontinue treatment if serious infection, sepsis, or hematologic abnormality develops.

Drug Interactions. None known.

Notes. Etanercept sterile powder must be refrigerated at 2–8°C (38–46°F); do not freeze. Reconstitute 25 mg vial with 1 mL of bacteriostatic sterile water (included); inject diluent slowly to avoid foaming. Administer the solution as soon as possible after reconstitution; however, the solution may be stored under refrigeration for up to 6 hr in the vial. Etanercept may be used concurrently with other rheumatoid arthritis therapies such as analgesics, corticosteroids, or methotrexate. Etanercept is also being studied for the treatment of CHF, endometriosis, organ transplantation, and cachexia.31

<table>
<thead>
<tr>
<th>INFLIXIMAB</th>
<th>Remicade</th>
</tr>
</thead>
<tbody>
<tr>
<td>Pharmacology. Infliximab is a chimeric monoclonal antibody that binds to soluble and transmembrane forms of TNFα, thereby neutralizing the activity of TNFα and inhibiting TNFα binding to its receptor sites. It has no effect on lymphotoxin (TNFβ).32,33 Infliximab induces pro-inflammatory cytokines including interleukins 1 and 6 and increases endothelial cell permeability by enhancing leukocyte migration.</td>
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</tr>
<tr>
<td>Administration and Adult Dosage. IV infusion for rheumatoid arthritis 3 mg/kg, with repeat infusions at weeks 2 and 6, then q 8 weeks thereafter. For rheumatoid arthritis, infliximab is indicated to be used with methotrexate. IV for moderately to severely active Crohn’s disease 5 mg/kg as a single IV infusion. Some patients might benefit from treatment q 8 weeks after the single infusion.34 IV for fistulizing Crohn’s disease 5 mg/kg at weeks 0, 2, and 6. (See Notes.)</td>
<td></td>
</tr>
<tr>
<td>Special Populations. Pediatric Dosage. Safety and efficacy not established.</td>
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<tr>
<td>Geriatric Dosage. Same as adult dosage.</td>
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</tr>
<tr>
<td>Dosage Forms. Inj 100 mg.</td>
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</tr>
<tr>
<td>Patient Instructions. Infliximab is administered intravenously by your health care professional. Notify your physician if chest pain, fever, chills, facial flushing, itching, hives, or difficult breathing occurs within a few hours of administration.</td>
<td></td>
</tr>
<tr>
<td>Pharmacokinetics. Fate. Infliximab is distributed primarily within the vascular compartment. Direct and linear relationship between dose, maximum serum concentration, and AUC. Age and weight do not affect Cl or Vd. No systemic accumulation of infliximab occurs. t½ 8–9.5 days.</td>
<td></td>
</tr>
</tbody>
</table>
| Adverse Reactions. Serious infections have been reported. Infusion-related reactions such as fever, chills, pruritus, urticaria, chest pain, hypotension, hypertension, and dyspnea have occurred during or within the 2-hr postinfusion period. If these reactions occur, slow the infusion rate. Reactions occurring in ≥5% of patients include headache, nausea, abdominal pain, fatigue, fever, pharyngitis, vomiting, pain, dizziness, bronchitis, rash, rhinitis, chest pain, coughing, pruritus, sinusitis, myalgia, and back pain. Hypersensitivity reactions to infliximab can
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occur and antibodies to infliximab develop in about 13% of patients. Patients most likely to experience infusion-related reactions are those who developed antibodies. Have medications (eg, acetaminophen, antihistamine, corticosteroid, and epinephrine) available for immediate use in the event of a hypersensitivity reaction. Lupus-like syndrome (1 in 340 patients) and lymphoproliferative disorders occur rarely.

Contraindications. Hypersensitivity to murine proteins; presence of serious infection.

Precautions. Women should use adequate contraception for the duration of and at least 6 months after therapy. Use caution when infliximab is administered with immunosuppressive therapy or to patients who have a history of infections. Avoid use in patients with known GI luminal strictures.

Parameters to Monitor. Monitor patients closely for adverse effects, especially for infusion-related reactions during or within the 2-hr postinfusion period and for infection during therapy. (Crohn’s disease) Observe for improvement in abdominal cramping and in bowel consistence and rectal bleeding.

Drug Interactions. None known.

Notes. Dilute the total volume of the reconstituted infliximab solution dose to 250 mL with 0.9% NaCl. Gently mix. Administer over at least 2 hr through a non-pyrogenic, low protein-binding filter with a pore size of ≤1.2 µ. Infliximab has been reported to be effective in the treatment of severe esophageal Crohn’s disease and refractory perineal cutaneous Crohn’s disease.

LEFLUNOMIDE

Pharmacology. Leflunomide’s active metabolite (M1) inhibits dihydro-oratate dehydrogenase, thereby inhibiting pyrimidine biosynthesis. M1 exhibits immunomodulating and anti-inflammatory effects.

Administration and Adult Dosage. PO for rheumatoid arthritis 100 mg/day for 3 days, then 20 mg/day. Reduce dose to 10 mg if 20 mg is not tolerated.

Special Populations. Pediatric Dosage. Safety and efficacy not established.

Geriatric Dosage. Same as adult dosage.

Dosage Forms. Tab 10, 20, 100 mg.

Patient Instructions. Do not use if you are pregnant or planning to become pregnant. Men should use condoms because leflunomide can cause birth defects. Also, men planning on fathering children should discontinue leflunomide therapy and consult with their physicians. If you experience any major medical problems while on therapy, notify your physician. Avoid alcohol because this medication with alcohol can increase the risk of liver damage. Avoid immunizations unless approved by your physician.

Missed Doses. Take a missed dose as soon as you remember; if it is near the time for next dose, skip the dose; do not take a double dose.

Pharmacokinetics. Fate. Leflunomide is 80% bioavailable, with peak plasma levels achieved in 6–12 hr. Because of its long half-life, an oral loading dosage is
given over 3 days. Leflunomide is metabolized to a primary active metabolite (M1), with the parent drug rarely detectable in plasma. The specific site of metabolism is unknown; however, hepatic cytosolic and microsomal cellular fractions have been identified. \( V_{\text{dss}} \) of M1 is 0.13 L/kg; 99.3% is bound to albumin. M1 is eliminated by renal and biliary routes. Approximately 45% is eliminated as glucuronide and oxanilic acid metabolites in the urine and 48% as M1 in the feces. \( t_{1/2} \) (M1) 18 ± 9 days.

**Adverse Reactions.** Diarrhea, dyspepsia, hypertension, headache, rash, alopecia, and elevated liver function tests occur frequently. (See Notes.)

**Contraindications.** Immunocompromised patients; those positive for hepatitis B or C; pre-existing hepatic impairment; women planning to conceive.

**Precautions.** Caution in patients with renal insufficiency. Do not give live vaccines to patients receiving leflunomide.

**Drug Interactions.** Potentially hepatotoxic medications such as methotrexate can increase risk of hepatotoxicity. Rifampin increases peak plasma levels of M1. M1 inhibits CYP2C9. Plasma-free fraction of NSAIDs and tolbutamide levels might be increased. Co-administration with cholestyramine or activated charcoal decreases M1 levels.

**Parameters to Monitor.** Monitor ALT at baseline and then monthly. If ALT levels are stable, monitor per clinical judgment.

**Notes.** If toxicity develops or if plasma levels must be decreased quickly, follow this drug elimination protocol: administer cholestyramine 8 g tid for 11 days. Verify that plasma levels are <0.02 mg/L by 2 separate tests at least 14 days apart. Without this procedure, drug elimination can take up to 2 yr. Leflunomide is equally or more effective than traditional antirheumatic agents such as methotrexate, sulfasalazine, injectable gold, and cyclosporine.\(^{39}\)
<table>
<thead>
<tr>
<th>DRUG</th>
<th>DOSAGE FORMS</th>
<th>ADULT DOSAGE</th>
<th>ADVERSE EFFECTS</th>
<th>LABORATORY MONITORING</th>
</tr>
</thead>
<tbody>
<tr>
<td>Auranofin</td>
<td>Cap 3 mg.</td>
<td>PO 3–9 mg/day (3 mg as a single dose and 6 and 9 mg/day as 2 and 3 divided doses, respectively).</td>
<td>Loose stools, diarrhea, abdominal pain or cramping, rash, pruritus, stomatitis.</td>
<td>CBC, platelets, urine dipstick for protein q 4–12 weeks.</td>
</tr>
<tr>
<td>Ridaura</td>
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<tr>
<td>Aurothioglucose</td>
<td>Inj 50 mg/mL.</td>
<td>IM 25–50 mg at 3–4 week intervals.</td>
<td>Cutaneous reactions, stomatitis, gingivitis, glossitis, hematologic toxicity, nephrotoxicity, hepatotoxicity.</td>
<td>CBC, platelets, urine dipstick q 1–2 weeks for first 20 weeks, then q 1–2 months.</td>
</tr>
<tr>
<td>Solganal</td>
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<tr>
<td>Azathioprine</td>
<td>Tab 50 mg.</td>
<td>PO 1–2.5 mg/kg/day.</td>
<td>Myelosuppression, nausea, vomiting, anorexia, diarrhea, hepatotoxicity.</td>
<td>CBC, platelets q 1–2 weeks with changes in dosage and q 1–3 months thereafter.</td>
</tr>
<tr>
<td>Imuran</td>
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<tr>
<td>Cyclosporine</td>
<td>Cap 25, 100 mg.</td>
<td>PO 1.2–7.5 mg/kg/day in divided doses.</td>
<td>Nephrotoxicity, hypertension, tremor, hirsutism, gingival hyperplasia, diarrhea, nausea, vomiting.</td>
<td>Cr, q 2 weeks until stable dosage, then monthly; periodic CBC, K⁺, and LFTs.</td>
</tr>
<tr>
<td>Neoral(^\text{a})</td>
<td>Soln 100 mg/mL.</td>
<td></td>
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</tr>
<tr>
<td>Etanercept</td>
<td>Inj 25 mg.</td>
<td>IV 25 mg twice weekly.</td>
<td>Erythema, itching, pain, swelling at inj site, headache, rhinitis, dizziness, cough.</td>
<td>None.</td>
</tr>
<tr>
<td>Enbrel</td>
<td></td>
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<tr>
<td>Gold Sodium Thiomalate</td>
<td>Inj 10, 25, 50 mg/mL.</td>
<td>IM 25–50 mg q 2 weeks for 2–20 weeks (may increase interval to 3–4 weeks if stable).</td>
<td>See aurothioglucose.</td>
<td>See aurothioglucose.</td>
</tr>
<tr>
<td>Aurolate</td>
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<tr>
<td>Hydroxychloroquine Sulfate</td>
<td>Tab 200 mg.</td>
<td>PO 200–400 mg/day.</td>
<td>Retinopathy, nausea, vomiting, diarrhea, pruritus.</td>
<td>None.</td>
</tr>
<tr>
<td>Plaquenil</td>
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<thead>
<tr>
<th>DRUG</th>
<th>DOSAGE FORMS</th>
<th>ADULT DOSAGE</th>
<th>ADVERSE EFFECTS</th>
<th>LABORATORY MONITORING</th>
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<tbody>
<tr>
<td>Infliximab(^a) Remicade</td>
<td>Inj 100 mg.</td>
<td>IV 3 mg/kg, repeat at weeks 2 and 6, then q 8 weeks. Can be given to patients on methotrexate.</td>
<td>Infusion reactions, headache, nausea, fatigue, myalgia, rhinitis, pain, pruritus, urticaria, hypo- or hypertension, chest pain, vomiting, dyspnea.</td>
<td>None.</td>
</tr>
<tr>
<td>Leflunomide Arava</td>
<td>Tab 10, 20, 100 mg.</td>
<td>20 mg once daily.</td>
<td>Diarrhea, respiratory infection, headache, nausea, rash, liver enzyme elevations, dyspepsia, alopecia, hypertension, teratogenicity.</td>
<td>ALT monthly during initial therapy, then periodically.</td>
</tr>
<tr>
<td>Methotrexate Mexate-AQ Rheumatrex Various</td>
<td>Tab 2.5 mg</td>
<td>PO 7.5–25 mg (as a single dose or 3 divided doses) once weekly; or SC or IM 7.5–25 mg once weekly.</td>
<td>Myelosuppression, stomatitis, abdominal distress, diarrhea, nausea, vomiting, hepatotoxicity, pulmonary toxicity.</td>
<td>CBC, platelets, AST, serum albumin, Cr (_o) q 4–8 weeks.</td>
</tr>
<tr>
<td>Penicillamine Cuprimine Depen</td>
<td>Cap 125, 250 mg</td>
<td>PO 500–750 mg/day as a single daily dose (up to 500 mg) or in divided doses if &gt;500 mg.</td>
<td>Sensitivity reaction with skin rash, renal and hematologic toxicity.</td>
<td>CBC, urine dipstick for protein q 2 weeks until dosage is stable, then q 1–3 months.</td>
</tr>
<tr>
<td>Sulfasalazine Azulfidine Various</td>
<td>Tab 500 mg.</td>
<td>PO 2 g/day in 2 divided doses.</td>
<td>Nausea, vomiting, heartburn, dizziness, headache, hypersensitivity, skin rash, leukopenia.</td>
<td>CBC q 2–4 weeks for first 3 months, then q 3 months.</td>
</tr>
</tbody>
</table>

\(^a\)Neoral, a nonaqueous liquid formulation forms an emulsion in aqueous fluids and has a higher oral bioavailability than conventional formulations (ie, Sandimmune, which is not indicated for rheumatoid arthritis). Do not use these products interchangeably.

\(^b\)FDA-approved for use in patients taking methotrexate.

Adapted from references 40–44.
Nonsteroidal Anti-inflammatory Drugs

ACETAMINOPHEN Various

**Pharmacology.** Acetaminophen possesses analgesic and antipyretic activities with few anti-inflammatory effects. It has the same effectiveness as aspirin in inhibiting brain prostaglandin synthetase but very little activity as a peripheral prostaglandin inhibitor. This difference from aspirin and other NSAIDs might explain its relative lack of effectiveness as an anti-inflammatory, antirheumatic agent. Acetaminophen does not inhibit normal platelet action, prothrombin activity, or adversely affect GI mucosal health.

**Administration and Adult Dosage.** PO for pain or fever (non-SR) 325–1000 mg q 4–6 hr, to a maximum of 4 g/day; (SR Tab) 1300 mg q 8 hr. PR for pain or fever 650 mg q 4–6 hr, to a maximum of 4 g/day.

**Special Populations.** Pediatric Dosage. PO for pain or fever 10–15 mg/kg q 4–8 hr, may repeat dose q 4 hr, not to exceed 5 doses per day; or (up to 3 months) 40 mg/dose, (4–11 months) 80 mg/dose, (12–23 months) 120 mg/dose, (2–3 yr) 160 mg/dose, (4–5 yr) 240 mg/dose, (6–8 yr) 320 mg/dose, (9–10 yr) 400 mg/dose, (11 yr) 480 mg/dose, (12–14 yr) 640 mg/dose, (>14 yr) 650 mg/dose. PR for pain or fever (3–11 months) 80 mg q 6 hr, (1–3 yr) 80 mg q 4 hr, (3–6 yr) 120 to 125 mg q 4–6 hr, to a maximum of 720 mg/day; (6–12 yr) 325 mg q 4–6 hr, to a maximum of 2.6 g/day; (>12 yr) same as adult dosage.

Geriatric Dosage. Same as adult dosage.

**Dosage Forms.** Cap 325, 500 mg; Gelcap 500 mg; Chew Tab 80, 160 mg; SR Tab 650 mg; Tab 160, 325, 500, 650 mg; Drp 48, 100 mg/mL; Elxr 16, 24, 26, 32, 65 mg/mL; Syrup 32 mg/mL; Supp 80, 120, 125, 300, 325, 650 mg.

**Patient Instructions.** Do not exceed the maximum recommended daily dosage of 4 g (2 g in alcoholics). Report unresponsive fever or continued pain persisting for more than 3–5 days to your physician. Do not use with other anti-inflammatory agents unless directed by your physician.

**Missed Doses.** If you take this drug on a regular schedule, take a missed dose as soon as you remember. If it is about time for the next dose, take that dose only; do not double the dose or take extra.

**Pharmacokinetics.** Serum Levels. (Analgesia, antipyresis) 10–20 mg/L (66–132 µmol/L). Serum concentrations >300 mg/L (2 mmol/L) at 4 hr or 45 mg/L (300 µmol/L) at 12 hr after acute overdosage are associated with severe hepatic damage, whereas toxicity is unlikely if levels are <20 mg/L (100 µmol/L) at 4 hr or 30 mg/L (200 µmol/L) at 12 hr.21 (See Notes.)

Fate. Rapid absorption from the GI tract, with peak plasma concentrations being achieved within 0.5–2 hr. Absorption of liquid preparations is more rapid. Unbound to plasma proteins at therapeutic doses; 20–50% bound in overdose. Extensively metabolized in the liver to inactive conjugates of glucuronic and sulfuric acids and cysteine (saturable) and to a hepatotoxic intermediate metabolite (first-order) by CYP1A2 and CYP2E1. The intermediate is detoxified by glutathione
(saturable). $V_d$ is 0.95 ± 0.12 L/kg; Cl is 0.3 ± 0.084 L/hr/kg, decreased in hepatitis and increased in hyperthyroidism, pregnancy, and obesity; 2–3% excreted unchanged in urine.\textsuperscript{21}

$t_{1/2}$. 2 ± 0.5 hr, decreased in hyperthyroidism and pregnancy, and increased in hepatitis and neonates.\textsuperscript{21}

**Adverse Reactions.** Nontoxic at therapeutic doses. In acute overdose (single dose equaling or exceeding 10 g or 7.5–10 g daily for 1–2 days), potentially fatal hepatic necrosis and possible renal tubular necrosis can occur, but clinical and laboratory evidence of hepatotoxicity might be delayed for several days. (See Serum Levels.) Toxic hepatitis also has been associated with long-term ingestion of 5–8 g/day for several weeks or 3–4 g/day for a year. Occasionally, maculopapular rash or urticarial skin reactions occur; methemoglobinemia, neutropenia, and thrombocytopenic purpura are rarely reported. Analgesic nephropathy has been associated with the consumption of 1–15.3 kg of acetaminophen over 3–23 yr.\textsuperscript{45}

**Contraindications.** G6PD deficiency.

**Precautions.** Use with caution in chronic alcoholics (not to exceed 2 g/day) and patients with phenylalanine hydroxylase deficiency (phenylketonuria) or G6PD deficiency. Some formulations contain aspartame, which is metabolized to phenylalanine; therefore do not use these products in patients with phenylketonuria. Also, some products contain sulfites.

**Drug Interactions.** Chronic alcoholics might be at increased risk for hepatic toxicity.\textsuperscript{46} The risk of hepatotoxicity also is increased by long-term use of other enzyme inducers (eg, barbiturates, carbamazepine, phenytoin, rifampin, sulfinpyrazone) and acetaminophen's efficacy also can be decreased by these agents. Co-administration with isoniazid increases the risk of hepatotoxicity; therefore, avoid acetaminophen in persons on isoniazid. Acetaminophen occasionally increases the anticoagulant effect of warfarin; therefore, monitor INR closely when adding or discontinuing long-term acetaminophen use.\textsuperscript{47}

**Notes.** Management of acute overdosage includes emesis and/or gastric lavage, if no more than a few hours have elapsed since ingestion. Supportive measures such as respiratory support and fluid and electrolyte therapy are recommended in addition. Administration of activated charcoal is not recommended because it can interfere with the absorption of acetylcysteine, which is used in the treatment of severe acute overdosage. Potentially dangerous acetaminophen levels (see Serum Levels) can be managed by the administration of 140 mg/kg acetylcysteine diluted 1:3 in a soft drink or plain water; follow with 70 mg/kg q 4 hr for 17 doses. If administered within 8–16 hr of ingestion, this therapy has been shown to minimize the expected hepatotoxicity, but treatment is still indicated as late as 24 hr after ingestion, with some data showing effectiveness up to 36 hr postingestion.\textsuperscript{48}

For the short-term treatment of osteoarthritis of the knee, acetaminophen 2.6 and 4 g/day are comparable to naproxen 750 mg/day and ibuprofen 1.2–2.4 g/day, respectively.\textsuperscript{49}
Pharmacology. Aspirin is an analgesic, antipyretic, and anti-inflammatory agent. Anti-inflammatory properties are related to the inhibition of prostaglandin biosynthesis. Aspirin nonselectively inhibits cyclo-oxygenase-1 (COX-1), which is associated with GI and renal effects and inhibition of platelet aggregation, and cyclo-oxygenase-2 (COX-2), which is associated with the inflammatory response. Unlike other NSAIDs, its antiplatelet effect is irreversible and permanent (because of transacetylation of platelet COX) for the life of the platelet (8–11 days). Salicylates without acetyl groups (e.g., sodium salicylate) have essentially no antiplatelet effect but retain analgesic, antipyretic, and anti-inflammatory activities. Low dosages (1–2 g/day) decrease urate excretion; high dosages (>5 g/day) induce uricosuria.

Administration and Adult Dosage. PO or PR for fever or minor pain 325–1000 mg q 4-6 hr, to a maximum of 4 g/day. PO for arthritis and rheumatic conditions 3.6–5.4 g/day in 3–4 divided doses. PO for acute rheumatic fever 5–8 g/day in divided doses. PO for prevention of TIAs or stroke 81–325 mg/day. PO for myocardial infarction risk reduction (primary prevention in healthy men >50 yr with at least one major cardiovascular risk factor) 81–325 mg/day; (secondary prevention) 162–325 mg/day. PO for unstable angina 162–325 mg/day. PO for prevention of coronary artery bypass graft occlusion 325 mg/day started 6 hr postoperatively and continued for 1 yr. PO for nonrheumatic atrial fibrillation (patients who are poor candidates for, or decline, oral anticoagulants) 325 mg/day; (patients <75 yr with no risk factors for stroke) 325 mg/day. The optimum dosage for platelet inhibition has not been determined; doses as low as 50 mg/day inhibit platelet aggregation and provide effective protection against thrombosis.

Special Populations. Pediatric Dosage. PO for juvenile rheumatoid arthritis 60–110 mg/kg/day in divided doses. PO for acute rheumatic fever 100 mg/kg/day in divided doses initially for 2 weeks, then 75 mg/kg/day in divided doses for 4–6 weeks. PO for Kawasaki disease 80–120 mg/kg/day; decrease to 10 mg/kg/day after fever resolves. (See Precautions.) PO as an analgesic/antipyretic 10–15 mg/kg/dose q 4 hr, to a maximum of 60–80 mg/kg/day. Alternatively, (2–3 yr) 162 mg q 4 hr; (4–5 yr) 243 mg q 4 hr; (6–8 yr) 325 mg q 4 hr; (9–10 yr) 405 mg q 4 hr; (11 yr) 486 mg q 4 hr; (≥12 yr) 650 mg q 4 hr. (See Precautions.)

Geriatric Dosage. Use minimal effective dosages; elderly are more susceptible to GI bleeding and acute renal insufficiency. PO for MI risk reduction (healthy men >50 yr for primary prevention with cardiovascular risk factors) 81–325 mg/day.

Other Conditions. Uremia or reduced albumin levels are likely to produce higher unbound drug levels that can increase pharmacologic or toxic effects. Dosage reduction might be required in these patients (e.g., kidney disease, malnutrition).

Dosage Forms. Chew Tab 81 mg; EC Tab 81, 165, 325, 500, 650, 975 mg; SR Tab 650, 800 mg; Tab 81, 325, 500 mg; Supp 120, 200, 300, 600 mg.
**Patient Instructions.** Children and teenagers (<16 yr) should not use aspirin-containing medications for chickenpox or flu symptoms because of the association with Reye’s syndrome, a rare but serious illness. Take this drug with food, milk, or a full glass of water to minimize stomach upset; report any symptoms of gastrointestinal ulceration or bleeding. Contact your physician if ringing in the ears or gastrointestinal pain occurs. Do not crush or chew enteric-coated or sustained-release preparations. Avoid other products containing aspirin or nonsteroidal anti-inflammatory drugs.

**Missed Doses.** If you take this drug on a regular schedule and you miss a dose, take it as soon as you remember. If it is about time for the next dose, take that dose only. Do not double the dose or take extra.

**Pharmacokinetics.**

**Onset and Duration.** PO onset of analgesia 30 min.²¹

**Serum Levels.** (Salicylate) 150–300 mg/L (1.1–2.2 mmol/L) for rheumatic diseases, often accompanied by mild toxic symptoms. Tinnitus occurs at 200–400 mg/L (1.5–2.9 mmol/L), hyperventilation at >350 mg/L (2.6 mmol/L), acidosis at >450 mg/L (3.3 mmol/L), and severe or fatal toxicity at >900 mg/L (6.6 mmol/L) 6 hr after acute ingestion.⁵⁵,⁵⁶

**Fate.** Rapidly absorbed from the GI tract; oral bioavailability of aspirin is 80–100%. Enteric coating does not adversely affect absorption.⁵⁷ A single analgesic/antipyretic dose produces peak salicylate levels of 30–60 mg/L (0.22–0.44 mmol/L). Aspirin is 49% plasma protein bound, decreased in uremia; \( V_d \) is 0.15 ± 0.03 L/kg; Cl is 0.56 ± 0.07 L/hr/kg. Aspirin is rapidly hydrolyzed to salicylate, which also is pharmacologically active. Salicylate is metabolized primarily in the liver to 4 metabolites (salicylic acid, phenolic and acyl glucuronides, and gentisic acid). Salicylate plasma protein binding is dose dependent, 95% at 15 mg/L and 80% at 300 mg/L, and decreased in uremia, hypoalbuminemia, neonates, and pregnancy; \( V_d \) is 0.17 ± 0.03 L/kg; Cl is dose dependent, 0.012 L/hr/kg at 134–157 mg/L, and decreased in hepatitis and neonates. Only 1% of a dose of aspirin is excreted unchanged in the urine.

\[ t_{1/2} \] (Aspirin) 0.25 ± 0.03 hr.²¹ (Salicylate) dose dependent: 2.4 hr with 0.25 g, 5 hr with 1 g, 6.1 hr with 1.3 g, 19 hr with 10–20 g.⁵⁸

**Adverse Reactions.** Hearing impairment, GI upset, and occult bleeding are frequent, with acute hemorrhage from gastric erosion also likely. As with other NSAIDs, aspirin can cause renal dysfunction, particularly in those with pre-existing renal disease or CHF. Rare hepatotoxicity occurs, primarily in children with rheumatic fever or rheumatoid arthritis and adults with SLE or pre-existing liver disease;⁵⁹ the syndrome of asthma, angioedema, and nasal polyps can be provoked in susceptible patients.⁶⁰ A single analgesic dose can suppress platelet aggregation and prolong bleeding time for up to 1 week; large dosages can prolong PT.⁶¹

**Contraindications.** Bleeding disorders; asthma; hypersensitivity to other NSAIDs or tartrazine dye.

**Precautions.** Use with caution in patients with renal disease, gastric ulcer, bleeding tendencies, hypoprothrombinemia, or history of asthma, or during anticoagulant therapy. Because of the association with Reye’s syndrome, the use of salicylates in children and teenagers with flu-like symptoms or chickenpox is not
recommended. Those developing bronchospasm with aspirin can develop similar reactions to other NSAIDs. Sodium salicylate and other nonacetylated salicylates (except diflunisal) are usually well tolerated in these patients.

**Drug Interactions.** Alkalinizing agents (e.g., acetazolamide, antacids) can reduce salicylate levels; acetazolamide also can enhance CNS penetration of salicylate. Corticosteroids can reduce serum salicylate levels. Large doses of salicylates can increase oral anticoagulant effect; even small doses can increase risk of bleeding with oral anticoagulants or heparin because of the antiplatelet effect of aspirin. Alcohol and salicylate increase the risk of GI blood loss. Salicylates can cause an increased response to sulfonylureas, especially chlorpropamide. Salicylate decreases the uricosuric effect of uricosuric agents (e.g., probenecid, sulfipyrazone). Salicylate, especially in large doses, can decrease renal elimination of methotrexate and displace it from plasma protein binding sites.

**Parameters to Monitor.** Monitor for abnormal bleeding or bruising and occult GI blood loss (periodic hematocrit) in patients who ingest salicylates regularly. Serum salicylate level determinations are recommended with higher dosage regimens because of the wide variation among patients in serum levels produced. Monitor renal function and hearing changes (tinnitus); however, using tinnitus as an index of maximum salicylate tolerance is not recommended.

**Pharmacology.** Ibuprofen is an NSAID with analgesic and antipyretic properties. It is a nonselective inhibitor of cyclo-oxygenase-1 (COX-1) and cyclo-oxygenase-2 (COX-2) and reversibly alters platelet function and prolongs bleeding time.

**Administration and Adult Dosage.** PO for mild to moderate pain 400 mg q 4–6 hr prn. PO for primary dysmenorrhea 400 mg q 4 hr prn. PO for rheumatoid arthritis and osteoarthritis 400–800 mg tid or qid, to a maximum of 3.2 g/day.

**Special Populations.** *Pediatric Dosage.* PO for fever (6 months–12 yr) 5 mg/kg for fever <102.5°F or 10 mg/kg for fever >102.5°F given q 6–8 hr, to a maximum of 40 mg/kg/day. PO for pain (6 months–12 yr) 10 mg/kg q 6–8 hr prn, to a maximum of 40 mg/kg/day. PO for juvenile arthritis 30–40 mg/kg/day in 3 or 4 divided doses; 20 mg/kg/day in milder disease.

**Geriatric Dosage.** Use minimal effective dosages because the elderly are more susceptible to GI bleeding and acute renal insufficiency.

**Dosage Forms.** Cap 200, 400 mg; Chew Tab 50, 100 mg; Tab 100, 200, 400, 600, 800 mg; Drp 40 mg/mL; Susp 20, 40 mg/mL.

**Patient Instructions.** This drug may be taken with food, milk, or antacid to minimize stomach upset. Report any symptoms of gastrointestinal ulceration or bleeding, skin rash, weight gain, or edema. Dizziness can occur; until the extent of this effect is known, use appropriate caution.

**Missed Doses.** If you take this drug on a regular schedule and you miss a dose, take it as soon as you remember. If it is about time for the next dose, take that dose only. Do not double the dose or take extra.
Pharmacokinetics. Serum Levels. 10 mg/L (48 µmol/L) for antipyretic effect.\textsuperscript{21} Serum concentrations over 200 mg/L (971 mmol/L) 1 hr after acute overdosage may be associated with severe toxicity (apnea, metabolic acidosis, and coma).\textsuperscript{65}

Fate. Rapidly absorbed from the GI tract with bioavailability over 80%.\textsuperscript{21} Peak serum levels in children of 17–42 mg/L (82–204 µmol/L) after a dose of 5 mg/kg and 25–53 mg/L (121–257 µmol/L) after a dose of 10 mg/kg are achieved in 1.1 ± 0.3 hr.\textsuperscript{66} Greater than 99% plasma protein bound; metabolized to at least 2 inactive metabolites; $V_d$ is 0.15 ± 0.02 L/kg, increased in cystic fibrosis; $Cl$ is 0.045 ± 0.012 L/hr/kg, increased in cystic fibrosis. Less than 1% is excreted unchanged in the urine.\textsuperscript{21}

$\frac{t_1}{2}$. 2 ± 0.5 hr.\textsuperscript{21}

Adverse Reactions. Gastric distress, blood loss, diarrhea, vomiting, dizziness, and skin rash occur occasionally; GI ulceration (for all NSAIDs there is a greater risk in the elderly and with higher dosages) and fluid retention have been reported.\textsuperscript{67} Ibuprofen occasionally causes renal dysfunction, particularly in those with pre-existing renal disease, CHF, or cirrhosis.\textsuperscript{68} A slight rise in the bleeding time, elevation of liver enzymes, lymphopenia, agranulocytosis, aplastic anemia, and aseptic meningitis have been reported rarely.\textsuperscript{69,70}

Contraindications. Syndrome of nasal polyps; angioedema; bronchospastic reactivity to aspirin or other NSAIDs.

Precautions. Avoid during pregnancy. Use with caution in patients with pre-existing renal disease, CHF, or cirrhosis;\textsuperscript{68} a history of ulcer disease or bleeding; or risk factors associated with peptic ulcer disease (eg, advanced age).

Drug Interactions. NSAIDs may inhibit the antihypertensive response to ACE inhibitors, β-blockers, diuretics, and hydralazine, and the natriuretic effect of diuretics. Possible GI bleeding and the antiplatelet effect of NSAIDs can increase the risk of serious bleeding during anticoagulant therapy. NSAIDs can decrease renal lithium clearance. Some NSAIDs (especially indomethacin and ketoprofen) reduce methotrexate clearance. Indomethacin (and probably other NSAIDs) can reduce renal function.

Parameters to Monitor. Monitor for blood loss, weight gain, and renal function during long-term use.

Notes. Misoprostol is effective in preventing NSAID-associated GI ulceration; H$_2$-receptor antagonists, however, prevent duodenal but not gastric ulcerations and may mask the signs and symptoms of NSAID-induced GI ulceration. Proton-pump inhibitors (eg, omeprazole) are effective in treating NSAID-related dyspepsia and preventing NSAID-induced ulcers.\textsuperscript{71}

INDOMETHACIN Indocin, Various

Pharmacology. Indomethacin is an indoleacetic acid NSAID that is one of the most potent nonselective inhibitors of cyclo-oxygenase available. In addition to its anti-inflammatory effects, indomethacin has prominent analgesic and antipyretic properties. It also has been used to suppress uterine activity and prevent premature labor.
Adult Dosage. PO for rheumatoid arthritis, rheumatoid (ankylosing) spondylitis, and osteoarthritis of the hip 25 mg bid or tid initially. Increase in 25 mg/day increments at weekly intervals until satisfactory response or to a maximum of 150–200 mg/day. Alternatively, up to 100 mg of the daily dosage may be given hs for persistent night or morning stiffness. PO for acute gouty arthritis 100 mg, followed by 50 mg tid until resolved. SR Cap 75 mg 1–2 times/day can be substituted for all uses except gouty arthritis, based on the non-SR dosage.

Pediatric Dosage. IV for pharmacologic closure of persistent patent ductus arteriosus in premature infants 0.2 mg/kg, followed by 2 additional IV doses of 0.1–0.25 mg/kg (depending on age) at 12- to 24-hr intervals. Alternatively, give 0.3 mg/kg as a single dose, or 1 or more doses of 0.1 mg/kg as a retention enema or via orogastric tube.

Dosage Forms. Cap 25, 50 mg; SR Cap 75 mg; Supp 50 mg; Susp 5 mg/mL; Inj 1 mg.

Pharmacokinetics. Indomethacin is rapidly and well absorbed from the GI tract, with a bioavailability of 98%. Peak serum levels are reached within 2 hr with effective concentrations in the range of 0.3–3 mg/L (0.8–8 µmol/L). It is 90% plasma protein bound and has extensive O-demethylation and N-deacylation to inactive metabolites; Vd is 0.29 ± 0.04 L/kg; Cl is 0.084 ± 0.012 L/hr/kg, lower in premature infants, neonates, and the aged; 15 ± 8% is excreted unchanged in the urine. The half-life of the drug is 2.4 ± 0.4 hr, higher in premature infants, neonates, and the aged.

Adverse Reactions. Adverse effects are frequent, and about 20% of patients cannot tolerate the drug. Frontal lobe headache, drowsiness, dizziness, mental confusion, and GI distress are frequent, especially with dosages >100 mg/day; occasional peripheral neuropathy, occult bleeding, and peptic ulcer occur. Pancreatitis, corneal opacities, hepatotoxicity, aplastic anemia, agranulocytosis, thrombocytopenia, aggravation of psychiatric disorders, and allergic reactions are reported rarely. The syndrome of asthma, angioedema, and nasal polyps may be provoked in susceptible patients. Precautions, drug interactions, and monitoring are similar to other NSAIDs. (See Ibuprofen.)

Pharmacology. (See Ibuprofen.)

Administration and Adult Dosage. PO for mild to moderate pain, dysmenorrhea, or acute tendinitis or bursitis (naproxen) 500 mg, followed by 250 mg q 6–8 hr, to a maximum of 1250 mg/day; (naproxen sodium) 550 mg, followed by 275 mg q 6–8 hr, to a maximum of 1375 mg/day. PO for rheumatoid arthritis, osteoarthritis, and ankylosing spondylitis (naproxen) 250–500 mg bid initially, to a maximum of 1500 mg/day for limited periods; (naproxen sodium) 275–550 mg bid or 275 mg q morning and 550 mg q evening initially, to a maximum of 1650 mg/day for limited periods. If no improvement has occurred after 4 weeks of therapy, consider other drug therapy. PO for acute gout (naproxen) 750 mg,
followed by 250 mg q 8 hr until resolved; (naproxen sodium) 825 mg, followed by 275 mg q 8 hr until resolved.

**Special Populations.** *Pediatric Dosage.* PO for juvenile arthritis 10 mg/kg/day in 2 divided doses.

*Geriatric Dosage.* Use minimal effective dosages because the elderly are more susceptible to GI bleeding and acute renal insufficiency.

**Dosage Forms.** *Tab* (naproxen) 250, 375, 500 mg; (naproxen sodium) 220, 275, 550 mg; *EC Tab* (naproxen) 375, 500 mg; *SR Tab* (naproxen sodium) 375, 500, 750 mg; *Susp* (naproxen) 25 mg/mL.

**Patient Instructions.** (See Ibuprofen.)

**Pharmacokinetics.** *Serum Levels.* Trough concentrations >50 mg/L (>217 µmol/L) are associated with response in rheumatoid arthritis.²¹

*Fate.* Rapidly absorbed from the GI tract with a bioavailability of about 99%. Greater than 99.7% plasma protein bound, saturable with increasing dosage, increased with uremia, cirrhosis, and in the elderly, and decreased in rheumatoid arthritis and hypoalbuminemia; *Vd* is 0.16 ± 0.02 L/kg, increased in uremia, cirrhosis, and rheumatoid arthritis. *Cl* is 0.0078 ± 0.0012 L/hr/kg, increased in rheumatoid arthritis, and decreased in uremia; less than 1% is excreted unchanged in urine.²¹

\[ t_{1/2} = 14 ± 1 \text{ hr}, \text{ increased in the elderly}. \]

**Adverse Reactions.** Naproxen can occasionally cause renal dysfunction, particularly in those with pre-existing renal disease, CHF, or cirrhosis. Interstitial nephritis and nephrotic syndrome have been reported.⁷²,⁷³ (See also Ibuprofen.) Contraindications, precautions, drug interactions, and monitoring are similar to other NSAIDs. (See Ibuprofen.)

**SELECTIVE COX-2 INHIBITORS:**

<table>
<thead>
<tr>
<th>CELECOXIB</th>
<th>ROFECOXIB</th>
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<tbody>
<tr>
<td><em>Celebrex</em></td>
<td><em>Vioxx</em></td>
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**Pharmacology.** Inhibition of the COX-2 enzyme isoform is thought to be responsible for the anti-inflammatory effects of NSAIDs, whereas inhibition of COX-1 results in GI and possibly other side effects. A relatively selective COX-2 inhibitor should combine anti-inflammatory, analgesic, and antipyretic efficacies equivalent to older, nonselective NSAIDs with improved safety.⁷⁴

**Administration and Adult Dosage.** (Celecoxib) PO for osteoarthritis 100 mg bid or 200 mg daily; PO for rheumatoid arthritis 100–200 mg bid; PO for familial adenomatous polyposis 400 mg bid. (Rofecoxib) PO for osteoarthritis 12.5–25 mg once daily; PO for acute pain and primary dysmenorrhea 50 mg/day prn, to a maximum of 5 days of consecutive use.

**Special Populations.** *Pediatric Dosage.* (<18 yr) Safety and efficacy not established for either agent.
Geriatric Dosage. (Celecoxib) Dosage adjustment is usually not necessary; however, use the lowest effective dose; (<50 kg) initiate therapy at the lowest recommended dose. (Rofecoxib) dosage adjustment is not necessary; however, initiate with the lowest recommended dose.

Dosage Forms. (Celecoxib) Cap 100, 200 mg. (Rofecoxib) Tab 12.5, 25, 50 mg; Susp 2.5, 5 mg/mL.

Patient Instructions. This drug can cause headache, upset stomach, or diarrhea. Report edema, rash, unusual weight gain, or signs and symptoms of gastrointestinal bleeding to your physician. Avoid products that contain aspirin and non-steroidal anti-inflammatory drugs unless otherwise directed. Take without regard to meals (except take with food if taking celecoxib 400 mg bid).

Missed Doses. If you take this drug on a regular schedule, take a missed dose as soon as you remember. If it is about time for the next dose, take that dose only; do not double the dose or take extra.

Pharmacokinetics. Fate. (Celecoxib) Absolute bioavailability not studied. Peak plasma levels occur in 3 hr. With high-fat meals, peak levels are delayed 1–2 hr with accompanying increases in total absorption of 10–20%; 97% plasma protein bound. Predominantly metabolized hepatically by CYP2C9 to inactive metabolites with <3% excreted unchanged in urine or feces. (Rofecoxib) Rapidly absorbed from the GI tract with bioavailability of 93%. Peak plasma level occurs in 2–3 hr and is delayed 1–2 hr when taken with a high-fat meal, with no effect on peak plasma concentration or extent of absorption; 87% plasma protein bound. Metabolism is predominantly by cytosolic enzymes with minor P450 involvement. Inactive metabolites. Predominantly eliminated via hepatic metabolism with <1% unchanged drug excreted in urine.

\[ t_{1/2} \] (Celecoxib) 11 hr; (rofecoxib) 17 hr.

Adverse Reactions. COX-2 inhibitors can cause GI toxicity, dyspepsia, abdominal pain, nausea, vomiting, and diarrhea at a rate similar to placebo and less than conventional NSAIDs. Renal and liver effects are equivalent to other NSAIDs.\(^{75,76}\) Contraindications. (Celecoxib, rofecoxib) History of aspirin- or NSAID-induced asthma, urticaria, or allergic type reactions. (Celecoxib) allergy to sulfonamides.

Precautions. Use celecoxib and rofecoxib cautiously in patients with pre-existing asthma, renal or hepatic compromise, fluid retention, hypertension, or CHF.

Drug Interactions. NSAIDs can diminish the effects of ACE inhibitors, furosemide, and thiazide diuretics and increase lithium plasma levels. Concurrent use with anticoagulants can increase the risk of bleeding. (Celecoxib) Inhibitors of CYP2C9 (eg, fluconazole) can increase serum concentrations of celecoxib. (Rofecoxib) Increased serum concentrations (23%) and reduced renal clearance of methotrexate. Rifampin decreases rofecoxib serum levels by 50%.

Parameters to Monitor. Monitor for weight gain, renal function during long-term use, and occult blood loss if on concomitant aspirin or anticoagulant therapy.

Notes. Celecoxib 100 or 200 mg bid is as effective as naproxen 500 mg bid for the treatment of osteoarthritis and produces fewer gastroduodenal ulcers than naproxen, diclofenac, or ibuprofen.\(^{76}\) Likewise, rofecoxib 12.5 or 25 mg is as ef-
ective as ibuprofen 800 mg tid and diclofenac 50 mg tid for the treatment of osteoarthritis and produces fewer gastroduodenal ulcers than ibuprofen. Parecoxib (Pharmacia) is an injectable COX-2 inhibitor being studied for the treatment of acute pain. Doses of 20 and 40 mg have been used in clinical trials. It appears to be as effective as injectable ketorolac, but with improved safety. Parecoxib is a water-soluble prodrug of valdecoxib (Pharmacia) which is also pending FDA approval as an oral drug.
<table>
<thead>
<tr>
<th>CLASS AND DRUG</th>
<th>DOSAGE FORMS</th>
<th>ADULT DOSAGE</th>
<th>HALF-LIFE (HR)</th>
<th>COMMENTS</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>ACETIC ACIDS</strong></td>
<td></td>
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<tr>
<td>Diclofenac</td>
<td></td>
<td>PO (pain, dysmenorrhea) (Cataflam) 50 mg tid; PO (arthritis) 100–200 mg/day in 2 doses. PO SR 100 mg once or twice daily (dosages expressed as diclofenac).</td>
<td>1.1 ± 0.2</td>
<td>Although it is unclear whether the risk of hepatotoxicity is any greater than with other NSAIDs, careful monitoring of symptoms and liver function tests is recommended.</td>
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<tr>
<td></td>
<td>Tab (diclofenac potassium) 50 mg</td>
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<tr>
<td></td>
<td>Tab (diclofenac sodium) 50, 75 mg plus misoprostol 200 µg (Arthrotec).</td>
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<tr>
<td></td>
<td>SR Tab (diclofenac sodium) 25, 50, 75, 100 mg</td>
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<td></td>
<td></td>
<td>PO (pain) 200–400 mg q 6–8 hr; PO (arthritis) 600–1200 mg/day in 2–3 divided doses.</td>
<td>7.3 ± 4</td>
<td>Recommended for treatment of osteoarthritis; not as effective as other NSAIDs for rheumatoid arthritis.</td>
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<tr>
<td></td>
<td>Cap 200, 300, mg</td>
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<tr>
<td></td>
<td>Tab 400, 500 mg</td>
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<tr>
<td></td>
<td>SR Tab 500, 600 mg.</td>
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<tr>
<td>Etodolac</td>
<td></td>
<td>PO (gouty arthritis) 100 mg, then 50 mg tid; PO or PR (arthritis) 50–200 mg/day in 3 divided doses. SR in 1–2 doses, can substitute for equal daily dosage of non-SR.</td>
<td>2.4 ± 0.4</td>
<td>See monograph. Associated with a high frequency of CNS effects such as drowsiness, dizziness, mental confusion, and frontal lobe headache.</td>
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<tr>
<td></td>
<td>Cap 25, 50 mg</td>
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<tr>
<td></td>
<td>SR Cap 75 mg</td>
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<tr>
<td></td>
<td>Susp 5 mg/mL</td>
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<td></td>
<td>Supp 50 mg</td>
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<td></td>
<td>Inj 1 mg.</td>
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<tr>
<td>Indomethacin</td>
<td></td>
<td>PO (pain, short term) 10 mg q 4–6 hr prn, to a maximum of 40 mg/day for 5 days (including IM/IV). IM or IV (short-term management of pain) 30 or 60 (IM only) mg once, then 15–30 mg q 6 hr.</td>
<td>4.5</td>
<td>For short-term (up to 5 days) use only. Do not exceed 60 mg/day parenterally in patients 65 yr or older, under 50 kg, or with elevated Crs.</td>
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<tr>
<td></td>
<td>Tab 10 mg</td>
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<tr>
<td>Ketorolac</td>
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</table>
**NONSTEROIDAL ANTI-INFLAMMATORY DRUGS COMPARISON CHART (continued)**

<table>
<thead>
<tr>
<th>CLASS AND DRUG</th>
<th>DOSAGE FORMS</th>
<th>ADULT DOSAGE</th>
<th>HALF-LIFE (HR)</th>
<th>COMMENTS</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Sulindac</strong></td>
<td>Tab 150, 200 mg.</td>
<td>PO (arthritis) 300–400 mg/day in 2 divided doses.</td>
<td>15 ± 4 (active sulfide metabolite)</td>
<td>Purported “renal-sparing” effect has been questioned. Because the active sulfide metabolite has a relatively long half-life, renal effects may not be observed for several days.</td>
</tr>
<tr>
<td>Clinoril</td>
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<tr>
<td><strong>Tolmetin</strong></td>
<td>Cap 400 mg</td>
<td>PO (arthritis) 0.6–1.8 g/day in 3–4 divided doses.</td>
<td>4.9 ± 0.3</td>
<td>Higher frequency of anaphylactoid reactions than other NSAIDs.</td>
</tr>
<tr>
<td>Tolectin Tab 200, 600 mg</td>
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<tr>
<td>Various</td>
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</tr>
<tr>
<td><strong>ANTHRANILIC ACIDS (FENAMATES)</strong></td>
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<tr>
<td><strong>Meclofenamate</strong></td>
<td>Cap 50, 100 mg.</td>
<td>PO (pain) 50 mg q 4–6 hr; PO (arthritis) 200–400 mg/day in 3–4 divided doses.</td>
<td>3</td>
<td>The fenamates as a group are more toxic than other NSAIDs and associated with headache, dizziness, and hemolytic anemia.</td>
</tr>
<tr>
<td>Meclomen</td>
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<tr>
<td>Various</td>
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<tr>
<td><strong>Mefenamic Acid</strong></td>
<td>Cap 250 mg.</td>
<td>PO (pain, dysmenorrhea) 250 mg q 6 hr for up to 1 week.</td>
<td>3</td>
<td>Not recommended; see Meclofenamate Comments.</td>
</tr>
<tr>
<td>Ponstel</td>
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<tr>
<td><strong>NONACIDIC COMPOUNDS</strong></td>
<td></td>
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</tr>
<tr>
<td><strong>Nabumetone</strong></td>
<td>Tab 500, 750 mg.</td>
<td>PO (arthritis) 1–2 g/day in 1–2 doses.</td>
<td>23 ± 4 (active 6-MNA metabolite)</td>
<td>Reported to have less GI toxicity than other NSAIDs; however, additional well-controlled, double-blind studies are needed.</td>
</tr>
<tr>
<td>Relafen</td>
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<tr>
<td>CLASS AND DRUG</td>
<td>DOSAGE FORMS</td>
<td>ADULT DOSAGE</td>
<td>HALF-LIFE (HR)</td>
<td>COMMENTS</td>
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<tr>
<td><strong>OXICAMS</strong></td>
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<tr>
<td>Meloxicam</td>
<td>Tab 7.5 mg.</td>
<td>PO (arthritis) 7.5–15 mg once daily.</td>
<td>20</td>
<td>Less mucosal damage than with piroxicam.</td>
</tr>
<tr>
<td>Mobic</td>
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<tr>
<td>Piroxicam</td>
<td>Cap 10, 20 mg.</td>
<td>PO (arthritis) 20 mg/day in 1–2 doses.</td>
<td>48 ± 8</td>
<td>Based on postmarketing surveillance data, reported to cause about 12 times more GI adverse effects than ibuprofen. High frequency of phototoxic cutaneous eruptions.</td>
</tr>
<tr>
<td>Feldene</td>
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<tr>
<td>Various</td>
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<tr>
<td><strong>PROPIONIC ACIDS</strong></td>
<td></td>
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</tr>
<tr>
<td>Fenoprofen</td>
<td>Cap 200, 300 mg</td>
<td>PO (pain) 200 mg q 4–6 hr; PO (arthritis) 1.2–2.4 g/day in 3–4 divided doses.</td>
<td>2.5 ± 0.5</td>
<td>Similar to ibuprofen.</td>
</tr>
<tr>
<td>Nalfon</td>
<td>Tab 600 mg.</td>
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</tr>
<tr>
<td>Flurbiprofen</td>
<td>Tab 50, 100 mg.</td>
<td>PO (arthritis) 200–300 mg/day in 2–4 divided doses.</td>
<td>3.8 ± 1.2</td>
<td>Similar to ibuprofen.</td>
</tr>
<tr>
<td>Ansaid</td>
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</tr>
<tr>
<td>Ibuprofen</td>
<td>Cap 200, 400 mg</td>
<td>PO (pain, dysmenorrhea) 400 mg q 4–6 hr; PO (arthritis) 1.2–3.2 g/day in 3–4 divided doses.</td>
<td>2 ± 0.5</td>
<td>See monograph.</td>
</tr>
<tr>
<td>Advil</td>
<td>Chew Tab 50, 100 mg</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Motrin</td>
<td>Tab 100, 200, 400, 600, 800 mg</td>
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<tr>
<td>Nuprin</td>
<td>Drp 40 mg/mL</td>
<td></td>
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<tr>
<td>Various</td>
<td>Susp 20, 40 mg/mL</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Ketoprofen</td>
<td>Cap 25, 50, 75 mg</td>
<td>PO (pain) 25–50 mg q 6–8 hr; PO (arthritis) 150–300 mg/day in 3 divided doses.</td>
<td>1.8 ± 0.3</td>
<td>Similar to ibuprofen.</td>
</tr>
<tr>
<td>Orudis</td>
<td>Tab 12.5 mg.</td>
<td></td>
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<tr>
<td>Oruvail</td>
<td>SR Cap 100, 150, 200 mg.</td>
<td></td>
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<tr>
<td>Various</td>
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</tbody>
</table>
### NONSTEROIDAL ANTI-INFLAMMATORY DRUGS COMPARISON CHART (continued)

<table>
<thead>
<tr>
<th>CLASS AND DRUG</th>
<th>DOSAGE FORMS</th>
<th>ADULT DOSAGE</th>
<th>HALF-LIFE (HR)</th>
<th>COMMENTS</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Naproxen</strong></td>
<td>Tab (naproxen sodium) 220, 275, 550 mg</td>
<td>PO (pain) 500 mg, then 250 mg q 6–8 hr; PO (arthritis) 0.5–1.5 g/day in 2 divided doses.</td>
<td>14 ± 1</td>
<td>See monograph. Equal in efficacy and safety to ibuprofen</td>
</tr>
<tr>
<td>Aleve</td>
<td>Tab (naproxen) 250, 375, 500 mg</td>
<td></td>
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<tr>
<td>Anaprox</td>
<td>EC Tab (naproxen) 375, 500 mg</td>
<td>PO (acute gout) 750 mg, then 250 mg q 8 hr. (Doses expressed as naproxen.)</td>
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<tr>
<td>Naprelan</td>
<td>SR Tab (naproxen) 375, 500, 750 mg</td>
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<tr>
<td>Naprosyn</td>
<td>Susp (naproxen) 25 mg/mL.</td>
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</tr>
<tr>
<td>Various</td>
<td>Tab 600 mg.</td>
<td>PO (arthritis) 1.2 g/day in 1 dose.</td>
<td>50–60</td>
<td>Similar to other NSAIDs.</td>
</tr>
<tr>
<td><strong>Oxaprozin</strong></td>
<td>Tab 600 mg.</td>
<td>PO (arthritis) 1.2 g/day in 1 dose.</td>
<td>50–60</td>
<td>Similar to other NSAIDs.</td>
</tr>
</tbody>
</table>

### SALICYLATES

<table>
<thead>
<tr>
<th>Class and Drug</th>
<th>Dosage Forms</th>
<th>Adult Dosage</th>
<th>Half-Life (HR)</th>
<th>Comments</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Aspirin</strong></td>
<td>See monograph.</td>
<td>PO (pain) 325–1000 mg q 4 hr; PO (arthritis) 3.6–5.4 g/day in 3–4 divided doses.</td>
<td>0.25 ± 0.03 (aspirin)</td>
<td>See monograph.</td>
</tr>
<tr>
<td>Various</td>
<td></td>
<td></td>
<td>2–19 (salicylate, dose dependent)</td>
<td></td>
</tr>
<tr>
<td><strong>Choline Magnesium</strong></td>
<td>Tab 500, 750 mg, 1g</td>
<td>PO (pain, arthritis) 1.5–3 g/day in 1–2 divided doses.</td>
<td>2–19 (salicylate, dose dependent)</td>
<td>Salicylate is only a weak inhibitor of cyclo-oxygenase. It therefore has no antiplatelet effect and can usually be administered safely to individuals with aspirin sensitivity. See also Aspirin monograph.</td>
</tr>
<tr>
<td><strong>Trisalicylate</strong></td>
<td>Liquid 100 mg/mL.</td>
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<tr>
<td>Trilisate</td>
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<tr>
<td><strong>Diflunisal</strong></td>
<td>Tab 250, 500 mg.</td>
<td>PO (arthritis) 250–500 mg bid.</td>
<td>11 ± 2 (dose dependent)</td>
<td>Not converted to salicylate; similar to other NSAIDs.</td>
</tr>
<tr>
<td>Dolobid</td>
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<tr>
<td>Various</td>
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(continued)
<table>
<thead>
<tr>
<th>CLASS AND DRUG</th>
<th>DOSAGE FORMS</th>
<th>ADULT DOSAGE</th>
<th>HALF-LIFE (HR)</th>
<th>COMMENTS</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Magnesium Salicylate</strong></td>
<td>Tab 500, 545, 600 mg.</td>
<td>PO (pain, arthritis) 3.6–4.8 g/day in 3–4 divided doses.</td>
<td>2–19 (salicylate, dose dependent)</td>
<td>See Choline Magnesium Trisalicylate comments and Aspirin monograph.</td>
</tr>
<tr>
<td>Doan’s</td>
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<td>Various</td>
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<tr>
<td><strong>Salsalate</strong></td>
<td>Cap 500 mg</td>
<td>PO (arthritis) 3 g/day in 2–3 divided doses.</td>
<td>2–19 (salicylate, dose dependent)</td>
<td>See Choline Magnesium Trisalicylate comments and Aspirin monograph.</td>
</tr>
<tr>
<td>Disalcid</td>
<td>Tab 500, 750 mg.</td>
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<tr>
<td>Various</td>
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<tr>
<td><strong>SELECTIVE COX-2 INHIBITORS</strong></td>
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<tr>
<td>Celecoxib</td>
<td>Cap 100, 200.</td>
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<tr>
<td>Celebrex</td>
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</tr>
<tr>
<td><strong>Rofecoxib</strong></td>
<td>Tab 12.5, 25, 50 mg</td>
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<tr>
<td>Vioxx</td>
<td>Susp 2.5, 5 mg/mL.</td>
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</tbody>
</table>

*Long-term dosage for arthritis should be guided by serum salicylate levels; see Aspirin monograph.

Adapted from references 21, 61, 72, 73, and 77–85, and product information.
Class Instructions. This drug can cause drowsiness. Until the extent of this effect is known, use caution when driving, operating machinery, or performing other tasks requiring mental alertness. Avoid excessive concurrent use of alcohol and other drugs that cause drowsiness. Prolonged use of this drug can cause constipation, and concurrent use of a stool-softening or stimulant laxative may be helpful.

For moderate to severe pain (pain rating >5 on a 0–10 scale), you must take doses at regular intervals around the clock to anticipate and prevent pain. When the drug is taken at the correct interval and pain relief does not last for this period, use additional “rescue” doses of a short-acting drug to maintain pain relief. When more than 4 rescue doses are used in a day, contact the prescriber for a dosage increase. Addiction does not occur when these drugs are used for legitimate painful conditions. Dependence, a condition in which the body may go through withdrawal when the drug is stopped suddenly, can occur with prolonged usage but can be managed by slowly decreasing the dosage when the drug is no longer needed.

Missed Doses. If you miss a dose, take it as soon as you remember. If it is about time for the next dose, take that dose only. Do not double the dose or take extra. Take subsequent doses at the same interval previously established for pain relief.

Pharmacology. Codeine is 3-methoxymorphine, a phenanthrene opioid with very low affinity for opioid receptors. Its analgesic activity appears to result from conversion to morphine. Poor metabolizers of debrisoquine/sparteine (approximately 7% of the Caucasian population) cannot convert appreciable amounts of codeine to morphine or obtain analgesia from codeine but are still subject to the same adverse effects. (See Morphine Sulfate.)

Administration and Adult Dosage. PO, SC, or IM for analgesia 15–60 mg q 4–6 hr. PO or SC for antitussive action 10–20 mg q 4–6 hr, to a maximum of 120 mg/day. IV not recommended. (See Precautions.)

Special Populations. Pediatric Dosage. PO, SC, or IM for analgesia (≥1 yr) 0.5 mg/kg q 4–6 hr. PO for antitussive action (2–6 yr) 2.5–5 mg q 4–6 hr, to a maximum of 30 mg/day; (7–12 yr) 5–10 mg q 4–6 hr, to a maximum of 60 mg/day; (>12 yr) same as adult dosage. (See Notes.)

Geriatric Dosage. Same as adult dosage.90

Other Conditions. Reduce initial dosage in debilitated patients or those with hypoxia or hypercapnia.

Dosage Forms. Tab 15, 30, 60 mg; Inj 15, 30, 60 mg/mL; Oral Liquid 2, 2.4, 3 mg/mL in various combinations. Formulated as phosphate or sulfate salt.

Patient Instructions. (See Opioids Class Instructions.)

Pharmacokinetics. Onset and Duration. PO, SC onset 15–30 min; IM peak analgesia 0.5–1 hr; duration (all routes) 4–6 hr.91

Fate. Systemic availability averages 40% but with a wide range (12–84%), reflecting large variability in hepatic enzyme activity.92 A single PO 15 mg dose
produces serum levels of 26–33 µg/L (82–104 nmol/L) in 2 hr and 13–22 µg/L (41–69 nmol/L) in 5 hr. The drug is 7% plasma protein bound. Vd is 2.6 ± 0.3 L/kg; Cl is 0.66 ± 0.12 L/hr/kg. Metabolized in the liver to codeine-6-glucoronide, N-demethylated to norcodeine, and O-demethylated to morphine by genetic polymorphic CYP2D6. Codeine-6-glucuronide is the major metabolite, and norcodeine and morphine are minor metabolites, each accounting for approximately 10% of the dose. Accumulation of morphine occurs with repeated administration, resulting in a morphine:codeine AUC ratio of 0.29:1. Variation in the reported rates of codeine conversion to morphine may be related to the assays used, with much higher concentrations of morphine reported with radioimmunoassays than with HPLC or GC-MS. Primarily urinary excretion of inactive forms; 3–16% is excreted unchanged in urine.

\[ t_{1/2} \] = 2.9 ± 0.7 hr.

**Adverse Reactions.** Sedation, dizziness, nausea, vomiting, constipation, and respiratory depression occur frequently. Dose-related signs of intoxication are miosis, drowsiness, decreased rate and depth of respiration, bradycardia, and hypotension. Dose-related adverse reactions in children are somnolence, ataxia, miosis, and vomiting at 3–5 mg/kg/day and respiratory depression at >5 mg/kg/day. Because hepatic glucuronidation is incomplete in infants, they are at particular risk for dose-related adverse effects.

**Precautions.** Because it can cause severe hypotension, do not administer codeine phosphate IV.

**Drug Interactions.** Potent CYP2D6 inhibitors (eg, quinidine, fluoxetine) can abolish the conversion to morphine and the pharmacologic effects of codeine.

**Notes.** Codeine is no more effective than placebo in suppressing nighttime cough in children. The American Academy of Pediatrics recommends that parents be educated about the lack of proven antitussive effects and the potential risks of codeine-containing products because overdosage has been reported.

**FENTANYL**

**Pharmacology.** Fentanyl is a phenylpiperidine opioid agonist with predominant effects on the mu opioid receptor and is about 50–100 times more potent as an analgesic than morphine. Other related compounds are sufentanil (Sufenta), which is 5–7 times more potent than fentanyl; alfentanil (Alfenta), which is less potent than fentanyl but acts more rapidly and has a shorter duration of action; and remifentanil (Ultiva), which is more potent than fentanyl and is extremely short acting because of its rapid ester hydrolysis. (See Morphine Sulfate.)

**Administration and Adult Dosage.** IV patient-controlled analgesia (PCA) 20–100 µg per activation with 3–10-min lockout period, both titrated to patient response. (See Patient-Controlled Analgesia Guidelines Chart, page 44.) Epidurally for analgesia 25–150 µg as an intermittent bolus dose or 25–150 µg/hr as a continuous infusion, titrated to patient response. (See Notes and Intraspinal Narcotic Administration Guidelines Chart, page 44.) Transdermal for analgesia calculate the previous 24-hr analgesic requirement and convert this amount to the equal
analgesic oral morphine dosage from the Opioid Analgesics Comparison Chart. A short-acting opioid or the fentanyl lozenge (Actiq) must be used for control of breakthrough pain until sufficient transdermal fentanyl is absorbed to achieve adequate analgesia. Use the following table to determine the fentanyl transdermal dosage from the daily equivalent oral morphine dosage:

<table>
<thead>
<tr>
<th>24-HR ORAL MORPHINE DOSAGE (MG/DAY)</th>
<th>FENTANYL TRANSDERMAL DOSAGE (µG/HR)</th>
</tr>
</thead>
<tbody>
<tr>
<td>45–134</td>
<td>25</td>
</tr>
<tr>
<td>135–224</td>
<td>50</td>
</tr>
<tr>
<td>225–314</td>
<td>75</td>
</tr>
<tr>
<td>315–404</td>
<td>100</td>
</tr>
<tr>
<td>405–494</td>
<td>125</td>
</tr>
<tr>
<td>495–584</td>
<td>150</td>
</tr>
<tr>
<td>585–674</td>
<td>175</td>
</tr>
<tr>
<td>675–764</td>
<td>200</td>
</tr>
<tr>
<td>765–854</td>
<td>225</td>
</tr>
<tr>
<td>855–944</td>
<td>250</td>
</tr>
<tr>
<td>945–1034</td>
<td>275</td>
</tr>
<tr>
<td>1035–1124</td>
<td>300</td>
</tr>
</tbody>
</table>

Assumes morphine 10 mg IM is equivalent to morphine 60 mg orally; however, because of individual variability, equivalent dosages can vary among patients. These conversion dosages are conservative, and approximately 50% of patients are likely to require a dosage increase after initial application. (See Opioid Analgesics Comparison Chart.)

Initiate treatment using the recommended transdermal fentanyl dosage and increase based on response no more frequently than q 3–6 days. Multiple transdermal patches can be used to achieve appropriate dosage (do not cut patches for a partial dosage). To change treatment to another opioid, discontinue the transdermal patch for 12–18 hr and start treatment with the new opioid at about one-half the equianalgesic dosage. **IV for induction and maintenance anesthesia** (loading) 4–20 µg/kg, (maintenance) 2–10 µg/kg/hr, (additional bolus) 25–100 µg. **IV for postoperative (recovery room) pain control** 50–100 µg q 1–2 hr as needed; **Lozenge (Oralet) for anesthesia premedication or induction of conscious sedation** 5 µg/kg (provides effects similar to 0.75–1.25 µg/kg given IM), to a maximum of 400 µg. **Lozenge for the management of breakthrough cancer pain (Actiq) in patients already receiving >60 mg of oral morphine/day or >50 µg/hr of transdermal fentanyl** initial dose of 200 µg. Until the appropriate dose is reached, an additional dose can be used to treat an episode of breakthrough pain. Re-administration can start 15 min after the previous lozenge has been com-
pleted. Do not give >2 units for a breakthrough pain episode while a patient is in the titration phase. Evaluate each new dose in the titration period over several breakthrough pain episodes. If >4 units/day are needed, increase the dosage of the long-acting opioid.

**Special Populations. Pediatric Dosage.** IV for sedation in neonates 9–20 µg/kg/hr; tolerance limits its usefulness for prolonged sedation.101 IV for induction and maintenance anesthesia (2–12 yr) 2–3 µg/kg initially, followed by 1–5 µg/kg/hr. Lozenge for anesthesia premedication or induction of conscious sedation (<15 kg) contraindicated; (≥15 kg) 5–15 µg/kg, to a maximum of 400 µg.

**Geriatric Dosage.** Lozenge for anesthesia premedication or induction of conscious sedation (>65 yr) 2.5–5 µg/kg, to a maximum of 400 µg. Altered pharmacodynamics rather than pharmacokinetics appear to be responsible for increased sensitivity in elderly patients.102

**Other Conditions.** In patients with head injury, cardiovascular, pulmonary, or hepatic disease, consider a lower dosage of 2.5–5 µg/kg, to a maximum of 400 µg.

**Dosage Forms.** Inj 50 µg/mL; SR Patch 25, 50, 75, 100 µg/hr; Lozenge for anesthesia (Oralet) 100, 200, 300, 400 µg; Lozenge (on a stick) for breakthrough cancer pain (Actiq) 200, 400, 600, 800, 1200, 1600 µg.

**Patient Instructions.** (See Opioids Class Instructions.) (Fentanyl Actiq) once an effective dosage is determined, limit consumption to ≤4 units/day.

**Pharmacokinetics. Onset and Duration.** IM onset 7–15 min; duration 1–2 hr. Epidural onset 5 min; duration 4–6 hr.91 Transdermal onset 6–8 hr; peak 24–72 hr; duration after a single application 72 hr.103,104 More than 17 hr is required for serum levels to fall by one-half after patch removal.

**Serum Levels.** (Analgesia) 1–3 µg/L (3–9 nmol/L);103,104 (balanced anesthesia) 6–20 µg/L (18–60 nmol/L).100

**Fate.** Bioavailability is 52% with lozenge. Of the fentanyl released by the transdermal system, 92% is absorbed, but overall systemic bioavailability of the transdermal preparation is approximately 30%. The drug is 84 ± 2% plasma protein bound; it is metabolized rapidly primarily by the liver to norfentanyl and other inactive metabolites; Vd is 4 ± 0.4 L/kg; Cl is 0.78 ± 0.12 L/hr/kg, decreased in the elderly and increased in neonates and children. Pharmacokinetics are not altered in renal insufficiency or compensated hepatic cirrhosis. Less than 10% is excreted unchanged in the urine,86,102–105 t ½ 6.1 ± 2 hr,105 7.1–11 hr during cardiopulmonary bypass surgery.100

**Adverse Reactions.** (See Morphine Sulfate.) Unlike other opioids, fentanyl, alfentanil, remifentanil, and sufentanil are not associated with histamine release and may be preferable when cardiovascular stability is an issue.100 The frequency of pruritus is lower than that of morphine but not as low as that of meperidine.106,107 PCA fentanyl produces less depression of postoperative cognitive function in elderly patients than does PCA morphine.108 Development of withdrawal reactions after use for sedation in neonates and infants is likely with a total dosage >2.5 mg/kg or duration of infusion >9 days.109
**Contraindications.** (See Morphine Sulfate.) (Fentanyl SR patch) acute or postoperative pain, including outpatient surgery; patients <12 yr or <50 kg; pain that can be managed by conventional analgesics; and doses >25 μg/hr at the initiation of opioid therapy. (Oralet) management of acute and chronic pain. (Actiq) management of acute or postoperative pain.

**Precautions.** (See Morphine Sulfate.) Analyses of fentanyl transdermal systems after 3 days of continuous application demonstrated a considerable amount of remaining drug (28–84%), which is a potentially lethal dose (1036 μg) for a 70-kg individual. Cutting the membrane-controlled fentanyl transdermal system to achieve a different dosage is not recommended because it can damage the integrity of the semipermeable membrane. Placing a piece of impermeable material (eg, adhesive bandage) on the skin proportionate in surface area to the intended reduction in dosage may be effective.

**Drug Interactions.** (See Morphine Sulfate.) The effects of fentanyl may be potentiated by other CNS depressant drugs (eg, barbiturates, general anesthetics, narcotics, and tranquilizers) and ritonavir, the latter by inhibition of CYP2D6. Carbamazepine may decrease fentanyl’s effect during anesthesia for craniotomy.

**Parameters to Monitor.** Monitor vital signs and pain ratings routinely.

**Notes.** Epidural administration has not been shown to be more advantageous than IV administration during surgery. Lack of rapid titratability precludes the usefulness of the transdermal fentanyl system for pain control in patients with rapidly changing analgesic requirements. Transdermal fentanyl for cancer pain causes a lower frequency of constipation than SR morphine. IV is the parenteral route of choice after major surgery. This route is suitable for titrated bolus or continuous administration but requires close monitoring because there is a great risk of respiratory depression with inappropriate dosage.

**MEPERIDINE HYDROCHLORIDE**

**Pharmacology.** Meperidine is a phenylpiperidine opioid agonist with important antimuscarinic activity and negative inotropic effects on the heart. Its major metabolite, normeperidine, has excitant effects that can precipitate tremors, myoclonus, or seizures. Meperidine’s antimuscarinic activity might negate the miosis that occurs with other opioids. (See Morphine Sulfate.)

**Administration and Adult Dosage.** PO, IV, or SC for analgesia 50–150 mg q 3–4 hr. (See Notes.) Oral doses are about one-half as effective as parenteral doses. Reduce dosage when given concomitantly with a phenothiazine or other drugs that potentiate the depressant effects of meperidine. IV for shaking caused by general anesthesia or amphotericin B 25–50 mg. (See Notes.) IM not recommended.

**Special Populations.** Pediatric Dosage. PO, IV, or SC for analgesia 1–1.8 mg/kg q 3–4 hr, to a maximum of 100 mg/dose. IM painful and should not be used in children. (See Notes.)

**Geriatric Dosage.** Same as adult dosage.
**Dosage Forms.** Syrup 10 mg/mL; Tab 50, 100 mg; Inj 10, 25, 50, 75, 100 mg/mL.

**Patient Instructions.** (See Opioids Class Instructions.)

**Pharmacokinetics.** **Onset and Duration.** PO onset about 15 min; duration 2–3 hr. SC or IM onset about 10 min; peak analgesia 0.5–1 hr; duration 2–3 hr.\(^{19,91}\)

**Serum Levels.** 500–700 µg/L (2–2.8 µmol/L) appear to be required for analgesia.\(^{103}\)

**Fate.** Oral bioavailability is about 52 ± 3%, increasing to 80–90% in cirrhosis caused by decreased first-pass metabolism.\(^{19,117}\) After a single 100 mg IM dose, mean serum levels of 670 µg/L (2.7 µmol/L) and 650 µg/L (2.6 µmol/L) are attained in 1 and 2 hr, respectively\(^{18,119}\); 58 ± 9% plasma protein bound, largely to \(\alpha_1\)-acid glycoprotein; decreased in the elderly and in uremia.\(^{19,120}\) \(V_d\) is 4.4 ± 0.9 L/kg, increased in the elderly and premature infants; \(Cl\) is 1.02 ± 0.3 L/hr/kg, reduced by 25% in surgical patients and 50% in cirrhosis, and reduced in acute viral hepatitis.\(^{19}\) Hydrolyzed and metabolized in the liver to normeperidine (an active metabolite), which is also hydrolyzed. An average of 2% unchanged drug and 1–21% (average 6%) normeperidine are excreted in urine.\(^{120}\)

\(t_{1/2}\). (Meperidine) \(\alpha\) phase 12 min, \(\beta\) phase 3.2 hr, increasing to 7 hr in patients with cirrhosis or acute liver disease and 14–21 hr in patients with moderate to severe renal dysfunction.\(^{110,112,122}\) (Normeperidine) 14–21 hr in normals, increasing to 35 hr in renal failure.\(^{123}\)

**Adverse Reactions.** (See Morphine Sulfate.) Factors that can predispose to normeperidine-induced seizures are dosage >400–600 mg/day, renal failure, history of seizures, long-term administration to cancer patients, and co-administration of agents that increase \(N\)-demethylation to normeperidine.\(^{124}\) (See Drug Interactions.) Local irritation and induration occur with repeated SC injection.

**Contraindications.** MAO inhibitors within the past 14–21 days; chronic pain.

**Precautions.** (See Morphine Sulfate.) Avoid in patients with reduced renal function and avoid continuous administration for more than a few days. The combination of meperidine with promethazine and chlorpromazine (DPT) for painful procedures is not recommended because it has poor efficacy compared with alternative approaches and is associated with a high frequency of adverse effects.\(^{115}\)

**Drug Interactions.** (See Morphine Sulfate.) Concurrent use with an MAO inhibitor can cause marked blood pressure alterations, sweating, excitation, and rigidity. Barbiturates, chlorpromazine, and phenytoin can decrease meperidine serum concentrations and increase normeperidine, reducing analgesia and increasing the risk of stimulation and seizures.\(^{125}\) Ritonavir can increase meperidine AUC via CYP2D6 inhibition.\(^{112}\)

**Parameters to Monitor.** Monitor vital signs and pain scores at regular intervals. Jerking and twitching movements may be signs of normeperidine accumulation and impending toxicity.\(^{126}\)

**Notes.** All opioids including meperidine and morphine increase biliary tract pressure. Sphincter of Oddi spasm may be less with meperidine than with morphine, but there is little evidence that this has clinical relevance. Unlike other opioids,
meperidine is useful in treating the shaking and shivering associated with general anesthesia or amphotericin B administration. Because of its low therapeutic index, reserve meperidine for very brief courses in otherwise healthy patients who have demonstrated untoward effects during treatment with other opioids such as morphine or hydromorphone. Because of its unreliable absorption and breakthrough pain when meperidine is administered IM, more rapid and predictable routes (eg, IV) are recommended. Oral meperidine is not recommended for cancer pain because the high dosage required to relieve severe pain increases the risk of CNS toxicity.

**METHADONE HYDROCHLORIDE**

**Pharmacology.** Methadone is a phenylethylamine opioid agonist qualitatively similar to morphine but with a chemical structure unrelated to the alkaloid-type structures of the opium derivatives. Analgesic activity of (R)-methadone is 8–50 times that of (S)-methadone, and (R)-methadone has a 10-fold higher affinity for opioid receptors. Methadone is lipophilic and has considerable tissue distribution; plasma concentrations during long-term treatment are sustained by this peripheral reservoir. It does not share cross-tolerance with other opioids, and the dosage required to achieve analgesia in opioid-tolerant patients is much lower than predicted by opioid conversion tables and single-dose studies. Unlike other opioids, methadone does not have active or toxic metabolites that are associated with CNS toxicity (eg, myoclonus, seizures). Because methadone is a long-acting narcotic agent, it can be substituted for short-acting narcotic agents for analgesia maintenance and detoxification. Methadone abstinence syndrome is similar to morphine; however, onset is slower and duration is longer. (See Morphine Sulfate.)

**Administration and Adult Dosage.** PO, IV, or SC for pain 5–80 mg/day in 1–3 divided doses. Dosage escalation is slower than with other opioids and averages approximately 2%/day. PO for maintenance and detoxification treatment the minimum effective dosage for reducing illicit heroin use is approximately 60 mg/day, and the optimum dosage range is 80–120 mg/day. Premature termination of treatment and use of suboptimal dosages remain common problems. If tapering is attempted, taper gradually over 4–12 months or longer. To convert from another opioid decrease the previous opioid dosage by one-third over 24 hr and replace it with methadone using a dosage ratio of 1 mg oral methadone = 10 mg oral morphine. During day 2, attempt another one-third decrease in the dosage of the previous opioid; on day 3, the final one-third of the dosage of the previous opioid may be discontinued. Maintain the patient on an q-8-hr schedule with approximately 10% of the daily methadone dosage as an extra dose for breakthrough pain.

**Special Populations.** Pediatric Dosage. IV for pain 0.1 mg/kg q 6–8 hr; PO for pain 0.2 mg/kg q 6–8 hr.

**Geriatric Dosage.** Same as adult dosage.

**Dosage Forms.** Tab 5, 10 mg; Dispersible Tab 40 mg; Soln 1, 2, 10 mg/mL; Inj 10 mg/mL; Pwdr 50, 100, 500, 1000 g.

**Patient Instructions.** (See Opioids Class Instructions.) Increase dosage cautiously with the assistance of your clinician.
Pharmacokinetics. Onset and Duration. (Analgesia) onset SC 10–20 min, PO 30–60 min; peak SC 0.5–1 hr; duration PO, SC, or IV 4–5 hr after a single dose, 8–48 hr with multiple doses.\(^\text{103,129,130}\)

Serum Levels. Best rehabilitation in methadone maintenance patients has been associated with serum levels >211 µg/L (682 nmol/L).\(^\text{132}\) There is no good correlation between serum levels and analgesia.\(^\text{130}\)

Fate. Oral bioavailability is 92 ± 21%; 89% plasma protein bound. Pharmacokinetics are best described by a 2-compartment model. \(V_{d}/H_{9252}\) is 3.8 ± 0.6 L/kg; Cl is 0.084 ± 0.03 L/hr/kg. Both \(V_{d}/H_{9252}\) and Cl are greater for (\(R\))-methadone.\(^\text{130}\) Extent of metabolism may increase with long-term therapy, resulting in a 15–25% decline in serum levels, although this has also been attributed to poor compliance. Metabolized in the liver to inactive metabolites via N-demethylation; metabolites are excreted in urine and bile.\(^\text{129}\) The drug is 24 ± 10% excreted unchanged in the urine, increased by urine acidification.\(^\text{19,132,133}\)

\(t_{1/2}\) phase 35 ± 12 hr;\(^\text{19}\) (\(R\))-methadone has a longer half-life (37.5 hr) than (\(S\))-methadone (28.6 hr).\(^\text{130}\)

Adverse Reactions. (See Morphine Sulfate.) Because of its long half-life and lack of cross-tolerance, patients receiving methadone are at greater risk for toxicity when inappropriate dosage increases are made.

Precautions. (See Morphine Sulfate.) The process of switching from another opioid to methadone is complex and should only be attempted by an experienced clinician in an inpatient setting over 3–6 days. (See Administration and Adult Dosage.)\(^\text{129}\)

Drug Interactions. (See Morphine Sulfate.) Carbamazepine, phenytoin, rifampin, and other drugs that induce CYP3A4 can decrease methadone serum levels and result in withdrawal symptoms in patients on methadone maintenance programs. Diazepam, erythromycin, fluvoxamine, ritonavir, and possibly other enzyme inhibitors can increase methadone levels and effects.\(^\text{125,134}\)

Parameters to Monitor. During analgesia, monitor vital signs and pain ratings routinely. During methadone maintenance, monitor for signs of withdrawal, which include lacrimation, rhinorrhea, diaphoresis, yawning, restlessness, insomnia, dilated pupils, and piloerection.\(^\text{131}\)

Notes. For treatment of narcotic addiction in detoxification or maintenance programs, methadone may be dispensed only by approved pharmacies. Maintenance therapy (treatment for longer than 3 weeks) may be undertaken only by approved methadone programs; this does not apply to addicts hospitalized for other medical conditions.

**MORPHINE SULFATE** Various

Pharmacology. Morphine and other opioids interact with stereospecific opiate receptors in the CNS and other tissues. (See Opioid Receptor Specificity Comparison Chart.) Opioid analgesia is caused by actions at several CNS sites. Morphine and other mu opioid agonists inhibit nociceptive reflexes through inhibition of neurotransmitter release, have inhibitory actions on neurons conveying nocicep-
tive information to higher brain centers, and enhance activity in descending pathways that exert inhibitory effects on the processing of nociceptive information in the spinal cord. Mu receptors are responsible for analgesia, respiratory depression, miosis, decreased GI motility, and euphoria. Stimulation of kappa receptors results in analgesia, less intense miosis and respiratory depression, dysphoria, and psychotomimetic effects. It is unclear what the consequences of delta receptor stimulation are in humans. The relief of pain is fairly specific; other sensory modalities are essentially unaffected, and mental processes are not impaired (unlike anesthetics), except when given in large doses or to opiate-naive individuals. These drugs also have antitussive effects, usually at dosages less than those required for analgesia.

**Administration and Adult Dosage.** With the exception of transdermal fentanyl, there is no ceiling or maximum dosage for morphine or other opioid agonists, and very large doses may be required for severe pain. PO for analgesia 8–20 mg q 4 hr; SR Tab, 12-hr (narcotic-naive patients) 30 mg q 8–12 hr initially; (narcotic-tolerant patients) total daily oral morphine dosage equivalent in 2 divided doses q 12 hr; SR Cap, 24-hr (narcotic-naive patients) 20 mg q 24 hr initially; (narcotic-tolerant patients) total daily oral morphine dosage equivalent q 24 hr; SC for analgesia 5–15 mg q 4 hr (10 mg/70 kg is the optimal initial dose); PR for analgesia 10–20 mg q 4 hr. IV for analgesia 4–10 mg, dilute and inject slowly over a 2–3-min period. IV infusion 1–10 mg/hr; IV PCA 1 mg per activation initially with 5–20 min lockout period, both titrated to patient response. Continuous infusion combined with PCA is effective in chronic cancer pain. Epidural for analgesia (unpreserved solution) (intermittent) 5 mg initially, may repeat with 1–2 mg after 1 hr; (continuous infusion) 0.05–0.1 mg/kg loading dose, then 0.005–0.01 mg/kg/hr. IT for cancer pain (unpreserved solution) 0.4–8.3 mg/day (average 1–23 mg/day); IT for cesarean section (unpreserved solution) 0.1 mg. Intraventricular (unpreserved solution) 0.1–2 mg, repeated approximately q 24 hr. Inh for dyspnea 5–15 mg in 2 mL sterile water or NS via nebulizer q 4 hr. IM is painful and is not recommended.

**Special Populations. Pediatric Dosage.** PO 0.3 mg/kg q 3–4 hr. IV 0.05–0.2 mg/kg q 4 hr. IV infusion 0.01–0.04 mg/kg/hr. Epidural 0.05–0.08 mg/kg. IT 0.01–0.03 mg/kg.

**Geriatric Dosage.** Reduce initial dosage in elderly patients and make smaller percentage incremental increases in total daily dosage (eg, 25%) than in younger patients.

**Other Conditions.** Reduce initial dosage in debilitated patients.

**Dosage Forms.** Cap 15, 30 mg; Soln 2, 4, 20 mg/mL; Supp 5, 10, 20, 30 mg; Tab 10, 15, 30 mg; SR Tab (8, 12 hr) 15, 30, 60, 100, 200 mg; SR Cap (24 hr) 20, 50, 100 mg; Inj (unpreserved solution) 0.5, 1, 10, 25, 50 mg/mL; (preserved solution) 2, 3, 4, 5, 8, 10, 15, 25, 50 mg/mL.

**Patient Instructions.** (See Opioids Class Instructions.)

**Pharmacokinetics. Onset and Duration.** (Analgesia) onset IM 10–30 min; peak 0.5–1 hr; duration 3–5 hr.
Serum Levels. It is speculated that moderate analgesia requires serum levels of at least 50 µg/L (88 nmol/L).

Fate. Well absorbed from the GI tract, but first-pass conjugation is extensive, reducing oral bioavailability to 24 ± 12%. Nebulized morphine by inhalation has a low bioavailability, 5 ± 3%, but a rapid peak at 10 min. After an IM dose of 10 mg, peak morphine levels of about 56 µg/L (98 nmol/L) are reached within 20 min. The drug is 35 ± 2% plasma protein bound and decreased in acute viral hepatitis, cirrhosis, and hypoalbuminemia. V_d is 2.12 L/kg in young normals and 1.16 L/kg in elderly patients; Cl is 2.02 L/hr/kg in young normals and 1.66 L/hr/kg in elderly patients. Morphine clearance reaches adult level by age 6 months–2.5 yr. Inactivated in the liver, primarily by conjugation to morphine–6–glucuronide (active) and morphine–3–glucuronide (inactive or antagonistic). Decreased clearance of glucuronide metabolites has been demonstrated in patients with renal insufficiency. Greater plasma concentrations of morphine–6–glucuronide are present with oral than with parenteral administration. Mostly excreted in urine; 14 ± 7% as the active morphine–6–glucuronide and 3.4% (oral) to 9% (parenteral) of a dose is excreted unchanged.

t_1/2. 1.9 ± 0.5 hr, increased in neonates and premature infants.

Adverse Reactions. Respiratory and circulatory depression and constipation are major adverse effects. Patients with renal failure are more prone to develop adverse reactions. Dose-related signs of intoxication are miosis, drowsiness, decreased rate and depth of respiration, bradycardia, and hypotension. Sedation, dizziness, nausea, vomiting, sweating, and constipation occur frequently. Euphoria, dysphoria, dry mouth, biliary tract spasm, postural hypotension, syncope, tachy- or bradycardia, urinary retention, and myoclonus occur occasionally. Myoclonus appears to be somewhat dose related and has been described after large doses via IV or intraspinal routes. Myoclonus can be managed by changing to another opioid or with a benzodiazepine or dantrolene. Frequent adverse effects from epidural administration are urinary retention and pruritus; the latter can be managed with naloxone or butorphanol. Possible allergic-type reactions are reported occasionally. Most allergic-type reactions consist of skin rash and wheal and flare over a vein, which can occur with IV injection; these are caused by direct stimulation of histamine release, are not allergic, and are not a sign of a more serious reaction. True allergy is rare. Confusion and disorientation have been linked to phenol and formaldehyde preservatives in epidural infusions, and seizures have been associated with high-dose IV infusions containing sodium bisulfite.

Precautions. Use with caution and in reduced dosage when giving concurrently with other CNS-depressant drugs. Use with caution in pregnancy; the presence of head injury, other intracranial lesions, or pre-existing increase in intracranial pressure; patients having an acute asthmatic attack; COPD or cor pulmonale; decreased respiratory reserve; pre-existing respiratory depression, hypoxia, or hypercapnia; patients whose ability to maintain blood pressure is already compromised; patients with atrial flutter or other supraventricular tachycardias; patients with prostatic hypertrophy or urethral stricture; elderly or debilitated patients; and patients with acute abdominal pain, when administration of the drug might obscure the diagnosis or clinical course. Use with caution in the elderly and neonates and in patients with...
renal dysfunction or elevated bilirubin or LDH levels.\textsuperscript{148,150,151,153} Infants >1 month eliminate morphine efficiently and are unlikely to be unusually sensitive to the respiratory depressant effects but may require longer dosage intervals.\textsuperscript{148} Do not administer IV, IT, or epidurally to opiate-naive patients unless a narcotic antagonist and facilities for assisted or controlled respiration are immediately available.

**Drug Interactions.** Concurrent use of opioids with other CNS depressants (eg, alcohol, antipsychotics, general anesthetics, heterocyclic antidepressants, and sedative-hypnotics) can cause respiratory depression. Cimetidine can increase serum concentration and duration of effect of the opioids.\textsuperscript{125}

**Parameters to Monitor.** Monitor for pain control and signs of respiratory or cardiovascular depression.

**NALOXONE HYDROCHLORIDE** Narcan, Various

**Pharmacology.** Naloxone, an N-allyl derivative of oxymorphone, is a narcotic antagonist that competitively binds at opiate receptors. Naloxone is essentially free of narcotic agonist properties and is used to reverse the effects of narcotic agonists and drugs with partial agonist properties.\textsuperscript{158}

**Administration and Adult Dosage.** IV (preferred) or SC for known or suspected narcotic overdose 0.1–0.2 mg as a first dose, then progressively double the dose q 2–3 min or 0.4 mg diluted in 9 mL saline and injected in 1-mL increments q 30–60 seconds, until respiration and consciousness have become normal or until 10 mg has been given. If response occurs, to prevent recurrent toxicity due to short naloxone half-life, IV infusion at an hourly rate equal to the initial dose required for arousal, with a possible repeat bolus of 50% required 20–30 min after start of infusion.\textsuperscript{159,160} If a total of 10 mg has been given and there is no response, the diagnosis of narcotic overdose should be questioned. The frequency of repeat doses is based on clinical evaluation of the patient. IV for postoperative narcotic depression 0.1–0.2 mg initially, may repeat q 2–3 min until desired level of reversal is reached. Subsequent doses might be needed if the effect of the narcotic outlasts the action of naloxone. (See Notes.) IV for epidural opioid-induced pruritus 0.005–0.01 mg/kg either in incremental doses or as an hourly infusion.\textsuperscript{140} PO for opioid-induced constipation 4–12 mg not more often than q 6 hr; more frequent administration might precipitate withdrawal. Give at a daily dose of approximately 20% of the 24-hr morphine dose. Initial doses should not exceed 5 mg.\textsuperscript{161,162}

**Special Populations.** Pediatric Dosage. IV for known or suspected narcotic overdose 0.01 mg/kg, may repeat as needed. IV for postoperative narcotic depression 0.005–0.01 mg initially, may repeat q 2–3 min until desired level of reversal is reached. IV (preferred) or SC for narcotic depression (neonates) 0.01 mg/kg initially, may repeat q 2–3 min until desired level of reversal is reached.

**Geriatric Dosage.** Same as adult dosage.

**Dosage Forms.** Inj 0.02, 0.4, 1 mg/mL.

**Pharmacokinetics.** **Onset and Duration.** Onset IV within 2–3 min, up to 15 min when given IM or SC; duration variable but usually 1 hr or less.\textsuperscript{163,164}
Fate. From 59% to 67% metabolized by hepatic conjugation and renal elimination of the conjugated compound. V_d is approximately 2–3 L/kg; Cl is about 1.3 L/hr/kg.

t_1/2. 64 ± 12 min in adults; 71 ± 36 min in neonates.

Adverse Reactions. Naloxone administration has been occasionally associated with life-threatening complications such as pulmonary edema, seizures, hypertension, arrhythmias, and violent behavior within 10 min of parenteral administration.

Contraindications. None known.

Precautions. Administration to narcotic-dependent persons (including neonates of dependent mothers) might precipitate acute withdrawal symptoms.

Drug Interactions. None known except for opioid antagonism.

Parameters to Monitor. Respiratory rate, pupil size (might not be useful in mixed-drug or narcotic partial agonist overdoses), heart rate, blood pressure, and symptoms of acute narcotic withdrawal syndrome.

Notes. Naloxone is effective when administered endotracheally to patients with difficult venous access. It is routinely used in the initial treatment of patients with coma of unknown origin. Its use in clonidine overdose has produced mixed results; use in septic and hemorrhagic shock has been disappointing.

OPIOID PARTIAL AGONISTS

Pharmacology. These agents can be classified based on their effects on the opioid receptors. Opioid partial agonists have analgesic effects but are characterized by an analgesic ceiling, such that, beyond a certain point, further increases in dosage do not result in additional analgesia but might produce adverse effects. Tramadol is partly metabolized by CYP2D6, thereby producing an active metabolite (M1) that binds to mu opioid receptors. Patients who are poor metabolizers of debrisoquine and sparteine have negligible M1 production and reduced analgesia, although some pain relief remains because of activation of monoaminergic antinociceptive pathways from tramadol enantiomers.

Administration, Dosage, and Dosage Forms. (See Opioid Analgesics Comparison Chart.)

Patient Instructions. (See Opioids Class Instructions.)

Pharmacokinetics. (See Opioid Analgesics Comparison Chart.)

Adverse Reactions. Sedation, sweating, dizziness, nausea, vomiting, euphoria, dysphoria (agents with delta receptor activity), and hallucinations are most frequent. Occasionally, insomnia, anxiety, anorexia, constipation, dry mouth, syncope, visual blurring, flushing, decreased blood pressure, and tachycardia are reported. After parenteral use, diaphoresis, sting on injection, respiratory depression, transient apnea in the newborn from administration to the mother during labor, shock, urinary retention, and alterations in uterine contractions during labor occur rarely. Other rarely reported effects are muscle tremor and toxic epidermal necrolysis. Local skin reactions and ulceration and fibrous myopathy at the injec-
tion site have been reported with long-term parenteral use of pentazocine.\textsuperscript{86} Tramadol adverse reactions include seizures (some after the first dose) with recommended and excessive dosages. Seizure risk is increased in patients taking concomitant medications that can reduce the seizure threshold (eg, heterocyclic antidepressants, selective serotonin reuptake inhibitors, MAO inhibitors, neuroleptics) and with certain medical conditions (eg, epilepsy, head trauma, metabolic disorders, alcohol and drug withdrawal, or CNS infection). In addition, naloxone administration for tramadol overdose can increase the risk of seizure. Anaphylactoid reactions also have been described in tramadol postmarketing surveillance.\textsuperscript{171-173} Dependence/addiction and major psychological disturbances have been reported with butorphanol nasal spray.\textsuperscript{174}

**Contraindications.** (Tramadol) prior allergy to any opiate; acute intoxication with alcohol, hypnotics, centrally acting analgesics, opioids, or psychotropic drugs. (See Notes.)

**Precautions.** (See Morphine Sulfate.) Also, use cautiously in MI patients because pentazocine and butorphanol increase cardiac workload. All of these agents can produce dependence and withdrawal symptoms after extended use.

**Drug Interactions.** (See Morphine Sulfate.) With the possible exception of tramadol, these agents can precipitate acute withdrawal in narcotic-dependent individuals.\textsuperscript{175}

**Notes.** Because of their ceiling effect, risk of precipitating opiate withdrawal, and marked adverse effects, these agents are not recommended for the management of cancer pain.\textsuperscript{116} Effects of pentazocine are antagonized by naloxone. Naloxone in Talwin NX tablets is not absorbed orally but theoretically prevents parenteral abuse of the oral dosage form; however, IV abuse of Talwin Nxs plus tripelennamine has been reported.\textsuperscript{176}

### OPIOID RECEPTOR SPECIFICITY COMPARISON CHART

<table>
<thead>
<tr>
<th>DRUG</th>
<th>RECEPTOR TYPE</th>
</tr>
</thead>
<tbody>
<tr>
<td>Buprenorphine</td>
<td>Partial agonist-antagonist</td>
</tr>
<tr>
<td>Butorphanol</td>
<td>Partial agonist-antagonist</td>
</tr>
<tr>
<td>Dezocine</td>
<td>Partial agonist-antagonist</td>
</tr>
<tr>
<td>Morphine</td>
<td>Agonist</td>
</tr>
<tr>
<td>Nalbuphine</td>
<td>Antagonist</td>
</tr>
<tr>
<td>Pentazocine</td>
<td>Partial agonist-antagonist</td>
</tr>
<tr>
<td>Tramadol\textsuperscript{a}</td>
<td>Partial or pure agonist\textsuperscript{b}</td>
</tr>
</tbody>
</table>

\textsuperscript{a}Also blocks norepinephrine and serotonin reuptake.

\textsuperscript{b}Not a classic agonist–antagonist; has little or no antagonist properties but appears to have partial mu receptor agonist activity.
**PATIENT-CONTROLLED ANALGESIA (PCA) GUIDELINES CHART**

<table>
<thead>
<tr>
<th>DRUG</th>
<th>IV BOLUS DOSE (MG)</th>
<th>LOCKOUT INTERVAL (MIN)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Buprenorphine</td>
<td>0.03–0.2</td>
<td>10–20</td>
</tr>
<tr>
<td>Fentanyl</td>
<td>0.02–0.1</td>
<td>3–10</td>
</tr>
<tr>
<td>Hydromorphone</td>
<td>0.1–0.5</td>
<td>3–15</td>
</tr>
<tr>
<td>Meperidine</td>
<td>5–30</td>
<td>5–15</td>
</tr>
<tr>
<td>Methadone</td>
<td>0.5–3</td>
<td>10–20</td>
</tr>
<tr>
<td>Morphine</td>
<td>0.5–3</td>
<td>5–20</td>
</tr>
<tr>
<td>Nalbuphine</td>
<td>1–5</td>
<td>5–15</td>
</tr>
<tr>
<td>Oxymorphone</td>
<td>0.2–0.8</td>
<td>5–15</td>
</tr>
<tr>
<td>Pentazocine</td>
<td>5–30</td>
<td>5–15</td>
</tr>
<tr>
<td>Sufentanil</td>
<td>0.003–0.015</td>
<td>3–10</td>
</tr>
</tbody>
</table>

*Some clinicians recommend combining PCA with a basal continuous infusion of the narcotic. The hourly dosage is determined by the patient’s previous narcotic dose requirements and adjusted q 8–24 hr based on the dose of PCA bolus administered, basal continuous infusion, and pain response. A typical starting hourly basal continuous infusion rate for morphine in a 70 kg adult is 0.5–3 mg/hr.*

**INTRASPINAL NARCOTIC ADMINISTRATION GUIDELINES CHART**

<table>
<thead>
<tr>
<th>ROUTE AND DRUG</th>
<th>INTRASPINAL BOLUS DOSE (MG)</th>
<th>ONSET (MIN)</th>
<th>DURATION (HR)</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>EPIDURAL</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Alfentanil</td>
<td>0.7–2<em>b</em></td>
<td>Rapid</td>
<td>1.5–1.7<em>c</em></td>
</tr>
<tr>
<td>Fentanyl</td>
<td>0.025–0.15</td>
<td>5</td>
<td>2–4</td>
</tr>
<tr>
<td>Hydromorphone</td>
<td>1–2</td>
<td>15</td>
<td>10–16</td>
</tr>
<tr>
<td>Methadone</td>
<td>1–10</td>
<td>10</td>
<td>6–10</td>
</tr>
<tr>
<td>Morphine</td>
<td>1–10</td>
<td>30</td>
<td>6–24</td>
</tr>
<tr>
<td>Sufentanil</td>
<td>0.015–0.05</td>
<td>15</td>
<td>4–6</td>
</tr>
</tbody>
</table>

*Use only preservative-free preparations for intraspinal narcotic administration.*

**INTRATHecal (SUBARACHNOID)**

| Morphine       | 0.1–0.5                   | 15          | 8–24          |

*aSome clinicians recommend combining PCA with a basal continuous infusion of the narcotic. The hourly dosage is determined by the patient’s previous narcotic dose requirements and adjusted q 8–24 hr based on the dose of PCA bolus administered, basal continuous infusion, and pain response. A typical starting hourly basal continuous infusion rate for morphine in a 70 kg adult is 0.5–3 mg/hr.*

*bBased on a 70 kg adult body weight (ie, 10–30 µg/kg).*

*cVery short duration of action; requires epidural infusion to obtain prolonged analgesia. Like fentanyl, prolonged epidural infusions produce high systemic concentrations and appear to have little advantage over IV infusion.*

*From references 91 and 100.*
<table>
<thead>
<tr>
<th>DRUG AND SCHEDULE</th>
<th>DOSAGE FORMS</th>
<th>EQUIVALENT PARENTERAL DOSAGE(^a) (MG)</th>
<th>EQUIVALENT ORAL DOSAGE(^a) (MG)</th>
<th>PARENTERAL/ORAL EFFICACY RATIO</th>
<th>DURATION OF ANALGESIA (HR)</th>
<th>PARTIAL ANTAGONIST ACTIVITY</th>
</tr>
</thead>
<tbody>
<tr>
<td>Alfentanil (C-II)</td>
<td>Inj 500 µg/mL</td>
<td>1</td>
<td>—</td>
<td>—</td>
<td>&lt;1</td>
<td>no</td>
</tr>
<tr>
<td>Buprenorphine (C-V)</td>
<td>Inj 0.324 mg/mL</td>
<td>0.3–0.6</td>
<td>—</td>
<td>—</td>
<td>6–8</td>
<td>yes</td>
</tr>
<tr>
<td>Butorphanol</td>
<td>Inj 1, 2 mg/mL</td>
<td>2</td>
<td>—</td>
<td>1/16</td>
<td>3–4</td>
<td>yes</td>
</tr>
<tr>
<td>Codeine (C-II)</td>
<td>Inj 30, 60 mg/mL</td>
<td>120</td>
<td>30</td>
<td>1/2–2/3</td>
<td>4–6</td>
<td>no</td>
</tr>
<tr>
<td>Dezocine (NC)</td>
<td>Inj 5, 10, 15 mg/mL</td>
<td>10–15</td>
<td>—</td>
<td>—</td>
<td>3–4</td>
<td>yes</td>
</tr>
<tr>
<td>Fentanyl (C-II)</td>
<td>Inj 50 µg/mL</td>
<td>0.1</td>
<td>—</td>
<td>1/5</td>
<td>1–2</td>
<td>no</td>
</tr>
<tr>
<td>Hydrocodone and Acetaminophen (C-III)</td>
<td>Tab 5, 7.5, 10 mg with acetaminophen 400 mg, 2.5, 5, 7.5 mg with acetaminophen 500 mg, 7.5 mg with acetaminophen 400, 500, 650, 750 mg, 10 mg with acetaminophen 325, 400, 500, 650, 660 mg</td>
<td>—</td>
<td>5</td>
<td>—</td>
<td>4–6</td>
<td>no</td>
</tr>
</tbody>
</table>

(continued)
<table>
<thead>
<tr>
<th>OPIOID ANALGESICS COMPARISON CHART (continued)</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>DRUG AND SCHEDULE</strong></td>
</tr>
<tr>
<td>Hydromorphone (C-II)</td>
</tr>
<tr>
<td>Dilaudid</td>
</tr>
<tr>
<td>Various</td>
</tr>
<tr>
<td></td>
</tr>
<tr>
<td></td>
</tr>
<tr>
<td>Levorphanol (C-II)</td>
</tr>
<tr>
<td>Levo-Dromoran</td>
</tr>
<tr>
<td>Meperidine (C-II)</td>
</tr>
<tr>
<td>Demerol</td>
</tr>
<tr>
<td>Various</td>
</tr>
<tr>
<td>Methadone (C-II)</td>
</tr>
<tr>
<td>Dolophine</td>
</tr>
<tr>
<td>Various</td>
</tr>
<tr>
<td></td>
</tr>
<tr>
<td></td>
</tr>
<tr>
<td>Morphine (C-II)</td>
</tr>
<tr>
<td>Various</td>
</tr>
<tr>
<td></td>
</tr>
<tr>
<td></td>
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<td></td>
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<tr>
<td></td>
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<td></td>
</tr>
</tbody>
</table>

(continued)
<table>
<thead>
<tr>
<th>DRUG AND SCHEDULE</th>
<th>DOSAGE FORMS</th>
<th>EQUIVALENT PARENTERAL DOSAGE (MG)</th>
<th>EQUIVALENT ORAL DOSAGE (MG)</th>
<th>PARENTERAL/ORAL EFFICACY RATIO</th>
<th>DURATION OF ANALGESIA (HR)</th>
<th>PARTIAL ANTAGONIST ACTIVITY</th>
</tr>
</thead>
<tbody>
<tr>
<td>Nalbuphine (NC)</td>
<td>Inj 10, 20 mg/mL</td>
<td>10</td>
<td>—</td>
<td>1/6</td>
<td>3–6</td>
<td>yes</td>
</tr>
<tr>
<td>Nubain Various</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Oxycodone (C-II)</td>
<td>Cap 5 mg</td>
<td>—</td>
<td>5</td>
<td>—</td>
<td>3–4</td>
<td>no</td>
</tr>
<tr>
<td>Oxycontin</td>
<td>Tab 5 mg</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Roxicodone</td>
<td>Tab 2.5, 5 mg with acetaminophen 325 mg, 5 mg with acetaminophen 500 mg, 7.5 mg with acetaminophen 500 mg, 10 mg with acetaminophen 650 mg.</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Soln 1, 20 mg/mL</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>SR Tab 10, 20, 40, 80 mg.</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Oxymorphone (C-II)</td>
<td>Inj 1, 1.5 mg/mL</td>
<td>1–1.5</td>
<td>—</td>
<td>1/6</td>
<td>4–5</td>
<td>no</td>
</tr>
<tr>
<td>Numorphan</td>
<td>Supp 5 mg.</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Pentazocine (C-IV)</td>
<td>Inj 30 mg/mL</td>
<td>30–60</td>
<td>25</td>
<td>1/3</td>
<td>2–3</td>
<td>yes</td>
</tr>
<tr>
<td>Talwin</td>
<td>Tab 50 mg with naloxone 0.5 mg.</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Talwin Nx</td>
<td>Tab 12.5 mg with aspirin 325 mg, 25 mg with acetaminophen</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Various</td>
<td>650 mg.</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Propoxyphene (C-IV)</td>
<td>Cap (HCl) 65 mg</td>
<td>—</td>
<td>65 (HCl)</td>
<td>—</td>
<td>4–6</td>
<td>no</td>
</tr>
<tr>
<td>Darvon</td>
<td>Tab (HCl) 65 mg with acetaminophen 650 mg.</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Various</td>
<td>Tab (Napsylate) 50, 100 mg</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Tab (Napsylate) 50 mg with acetaminophen 325 mg, 100 mg with acetaminophen 650 mg.</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Susp (Napsylate) 10 mg/mL.</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

(continued)
<table>
<thead>
<tr>
<th>DRUG AND SCHEDULE</th>
<th>DOSAGE FORMS</th>
<th>EQUIVALENT PARENTERAL DOSAGE</th>
<th>EQUIVALENT ORAL DOSAGE</th>
<th>PARENTERAL/ORAL EFFICACY RATIO</th>
<th>DURATION OF ANALGESIA</th>
<th>PARTIAL ANTAGONIST ACTIVITY</th>
</tr>
</thead>
<tbody>
<tr>
<td>Remifentanil (C-II)</td>
<td>Inj 3, 5, 10 mg.</td>
<td>0.1</td>
<td>—</td>
<td>—</td>
<td>&lt; 0.5</td>
<td>no</td>
</tr>
<tr>
<td>Sufentanil (C-II)</td>
<td>Inj 50 µg/mL.</td>
<td>0.01</td>
<td>—</td>
<td>—</td>
<td>2.5–3.5</td>
<td>no</td>
</tr>
<tr>
<td>Tramadol (NC)</td>
<td>Tab 50 mg. Tab 50 mg with acetaminophen (Ultracet)</td>
<td>—</td>
<td>25</td>
<td>—</td>
<td>4–6</td>
<td></td>
</tr>
</tbody>
</table>

---

*aControlled Substance Schedule designated after each drug (in parentheses); NC = not controlled.
*bParenteral dose equivalent to 10 mg morphine.
*cOral dose equivalent to 30 mg codeine. Not for SR products.
*dSubutex and Suboxone (buprenorphine plus naloxone) are used in treating addiction.
*eEquivalent sublingual dose.
*fRecommended dosage is one spray in one nostril, repeated prn in 60–90 min; this cycle may then be repeated q 3–4 hr prn pain.
*gSee Pharmacology and Notes in Methadone monograph.

From references 86, 91, 100, 103, 104, 117, 177–180 and product information.
REFERENCES

52  ANALGESIC AND ANTI-INFLAMMATORY DRUGS


142. Milner AR et al. Intrathecal administration of morphine for elective Caesarean section. A comparison between 0.1 mg and 0.2 mg. *Anaesthesia* 1996;51:871–3.


Antimicrobial Drugs

Aminoglycosides

AMINOGLYCOSIDES

Pharmacology. Aminoglycosides are aminocyclitol derivatives that have concentration-dependent bactericidal activity against Gram-negative aerobic bacteria via binding to the interface between the 30S and 50S ribosomal subunits; anaerobic bacteria are universally resistant because aminoglycoside transport into cells is oxygen dependent. Dibasic cations (eg, magnesium, calcium) and acidic conditions decrease their in vitro action. Streptomycin and kanamycin have poor activity against some Gram-negative bacteria, especially *P. aeruginosa*. Some Gram-positive organisms (eg, streptococci) are relatively resistant to all aminoglycosides; however, in combination with some penicillins or vancomycin, these organisms are often synergistically inhibited or killed. Aminoglycosides have a postantibiotic effect against Gram-negative bacteria, which can be exploited by using less frequent dosage intervals. Resistance is due to transferable plasmid-mediated enzymatic modification or decreased drug uptake.

Administration and Adult Dosage. IM or IV by slow intermittent infusion over 30–60 min, although 15-min infusions are safe. Newer dosage regimens combine the usual daily dosage into a single IV infusion administered over 60 min. This method takes advantage of the concentration-related bactericidal effects and postantibiotic effect of aminoglycosides and may result in less toxicity. IT or intraventricular administration is usually necessary to achieve therapeutic CSF levels. Special Populations. Pediatric Dosage. Same as adult dosage, but adjust for age-related reduction in renal function. Geriatric Dosage. Same as adult dosage, but adjust for age-related reduction in renal function. Other Conditions. Use of IBW for determining the mg/kg dosage appears to be more accurate than dosage based on TBW. In morbid obesity, dosage requirement may best be estimated using a dosing weight of IBW + 0.4 (TBW – IBW). With conventional dosage methods, serum drug levels should be in the range of 3–10 mg/L; high peaks (>6 mg/L with gentamicin and tobramycin) may be associated with better outcome in bacteremia, pneumonia, and other systemic infections. Critically ill patients with serious infections or in disease states known to markedly alter aminoglycoside pharmacokinetics (eg, cystic fibrosis, burns, or major surgery) often have variable distribution and excretion of the drugs. When the drug is administered once daily, higher peak concentrations (>10–20 mg/L with gentamicin and tobramycin) are targeted based on the patient’s disease state and pharmacokinetic parameters. (See Aminoglycosides Comparison Chart.)
Chart.) Adjust dosage based on renal function. Individualization is critical because these agents have a low therapeutic index. In renal impairment, the following guidelines may be used to determine initial dosage (modified from reference 6):

1. Select loading dose in mg/kg (LBW or dosing weight as above) to provide peak serum levels in the range listed below for the desired aminoglycoside.

<table>
<thead>
<tr>
<th>AMINOGLYCOSIDE</th>
<th>USUAL LOADING DOSE</th>
<th>EXPECTED PEAK SERUM LEVEL</th>
</tr>
</thead>
<tbody>
<tr>
<td>Tobramycin</td>
<td>1–2 mg/kg</td>
<td>3–10 mg/L</td>
</tr>
<tr>
<td>Gentamicin</td>
<td>1–2 mg/kg</td>
<td>3–10 mg/L</td>
</tr>
<tr>
<td>Amikacin</td>
<td>5–7.5 mg/kg</td>
<td>15–30 mg/L</td>
</tr>
</tbody>
</table>

2. Select maintenance dose (as percentage of chosen loading dose) to continue peak serum levels indicated above, according to desired dosage interval and the patient’s corrected Clcr.

<table>
<thead>
<tr>
<th>Clcr (ML/Min)</th>
<th>Half-Lifea (HR)</th>
<th>8 HR</th>
<th>12 HR</th>
<th>24 HR</th>
</tr>
</thead>
<tbody>
<tr>
<td>90</td>
<td>3.1</td>
<td>84%</td>
<td>—</td>
<td>—</td>
</tr>
<tr>
<td>80</td>
<td>3.4</td>
<td>80</td>
<td>91%</td>
<td>—</td>
</tr>
<tr>
<td>70</td>
<td>3.9</td>
<td>76</td>
<td>88</td>
<td>—</td>
</tr>
<tr>
<td>60</td>
<td>4.5</td>
<td>71</td>
<td>84</td>
<td>—</td>
</tr>
<tr>
<td>50</td>
<td>5.3</td>
<td>65</td>
<td>79</td>
<td>—</td>
</tr>
<tr>
<td>40</td>
<td>6.5</td>
<td>57</td>
<td>72</td>
<td>92%</td>
</tr>
<tr>
<td>30</td>
<td>8.4</td>
<td>48</td>
<td>63</td>
<td>86</td>
</tr>
<tr>
<td>25</td>
<td>9.9</td>
<td>43</td>
<td>57</td>
<td>81</td>
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<tr>
<td>20</td>
<td>11.9</td>
<td>37</td>
<td>50</td>
<td>75</td>
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<td>17</td>
<td>13.6</td>
<td>33</td>
<td>46</td>
<td>70</td>
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<td>15</td>
<td>15.1</td>
<td>31</td>
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<td>67</td>
</tr>
<tr>
<td>12</td>
<td>17.9</td>
<td>27</td>
<td>37</td>
<td>61</td>
</tr>
<tr>
<td>10b</td>
<td>20.4</td>
<td>24</td>
<td>34</td>
<td>56</td>
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<td>7b</td>
<td>25.9</td>
<td>19</td>
<td>28</td>
<td>47</td>
</tr>
<tr>
<td>5b</td>
<td>31.5</td>
<td>16</td>
<td>23</td>
<td>41</td>
</tr>
<tr>
<td>2b</td>
<td>46.8</td>
<td>11</td>
<td>16</td>
<td>30</td>
</tr>
<tr>
<td>0b</td>
<td>69.3</td>
<td>8</td>
<td>11</td>
<td>21</td>
</tr>
</tbody>
</table>

aAlternatively, 50% of the chosen loading dose can be given at an interval approximately equal to the estimated half-life.

bUse measured serum levels to adjust dosage for patients with Clcr <10 mL/min. Give supplemental doses of 50–75% of the loading dose after each hemodialysis period.
These guidelines are based on population data; serum levels in individual patients might deviate from guideline estimates. No guidelines have been developed for netilmicin or streptomycin.

**Dosage Forms.** (See Aminoglycosides Comparison Chart.)

**Patient Instructions.** Report any dizziness or sensations of ringing or fullness in the ears.

**Pharmacokinetics. Serum Levels.** (See Parameters to Monitor and Aminoglycosides Comparison Chart.)

**Fate.** Absorption after oral or rectal administration is about 0.2–2%; absorption across denuded skin can reach 5%. Irrigation of vascularized areas (eg, peritoneal cavity) results in absorption approximating IM use. IM administration is followed by rapid and complete absorption, with peak serum levels occurring after 0.5–1.5 hr. IV infusions over 0.5–1 hr produce serum levels similar to equal IM doses. Binding of aminoglycosides to plasma proteins is low. These agents distribute rapidly into the extracellular fluid compartment with a $V_d$ of about 0.3 ± 0.08 L/kg, which is increased by fever, edema, ascites, and fluid overload, and in neonates. Aminoglycosides accumulate markedly in some tissues, especially the renal cortex, to levels many times those found in the serum, particularly with frequent dosage intervals compared with the same dosage given at less frequent intervals. Levels in the CSF of patients with meningitis generally do not exceed 25% of serum levels, except in neonates; penetration into the eye is inadequate for treatment of intraocular infections. Penetration into lung tissues and sputum is low, and large doses might be necessary to optimally treat pneumonia with relatively insensitive organisms (eg, *P. aeruginosa*). Distribution of aminoglycosides into the peritoneal cavity of patients with peritonitis is therapeutically adequate. Elimination is via glomerular filtration of unchanged drug; $Cl$ is about 90% of $Cl_{cr}$. After discontinuation, low levels of aminoglycoside can be detected in the urine for several days caused by excretion of drug that had accumulated in deep tissue compartments.

$\frac{1}{2}$. Phase 5–15 min; $\beta$ phase (adults) about 2 ± 0.4 hr with normal renal function (1.5–9 hr in neonates <1 week and 3 hr in older infants); can be more variable in certain groups (eg, obstetric and burn patients) despite normal renal function; 50–70 hr in anuria. A prolonged $\gamma$ elimination phase is observed when concentrations fall to the lower range of detectability, representing egress from deep tissue compartments and subsequent renal elimination; the half-life of this phase is 60–350 hr (usually 150–200). $\beta$ Phase half-life is most important for use in calculating individualized dosage, but the $\gamma$ phase may account for the gradual rise of serum levels and apparent increase in half-life with continued therapy, despite stable renal function.

**Adverse Reactions.** Aminoglycoside-induced nephrotoxicity is usually mild and reversible; progression to severe renal disease and dependence on dialysis is rare. Nephrotoxicity is manifested by elevations in $Cr$, BUN, and aminoglycoside concentrations and appearance of renal tubular casts, enzymes, and $\beta_2$-microglobulin and occurs in 5–30% of patients, depending on the criteria used and the population risk factors present. Duration of therapy, prior aminoglycoside therapy,
advanced age, pre-existing renal disease, liver disease, volume depletion, and female sex have been identified as risk factors for nephrotoxicity. Concomitant use of nephrotoxic drugs also increases the risk of nephrotoxicity. Elevated trough levels are not a risk factor but often a result of nephrotoxicity. There is no evidence that there are clinically important differences in nephrotoxicity between gentamicin, tobramycin, netilmicin, and amikacin. Depletion of magnesium and other minerals caused by increased renal excretion occurs. Occasional, but often permanent, vestibular toxicity is reported, usually in association with streptomycin. Subclinical vestibular disturbances can be detected in 40% or more of patients receiving aminoglycosides. Early cochlear damage can be detected only by sequential audiometric examination because hearing loss in conversational frequencies is a sign of advanced auditory impairment. Furthermore, early auditory damage is not as apparent in the elderly or others with pre-existing high-tone deficits. Risk factors for ototoxicity are duration of therapy, bacteremia, hypovolemia, peak temperature, and liver disease. Elevated serum concentrations apparently are not associated with increased ototoxicity risk, and there are no apparent clinically important differences between gentamicin, tobramycin, netilmicin, and amikacin. Oral aminoglycosides, primarily neomycin, have been associated with a sprue-like malabsorption syndrome. Neuromuscular blockade with respiratory failure is rare, except in predisposed patients. 

**Precautions.** Pregnancy; pre-existing renal impairment; vestibular or cochlear impairment; myasthenia gravis; hypocalcemia; postoperative or other conditions that depress neuromuscular transmission.

**Drug Interactions.** Concurrent or sequential use of other nephro- or ototoxic agents can increase the risk of aminoglycoside toxicities. Concurrent use of aminoglycosides with neuromuscular blocking agents can potentiate neuromuscular blockade and cause respiratory paralysis. The action of oral anticoagulants can be potentiated by oral neomycin, presumably via reduced absorption or synthesis of vitamin K. Ticarcillin and acylampicillins can degrade aminoglycosides in vitro, resulting in artificially low levels; the extent of degradation is dependent on time, temperature, and β-lactam concentration. Degradation can occur in vivo in patients with renal insufficiency. Amikacin is the aminoglycoside least susceptible to β-lactam inactivation.

**Parameters to Monitor.** Renal function tests before and q 2–3 days during therapy. Audiometry and electronystagmography may be performed in patients able to cooperate. Monitor aminoglycoside serum concentrations carefully, especially in the elderly, those with renal impairment, hemodynamically unstable patients, and those requiring high peak serum concentrations or prolonged (>10 days) therapy. In adults receiving conventional therapy, monitor serum levels after steady state is achieved. With once-daily therapy targeting high peaks and undetectable troughs, obtain levels after the first dose. Obtain follow-up levels if renal function changes. In neonates or other patients with rapidly changing renal function, obtain serum drug concentrations initially and q 2–3 days until stable. However, with once- or twice-daily dosage and in pediatric patients, trough serum levels are often undetectable and other sampling strategies are necessary. (See also Special Populations, Other Conditions.)
Notes. Of the available aminoglycosides, gentamicin, tobramycin, netilmicin, and amikacin are the most clinically useful. Streptomycin use is largely restricted to the treatment of enterococcal endocarditis (in combination with ampicillin), tuberculosis, brucellosis, plague, and tularemia; it is currently available only for compassionate use from the manufacturer. Amikacin is often used as part of a combination regimen for treatment of Mycobacterium avium complex infection. Neomycin is much more toxic than the other aminoglycosides when given parenterally; it is restricted to oral use for gut sterilization and topical use for minor infections. Resistance among Gram-negative organisms, especially *P. aeruginosa*, has virtually eliminated the systemic use of kanamycin. Tobramycin is roughly equivalent to gentamicin therapeutically, although it is about 2–4 times more active against *P. aeruginosa* than is gentamicin, is often active against gentamicin-resistant *P. aeruginosa*, and might be preferred because of a superior peak-to-MIC ratio. Resistance of Gram-negative bacilli is lowest with amikacin; amikacin use does not appear to result in increased resistance to the drug.
## Aminoglycosides Comparison Chart

<table>
<thead>
<tr>
<th>Drug</th>
<th>Dosage Forms</th>
<th>Adult Dosagea</th>
<th>Pediatric Dosagea</th>
<th>Usual Therapeutic Serum Levels (MG/L)b</th>
</tr>
</thead>
<tbody>
<tr>
<td>Amikacin Sulfate</td>
<td>Inj 50, 250 mg/mL.</td>
<td>IM or IV 15–20 mg/kg/day in 2 equally divided doses; IT 5–20 mg/day.</td>
<td>IM or IV (&lt;1 week) 12–15 mg/kg q 36–48 hr; IM or IV (infants &gt;1 week) 12 mg/kg q 24 hr; IM or IV (children) same as adult mg/kg dosage.</td>
<td>Peakc 20–35  ≤10</td>
</tr>
<tr>
<td>Amikin</td>
<td></td>
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<td></td>
<td></td>
</tr>
<tr>
<td>Gentamicin Sulfate</td>
<td>Inj 10, 40 mg/mL.</td>
<td>IM or IV 5–6 mg/kg/day in equally divided doses q 8–12 hr or in a single-dose IV q 24 hr.</td>
<td>IM or IV (&lt;1 week) 4–5 mg/kg q 36–48 hr; IM or IV (infants &gt;1 week) 4 mg/kg q 24 hr; IM or IV (children) 6–7.5 mg/kg/day (7–10 mg/kg/day in cystic fibrosis) in 3–4 equally divided doses q 6–8 hr; IT 1–2 mg q 24 hr.</td>
<td>Peakc 6–12  ≤2</td>
</tr>
<tr>
<td>Garamycin</td>
<td>IT Inj 2 mg/mL.</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Various</td>
<td>Ophth Oint 3 mg/g</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Ophth Soln 3 mg/mL.</td>
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<tr>
<td></td>
<td>Top Crm 0.1%</td>
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</tr>
<tr>
<td></td>
<td>Top Oint 0.1%</td>
<td></td>
<td></td>
<td></td>
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<tr>
<td>Netilmicin Sulfate</td>
<td>Inj 100 mg/mL.</td>
<td>IM or IV 3–6.5 mg/kg/day in 1–3 equally divided doses q 8–24 hr.</td>
<td>Same as gentamicin.</td>
<td>Peakc 6–12  ≤2</td>
</tr>
<tr>
<td>Netromycin</td>
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</tbody>
</table>

(continued)
<table>
<thead>
<tr>
<th>DRUG</th>
<th>DOSAGE FORMS</th>
<th>ADULT DOSAGE^a</th>
<th>PEDIATRIC DOSAGE^a</th>
<th>USUAL THERAPEUTIC SERUM LEVELS (MG/L)^b</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>Peak^c</td>
<td>Trough</td>
<td>Peak^c</td>
</tr>
<tr>
<td><strong>Streptomycin Sulfate</strong></td>
<td>Inj 400 mg/mL.</td>
<td>IM 15–25 mg/kg/day (usually 1–2 g/day) in 2 equally divided doses q 12 hr; IM for TB 12–15 mg/kg/day to a maximum of 1 g or 25–30 mg/kg to a maximum of 1.5 g 2–3 times/week.</td>
<td>IM (neonates) 20–30 mg/kg/day in 2 equally divided doses q 12 hr; IM (children) 20–40 mg/kg/day in 2 equally divided doses q 12 hr; IM for TB 20–40 mg/kg/day or 25–30 mg/kg 2–3 times/week.</td>
<td>15–30</td>
</tr>
<tr>
<td>Various</td>
<td></td>
<td></td>
<td></td>
<td></td>
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</tbody>
</table>

| **Tobramycin Sulfate** | Inj 10, 40 mg/mL. | IM or IV same as gentamicin; IV for cystic fibrosis 10 mg/kg/day; Inhal for cystic fibrosis 300 mg q 12 hr for 28 days; IT 4–8 mg q 24 hr. | IM or IV same as gentamicin; IV for cystic fibrosis 10 mg/kg/day; Inhal for cystic fibrosis 300 mg q 12 hr for 28 days. | 6–12 | ≤2 |
| Nebcin                 | Inj 1.2 g            |         |        |         |        |
| TOBI                   | Ophth Oint 3 mg/g    |         |        |         |        |
| Various                | Ophth Soln 3 mg/mL.  |         |        |         |        |
|                        | Nebulizer Soln 60 mg/mL. |         |        |         |        |

^aFor systemic infections; UTIs are adequately treated with lower dosages.
^bBased on divided doses given q 8–12 hr; higher peaks and lower (or undetectable) troughs are seen when less frequent dosage intervals are used.
^cAs seen 30 min after a 30-min IV infusion or approximately 1 hr after IM administration of a usual adult dose. Uncomplicated UTIs can be treated with smaller doses that produce much lower serum levels; however, serious infections, such as Gram-negative bacteremia, pneumonia, or endocarditis might require doses resulting in serum levels in the higher part of the range. Clinical efficacy appears to increase as the ratio of the peak serum level to the MIC of the pathogen increases.15
^dThese doses conform to those used in published clinical trials, but higher dosages might be necessary in certain patient populations.
Antifungal Drugs

**AMPHOTERICIN B**
- Fungizone

**AMPHOTERICIN B CHOLESTERYL SULFATE**
- Amphotec

**AMPHOTERICIN B LIPID COMPLEX**
- Abelcet

**LIPOSOMAL AMPHOTERICIN B**
- AmBisome

**Pharmacology.** Amphotericin B is a polyene macrolide antifungal drug isolated from the bacteria *Streptomyces nodosus*. Drug binding to ergosterol constituents within the cytoplasmic membrane of fungi, with subsequent disruption of membrane integrity and function, is the pharmacologic mechanism of action for amphotericin B. Innate or acquired resistance to amphotericin B is rare. Sensitivity of fungi to amphotericin B is related to the concentration of ergosterol present in the cytoplasmic membrane.16,17

**Administration and Adult Dosage. Intravenous** (See Amphotericin B Formulations Comparison Chart.) A test dose may be given before the first amphotericin B dose. The greatest utility of a test dose is identification of patients particularly sensitive to infusion-related adverse effects of amphotericin B, or identification of patients with hypersensitivity to an alternative amphotericin B formulation. Conventional amphotericin B 1 mg in D5W 20 mL, or an adequate admixture volume to deliver 2–5% of the initial dose of any amphotericin B formulation, infused over 10–20 min without premedication can be used as a test dose. Monitor patients closely for 30–60 min after the test dose.16 The manufacturer of amphotericin B lipid complex recommends against a test dose. Initiate therapy with the full treatment dose for patients with life-threatening fungal disease. Some advocate initiation of amphotericin B at a fraction of the therapeutic dose with daily incremental increases to achieve the desired therapeutic dosage. Although it has not been evaluated in a controlled manner, the intent of this approach is improvement of patient tolerance to infusion-related adverse effects.16,18

**Maintenance therapy** conventional amphotericin B and amphotericin B lipid complex can be given every other day or Monday, Wednesday, and Friday.16,19

**IV for prophylaxis after bone marrow transplantation** (conventional amphotericin B) 0.1 mg/kg or 5–10 mg daily has been used.20 **Infusion time** the frequency and severity of infusion-related adverse effects is similar with administration of amphotericin B over 1–2 hr and 4–6 hr. To prevent drug-induced hyperkalemia, amphotericin B must be infused over 4–6 hr in patients with renal failure, pre-existing hyperkalemia, or markedly reduced potassium clearance.18 **Duration of therapy** with amphotericin B is not well defined. Patients with life-threatening mycotic disease must receive amphotericin B until resolution of clinical and microbiologic evidence of fungal infection, or until unacceptable drug-induced toxicity occurs. Cumulative total dosage of amphotericin B is generally 10–20 mg/kg.16,18 **PO for oral candidiasis** (amphotericin B suspension) 1 mL qid swished and held in mouth for 1 min, or as long as possible, then swallow. Continue therapy for at least 2 weeks. **Top** apply to affected area 2–4 times daily for 1–4 weeks. **IM or PO administration** is not recommended for injectable amphotericin B.
Alternative routes of administration of extemporaneously prepared amphotericin B for injection are infrequently used to facilitate drug availability to a sanctuary site or minimize systemic toxicity. Use of alternative routes of amphotericin B administration is based primarily on case reports, and the safety and efficacy of extemporaneously prepared amphotericin B administered by alternative routes have not been evaluated in a controlled manner. Subsequently, administration of amphotericin B by an alternative route should not replace standard therapy.

Intra-articular for fungal arthritis 5–50 mg q 2–7 days. The dose of intra-articular amphotericin B is determined by the size of the infected joint.21 Intracavitary for pulmonary aspergillomas 5–50 mg in D5W daily or 2–3 times weekly has been used in patients unable to undergo surgical resection.22 Inhalation for prophylaxis against Aspergillus sp. after bone marrow transplantation 0.15% in D5W nebulized to deliver 10 mg/day in 2 divided doses.23 Intranasal for prophylaxis in bone marrow transplant recipients amphotericin B 0.5% in sterile water 10 mg/day in divided doses.24 Intraperitoneal for the treatment of fungal peritonitis has been used in patients receiving peritoneal dialysis.16 Instillation is problematic because amphotericin B is physically incompatible with ionic solutions such as dialysate. Intrathecal administration of conventional amphotericin B 0.5–1 mg 2–3 times/week or 0.3 mg/day has been reported. The intrathecal dosage of conventional amphotericin B is generally started at 0.025–0.05 mg/dose, with subsequent doses increased at 0.025–0.05 mg/day increments to the desired therapeutic or maximum tolerated dosage. CNS administration is generally via an Ommaya reservoir. Although an Ommaya reservoir is not mandatory for intrathecal administration of amphotericin B, the device facilitates repeated drug administration with more precise drug delivery, improved patient tolerance, and clarified CSF diagnostic quality. Amphotericin B administration by lumbar puncture and intracisternal injection has been reported.16 Bladder irrigation for the treatment of uncomplicated fungal cystitis infuse 50 mg/L in sterile water over 24 hr.16 Topical ocular for the treatment of keratomycosis amphotericin B 0.15% (0.1–0.25%) in preservative-free sterile water has been given concurrently with atropine ophthalmic drops q 30–60 min for the initial 48–72 hr of treatment; subsequent to subjective improvement and ocular re-epithelization, the dosage interval may be changed to qid for at least 1 month.16,25 Intravitreal for fungal keratomycosis 5 μg/0.1 mL preservative-free sterile water has been used.26 Subtenonian injection for the treatment of postoperative fungal endophthalmitis 500–750 μg/day for 8 doses has been used.16

Special Populations. Pediatric Dosage. IV. Same as adult dosage for conventional amphotericin B, amphotericin B colloidal dispersion, amphotericin B lipid complex, and liposomal amphotericin. IV for prophylaxis after bone marrow or solid organ transplantation (liposomal amphotericin B) 1 mg/kg/day has been used.27 PO same as adult dosage. Top same as adult dosage for cream, lotion, and ointment.

Geriatric Dosage. Same as adult dosage for conventional amphotericin B, amphotericin B colloidal dispersion, amphotericin B lipid complex, and liposomal amphotericin. Long-term IV administration is more likely to be limited by renal impairment. Comorbid conditions might reduce patient tolerance to ancillary
medications used for management of infusion-related adverse effects (eg, corticosteroid-induced sodium retention).

**Other Conditions.** (All products) With pre-existing chronic renal dysfunction, no dosage adjustment is necessary, but the duration of the infusion must be 4–6 hr to prevent drug-related hyperkalemia. In acute renal dysfunction, interrupt treatment or extend dosage interval or decrease dosage to reduce exacerbation of renal impairment, as patient’s clinical condition allows.18 For patients ≥1.3 times IBW, calculate dose based on IBW or dosing weight of IBW + 0.4 x (TBW − IBW).28

**Dosage Forms.** **Inj** (see Amphotericin B Formulations Comparison Chart.) **Oral Susp** 100 mg/mL; **Top Crm** 30 mg/g; **Top Lot** 30 mg/mL; **Top Oint** 30 mg/g.

**Patient Instructions.** (Injection). Infusion reactions such as shaking, chills, fever, nausea, and other symptoms can occur when this medication is being given. Although uncomfortable, these effects are transient. Certain medications reduce infusion reactions for most people. Amphotericin B might affect your kidneys. If this occurs, you may need to take mineral supplements by mouth. (Oral Suspension). Shake container well before use. Swish and hold the product in your mouth for one minute, or as long as possible, and then swallow. Discontinue if mouth irritation occurs. (Topical). This preparation can stain clothing.

**Missed Doses.** Take a missed oral or topical dose as soon as it is remembered. If it is time for the next dose, do not double the dose.

**Pharmacokinetics.** Preclinical and phase 1 testing of conventional amphotericin B preceded development of high-performance liquid chromatography and refinement of pharmacokinetic methodology. Pharmacokinetic parameters quoted in tertiary literature might actually reflect drug concentration analysis using microbiologic assays.

**Serum Levels.** A correlation between serum levels and therapeutic or toxic drug effects has not been identified or defined for any commercially available amphotericin B formulation.

**Fate.** (Conventional amphotericin B) poor oral and IM absorption. End of infusion serum concentration was 0.984 ± 0.056 mg/L after 0.25 mg/kg to 8 normal healthy volunteers.16 V_dav is 0.74 ± 0.13 L/kg.29 Extensively bound (>90%) to plasma lipoproteins.16 Accumulates in hepatic, splenic, pulmonary, and renal tissue.28 V_dav of 4 ± 0.3 L/kg is derived from bioanalysis of serum from 2 patients completing chronic therapy with amphotericin B.16 Cl is 0.01 ± 0.001 L/hr/kg.29 Metabolites of amphotericin B have not been identified.16 Urinary elimination is 3–8%.16,20 (Amphotericin B cholesteryl sulfate) V_dav is 4.2 ± 1.4 L/kg in adults and 4.6 ± 1.7 L/kg in children; Cl is 0.11 ± 0.03 L/hr/kg in adults and 0.14 ± 0.02 L/hr/kg in children; AUC is 9.6 ± 2.6 mg/L/hr in adults and 7.1 ± 2.6 mg/L/hr in children.30 (Amphotericin B lipid complex) V_dav is 3.9 ± 0.3 L/kg; Cl is 0.08 ± 0.02 L/hr/kg; AUC is 2.76 ± 0.25 mg/L/hr.29 (Liposomal amphotericin B) V_dav is 0.37 L/kg; Cl is 0.023 L/hr/kg; AUC is 423 mg/L/hr.31

t½. (Conventional amphotericin B) ß phase 24–50 hr; γ phase 15 days;16,29 (amphotericin B cholesteryl sulfate) 32 ± 5.6 hr in adults and 32 ± 13 hr in children;30 (amphotericin B lipid complex) ß phase 45 ± 6.3 hr;29 (liposomal amphotericin B) α phase 1.74 hr; β phase 23.6 hr.31
**Adverse Reactions.** Frequent adverse effects include infusion-related reactions, nephrotoxicity, normochromic normocytic anemia and phlebitis. Infusion reactions ordinarily include rigors, chills, and fever. Less common infusion-related reactions include nausea, tachycardia, tachypnea, hypotension, hypertension, bradycardia, myalgia, and arthralgia. Symptoms generally occur during or within 60–90 min after completion of the infusion. Symptoms decrease with ancillary medications and repeated administration. **Meperidine** 25–50 mg IV reduces the duration and intensity of rigors and chilling. **Acetaminophen** 325–650 mg PO reduces hyperpyrexia, and is often administered as premedication. **Diphenhydramine** 25–50 mg PO or IV is often included as a premedication. **Hydrocortisone**, which reduces fever, chills, and nausea, is reserved for patients with infusion reactions refractory to other ancillary medications. Case reports describe the use of **dantrolene** for refractory rigors and chills. Although premedication with **ibuprofen** reduces the rigors and chills, most patients receiving amphotericin B are at risk for adverse effects from the nephrotoxic and antiplatelet effects of NSAIDs. The prevalence of infusion reactions is greater with conventional amphotericin B or amphotericin B cholesteryl sulfate than with amphotericin B lipid complex or liposomal amphotericin. Rapid infusion (≤60 min) of amphotericin B can cause hyperkalemia and cardiovascular collapse in anephric or hyperkalemic patients. Amphotericin B cholesteryl sulfate, amphotericin B lipid complex, and liposomal amphotericin are each less nephrotoxic than conventional amphotericin B. However, the lipid-based formulations are not devoid of nephrotoxicity. Nephrotoxicity is generally reversible. Permanent renal impairment can occur, particularly in patients receiving conventional amphotericin B at doses over 1 mg/kg/day or have pre-existing renal impairment, prolonged therapy, sodium depletion, or concurrent nephrotoxic drugs. Signs of nephrotoxicity are increased BUN and Cr, hypomagnesemia, hypokalemia, and renal tubular acidosis. Nephrotoxicity can be reduced with infusion of 0.9% NaCl 250–1000 mL over 30–45 min immediately before amphotericin B. The saline infusion may be repeated immediately after amphotericin B administration. The patient’s body size and cardiovascular status must be considered when selecting the volume and rate of 0.9% NaCl infusion. Normochromic normocytic anemia, which is secondary to amphotericin B–induced nephrotoxicity, is mild and transient and rarely requires intervention. Phlebitis is secondary to chronic peripheral administration of conventional amphotericin B. Some advocate adding heparin 1 IU/mL to minimize phlebitis.

Rare adverse effects reported with amphotericin B are anorexia, emesis, diarrhea, cramping epigastric pain, premature ventricular contraction, bradycardia, dilated cardiomyopathy, hypertension, diffuse alveolar hemorrhage, rhabdomyolysis, and parkinsonian syndrome. Intrathecal administration of amphotericin B causes headache, nausea, vomiting, abdominal pain, urinary retention, tinnitus, visual changes, ventriculitis, paresthesias, numbness, monos- or paraparesis, arachnoiditis, focal neurologic defects, and chemical or bacterial meningitis. Life-threatening brain puncture and hemorrhage can occur with intracisternal injection.

**Precautions.** Pregnancy. Impaired renal function. Avoid rapid infusions (<4 hr) in patients with Clr <20 mL/min, hyperkalemia, or reduced ability to excrete
potassium. Separate from neutrophil infusions by at least 6 hr. Complete infusion at least 2 hr before platelet transfusions.

**Drug Interactions.** Additive nephrotoxicity can occur with cyclosporine, tacrolimus, aminoglycosides, loop diuretics, or other nephrotoxic agents. Corticosteroids can enhance potassium loss.

**Parameters to Monitor.** Monitor infusion-related adverse effects with first 3 doses, then as indicated by severity of reactions. Monitor serum Cr, BUN, magnesium, potassium before therapy, and at least twice weekly during therapy. Monitor patients at great risk for renal dysfunction daily. Monitor Hb at least weekly. Monitor microbiologic, radiographic, and clinical signs of fungal infection. Ancillary use of hydrocortisone, acetaminophen, or aspirin might mask fevers.

**Notes.** To ensure even lipid complex distribution, invert admixtures of amphotericin B lipid complex several times immediately before starting the infusion and q 2 hr thereafter. Because amphotericin B has a propensity to precipitate, avoid admixture or Y-site administration of all amphotericin B formulations with IV fluids (except dextrose solution), other intravenous drugs, or blood products. Avoid admixture of conventional amphotericin B with lipid emulsion. Physical incompatibility of this admixture evolves >10 μ particles and phase separation. Acronyms for the various amphotericin B formulations are as follows: conventional amphotericin B, DAmB; amphotericin B cholesteryl sulfate, ABCD; amphotericin B lipid complex, ABLC; liposomal amphotericin B, L-AmB. Amphotericin B cholesteryl sulfate is also known as amphotericin B colloidal dispersion and Amphocil.
### AMPHOTERICIN B PRODUCTS COMPARISON CHART

<table>
<thead>
<tr>
<th></th>
<th>CONVENTIONAL AMPHOTERICIN B</th>
<th>AMPHOTERICIN B CHOLESTERYL SULFATE</th>
<th>AMPHOTERICIN B LIPID COMPLEX</th>
<th>LIPOSOMAL AMPHOTERICIN B</th>
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<tbody>
<tr>
<td><strong>LIPID CHEMISTRY</strong></td>
<td></td>
<td></td>
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<tr>
<td>Lipid component</td>
<td>Deoxycholate</td>
<td>Cholesteryl Sulfate</td>
<td>DMPG, DMPC</td>
<td>HSPC, DSPC</td>
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<tr>
<td>Diameter (nm)</td>
<td>50</td>
<td>120–140</td>
<td>1600–11,000</td>
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<tr>
<td>Configuration</td>
<td>Micelle</td>
<td>Discoid</td>
<td>Ribbon-like</td>
<td>Spherical liposome</td>
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<tr>
<td><strong>PHARMACEUTICAL CHARACTERISTICS</strong></td>
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<tr>
<td>Vial size (mg)</td>
<td>50</td>
<td>50, 100</td>
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<tr>
<td>Storage conditions</td>
<td>2–8°C</td>
<td>15–30°C</td>
<td>2–8°C</td>
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<tr>
<td><strong>ADMINISTRATION AND DOSAGE</strong></td>
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<tr>
<td>Daily dosage (mg/kg)</td>
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<tr>
<td>Sensitive fungi</td>
<td>0.5–1</td>
<td>3–4</td>
<td>2.5–5</td>
<td>1–3</td>
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<tr>
<td>Less-sensitive fungi</td>
<td>1–1.5</td>
<td>6</td>
<td>5</td>
<td>3–5</td>
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<tr>
<td>Infusion duration (hr)</td>
<td>1–6 (≤50 mg/hr)</td>
<td>2–4</td>
<td>2</td>
<td>1–2</td>
</tr>
<tr>
<td>In-line filter</td>
<td>Not recommended.</td>
<td>Do not filter.</td>
<td>Do not filter.</td>
<td>May use if pore size ≥1 µ.</td>
</tr>
<tr>
<td>Compatible IV fluids</td>
<td>D5W</td>
<td>D5W</td>
<td>D5W</td>
<td>D5W</td>
</tr>
<tr>
<td>Admixture concentration (mg/mL)</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Admixture expiration</td>
<td>0.5–0.25</td>
<td>0.16–0.83</td>
<td>1–2</td>
<td>1–2</td>
</tr>
<tr>
<td></td>
<td>Determined by lack of preservative</td>
<td>24 hr at 2–8°C</td>
<td>48 hr at 2–8°C, then an additional 6 hr at room temperature</td>
<td>6 hr at 2–8°C or at room temperature</td>
</tr>
</tbody>
</table>

(continued)
### AMphotericin B Products Comparison Chart (continued)

<table>
<thead>
<tr>
<th>PHARMACOKINETICS</th>
<th>CONVENTIONAL AMPHOTERICIN B</th>
<th>AMPHOTERICIN B CHOLESTeryl SULFATE</th>
<th>AMPHOTERICIN B LIPID COMPLEX</th>
<th>LIPOSOMAL AMPHOTERICIN B</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Vdss (L/kg)</strong></td>
<td>Fungizone: 0.74 ± 0.13 L/kg</td>
<td>Amphotec: (Adult) 4.2 ± 1.4</td>
<td>3.9 ± 0.3</td>
<td>AmBisome: 0.37</td>
</tr>
<tr>
<td></td>
<td>(Child) 4.6 ± 1.7</td>
<td>(Adult) 4.2 ± 1.4</td>
<td>3.9 ± 0.3</td>
<td></td>
</tr>
<tr>
<td><strong>Clearance (L/hr/kg)</strong></td>
<td>0.01 ± 0.001</td>
<td>(Adult) 0.11 ± 0.03</td>
<td>0.08 ± 0.02</td>
<td>0.023</td>
</tr>
<tr>
<td></td>
<td>(Child) 0.14 ± 0.02</td>
<td>(Adult) 0.11 ± 0.03</td>
<td>0.08 ± 0.02</td>
<td></td>
</tr>
<tr>
<td><strong>AUC (mg/L/hr)</strong></td>
<td>—</td>
<td>(Adult) 9.6 ± 2.6</td>
<td>2.8 ± 0.25</td>
<td>423</td>
</tr>
<tr>
<td></td>
<td></td>
<td>(Child) 7.1 ± 2.6</td>
<td>2.8 ± 0.25</td>
<td></td>
</tr>
<tr>
<td><strong>Half-life (hr)</strong></td>
<td>24–50</td>
<td>32 ± 5.6</td>
<td>45 ± 6.3</td>
<td>α phase 1.7</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>β phase 23.6</td>
</tr>
</tbody>
</table>

DMPC = dimyristoylphosphatidyl choline; DMPG = dimyristoylphosphatidyl glycerol; DSPC = distearoylphosphatidyl choline; HSPC = hydrogenated soy phosphatidyl choline.

*AUC values normalized to a dosage of 1 mg/kg/day.

From references 16, 28, 29, 30, and 31 and product information.*
**Pharmacology.** Caspofungin is an echinocandin antifungal that is a specific non-competitive inhibitor of β-(1-3) glucan synthetase in fungal cell membranes. This action leads to a weakened cell wall and eventual cell lysis and death. It is active against *Candida* and *Aspergillus* spp., and *Pneumocystis carinii* with little cross-resistance with the azoles.

**Adult Dosage.** *IV for refractory invasive aspergillosis* 70 mg on day 1, then 50 mg/day. Infuse doses over 1 hr. Do not mix with dextrose-containing solutions. Some evidence supports a 70 mg/day dose in patients unresponsive to 50 mg/day. In moderate hepatic impairment, give 35 mg/day after the 70 mg loading dose; no experience exists in severe hepatic impairment. Safety and efficacy not established under 18 yr.

**Dosage Forms.** *Inj* 50, 70 mg.

**Pharmacokinetics.** Caspofungin is about 97% plasma protein bound and extensively distributed in tissues. It is slowly metabolized by hydrolysis and N-acetylation. Less than 2% is excreted unchanged in urine. The principle half-life is 9–11 hr and accounts for most elimination; a longer 40–50 hr half-life is also reported.

**Adverse Reactions.** Caspofungin has been well tolerated in limited studies, with headache, fever, nausea, vomiting, flushing, pruritus and infusion vein complications most commonly reported. Some effects may be related to histamine release. One case of anaphylaxis has been reported. Elevation of liver function tests has been reported, especially with concurrent cyclosporine.

**Drug Interactions.** Caspofungin can reduce tacrolimus levels by about 20%. Cyclosporine increases caspofungin AUC by 35% and causes transient increases in ALT and AST. Concomitant use of cyclosporine and caspofungin is not recommended. Caspofungin does not inhibit any P450 enzymes, is not a substrate for these enzymes and does not induce CYP3A4. Some inducers of drug metabolism appear to decrease caspofungin levels; consider using the 70 mg/day dosage in patients who do not respond while on an inducer.

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**CLOTRIMAZOLE**

**Pharmacology.** Clotrimazole is an imidazole used for local therapy of fungal infections. The topical formulations are equivalent to other topical antifungals in the treatment of *Candida* spp. or dermatophyte skin infections. (See Topical Antifungals Comparison Chart.)

**Adult Dosage.** *Top for tinea infections* apply to affected area bid. *Vag Tab for vulvovaginal candidiasis* 100 mg/day at bedtime for 7 days; or 2 100 mg tablets once daily at bedtime for 3 days; or 1 500 mg tablet once at bedtime. *Vag Crm for vulvovaginal candidiasis* 1 applicatorful (50 mg) at bedtime for 6–14 days. *PO to treat oropharyngeal candidiasis* dissolve 10 mg troche in the mouth 5 times/day; *PO for prophylaxis of oral candidiasis in patients receiving immunosuppressive drugs* dissolve 10 mg troche in the mouth tid.
**Pediatric Dosage.** Top same as adult dosage. **Troche (<3 yr) safety and efficacy not established; (≥3 yr) same as adult dosage.** **Vag Crm, Tab (<12 yr) safety and efficacy not established; (≥12 yr) same as adult dosage.**

**Dosage Forms.** Top Crm, Top Lot, Top Soln, Vag Crm 1%; **Troche 10 mg; Vag Tab 100, 500 mg. Combination Packages Combination Packages (Gyne-Lotrimin 3) Vag Supp 200 mg (#3) and Vag Crm 1%; (Gyne-Lotrimin 7) Vag Supp 100 mg (#7) and Top Crm 1%.

**Adverse Reactions.** Nausea, vomiting, bad taste, and mildly abnormal liver function tests have occurred with oral troche. Vulvovaginal burning, itching, and irritation have been reported with vaginal products. Skin rash occurs occasionally with vaginal or topical use.

**FLUCONAZOLE**

**Pharmacology.** Fluconazole is a triazole antifungal agent that is highly water soluble and active in vivo against many fungal species (especially *Cryptococcus* spp.). The drug is active against *Candida* sp., *Blastoscyces dermatitidis*, *Coccidioides immitis*, and *Histoplasma capsulatum*. Antifungal effects are caused by inhibition of fungal cytochrome P450-dependent enzymes that prevent conversion of lanosterol to ergosterol.40-42

**Administration and Adult Dosage.** PO or IV for oropharyngeal or esophageal candidiasis 200 mg on day 1, then 100 mg/day for 10–14 days. Severe esophageal candidiasis may require up to 400 mg/day.40,43 **PO or IV for cryptococcal meningitis:** short-term therapy 400 mg/day for 6–10 weeks; maintenance therapy in patients with AIDS 200 mg/day indefinitely. Dosages up to 1 g/day have been used for cryptococcal meningitis. **PO for uncomplicated vaginal candidiasis** 150 mg as a single dose.42 **PO or IV for coccidioidal meningitis** 400 mg/day indefinitely;44 dosages up to 800 mg/day have been used. **PO or IV for prophylaxis of candidiasis in bone marrow transplantation** 400 mg/day and continued for 7 days after granulocyte count exceeds 1000/μL. Initiate therapy several days before onset of neutropenia.43

**Special Populations.** **Pediatric Dosage.** PO or IV for candidiasis 6 mg/kg once, then 3 mg/kg/day for at least 2 weeks for oropharyngeal candidiasis and at least 3 weeks (or 2 weeks after symptom resolution) for esophageal candidiasis; dosages up to 12 mg/kg/day have been used. **PO or IV for systemic candidiasis** 6–12 mg/kg/day. **PO or IV for treatment or prophylaxis of cryptococcal meningitis** 12 mg/kg once, then 6 mg/kg/day; continue treatment for at least 10–12 weeks after CSF cultures become negative. Prophylaxis in HIV-infected children continues indefinitely.

**Geriatric Dosage.** (>65 yr) although half-life is prolonged, dosage adjustment appears unnecessary, unless renal impairment is severe.51 (See Other Conditions.)

**Other Conditions.** Reduce dosage in impaired renal function: for Clr of 20–50 mL/min, give the usual dose q 48 hr; Clr of 10–19 mL/min, 50–200 mg q 48 hr; Clr <10 mL/min, 50–100 mg q 48 hr. Give a full dose after hemodialysis on dialysis days. Patients on chronic ambulatory peritoneal dialysis may receive 50–200 mg/day.

**Dosage Forms.** Tab 50, 100, 150, 200 mg; Susp 10, 40 mg/mL; Inj 2 mg/mL.
**Patient Instructions.** Take with a meal if stomach upset occurs. Report changes in appetite, dark urine, or light stools.

**Missed Doses.** Take this drug at regular intervals. If you miss a dose of this medicine, take it as soon as you remember. If it is almost time for your next dose, take that dose only and go back to your regular dosage schedule. Leave at least 12 hours between doses. Do not double the dose or take extra.

**Pharmacokinetics.** **Fate.** Rapidly and well absorbed (90%) orally, unaffected by gastric pH. Peak concentrations of 1.8–2.8 mg/L (5.9–9 μmol/L) achieved 2–4 hr after administration of 100–150 mg orally. Plasma protein binding is 11–12%; penetrates well into CSF (>60% of simultaneous serum levels). $V_d$ is 0.65 ± 0.2 L/kg; $Cl$ is 0.015 ± 0.006 L/hr/kg. About 64–90% of a dose is excreted unchanged in urine.41

$t_1/2$. 22 ± 4 hr; 37 hr in patients >65 yr; up to 125 hr in patients with renal impairment.41

**Adverse Reactions.** Occasional nausea, vomiting, diarrhea, abdominal pain, or elevations of liver transaminases occur. Severe hepatitis or exfoliative skin reactions occur rarely.40,43

**Precautions.** Observe patients who develop rash for worsening of the lesions and discontinue the drug if necessary.

**Drug Interactions.** Rifampin induces the metabolism of fluconazole and can lead to clinical failure. Fluconazole inhibits metabolism of phenytoin, warfarin, and, to a minor extent, cyclosporine. Low dosages have been shown to increase the serum levels of tolbutamide, glipizide, glyburide, and possibly other sulfonylureas. This could lead to a greater hypoglycemic effect, and dosage reduction might be necessary.43

**Parameters to Monitor.** Liver function tests weekly initially, then monthly. Monitor renal function tests weekly if abnormal at outset of therapy. (See Precautions). Monitor patients with elevated transaminases more carefully for hepatitis.

**Notes.** Combination therapy with fluconazole and fluycytosine for treatment of cryptococcal meningitis appears to be superior to single-agent therapy;46 further studies of this combination and of fluconazole plus amphotericin B are needed. Fluconazole-resistant *Candida albicans* has been clinically demonstrated; increased use of prophylactic fluconazole increases the likelihood of the emergence of resistant strains such as *Candida krusei*.43

**FLUCYTOSINE**

**Pharmacology.** Fluycytosine (5-FC) is a fluorinated cytosine analogue that appears to be deaminated to the cytotoxic antimetabolite fluorouracil by cytosine deaminase, an enzyme present in fungal but not in human cells. It has a narrow spectrum of activity and is used with other antifungals because resistance develops rapidly when used alone in *Candida* and *Cryptococcus* sp. infections.39

**Administration and Adult Dosage.** PO 50–150 mg/kg/day in 4 divided doses; the use of higher dosages has been suggested to prevent the emergence of resistance. Duration of therapy must be guided by the severity of infection and response to therapy.
Special Populations. Pediatric Dosage. PO same as adult dosage in mg/kg.

Geriatric Dosage. Same as adult dosage but adjust for age-related reduction in renal function.

Other Conditions. Reduce dosage in impaired renal function. An approximate dosage reduction can be determined by administering doses at intervals in hours equal to 4 times the Cr, in mg/dL. Alternative regimens such as reduced doses at 6-hr intervals have been recommended. In patients on maintenance hemodialysis q 48–72 hr, give 20–50 mg/kg after each dialysis.39,45 Use normal dosage in liver disease.

Dosage Forms. Cap 250, 500 mg.

Patient Instructions. Take the capsules required for a single dose over a 15-minute period with food to minimize stomach upset.

Missed Doses. Take this drug at regular intervals. If you miss a dose of this medicine, take it as soon as you remember. If it is almost time for your next dose, take that dose only and go back to your regular dosage schedule. Leave at least 4 hours between doses. Do not double the dose or take extra.

Pharmacokinetics. Serum Levels. Toxicity most likely >100 mg/L (780 μmol/L). (See also Precautions.)

Fate. Rapidly and well absorbed (about 90%), with peak about 1–2 hr after administration of a 500 mg dose to adults averaging 8–12 mg/L (62–93 μmol/L) in patients with normal renal function. Negligible binding to plasma proteins; Vd is 0.7 L/kg. Widely distributed throughout the body, including the CSF and eye. Eliminated almost entirely (average 90%) in the urine by glomerular filtration unchanged, with urine levels many times greater than serum levels. Low serum concentrations of fluorouracil have been found in patients taking flucytosine and may be responsible for hematologic toxicity.39,45

\[ t_{1/2} \approx 6 \pm 0.6 \text{ hr; up to 100 hr or greater with renal impairment.} \] \[ 39,45 \]

Adverse Reactions. Occasional nausea, vomiting, diarrhea, bone marrow suppression (often dose limiting in HIV-infected patients), and elevated liver function tests (usually asymptomatic and rapidly reversible). Diarrhea occurs occasionally; ulcerating enteritis occurs rarely.39,45

Precautions. Pregnancy; severe renal impairment (elimination is highly variable and monitoring of serum levels is recommended; keep peak concentrations <100 mg/L); impaired hepatic function; hematologic disorders; or history of therapy with myelosuppressive drugs (eg, zidovudine, ganciclovir, cancer chemotherapy) or radiation.39,45

Drug Interactions. Amphotericin B can increase the toxicity of flucytosine by increasing its cellular penetration and impairing its elimination secondary to nephrotoxicity.

Parameters to Monitor. Before and, frequently during, therapy, monitor BUN, Cr, Clcr, full hematology, and liver function tests. (See also Precautions.)

Notes. Flucytosine may be synergistic with amphotericin B, depending on the organism involved; the combination is useful in treating cryptococcal meningitis in AIDS and non-AIDS patients,46 although the superiority of the combination in
AIDS patients has not been established. Flucytosine might be additive or synergistic with fluconazole for the treatment of cryptococcal meningitis; however, further experience in clinical trials is needed before this combination can be recommended.

**GRISEOFULVIN**

**Pharmacology.** Griseofulvin is a fungistatic agent that appears to affect mitosis in fungal cells. It is active against dermatophytes and not useful in the treatment of yeast or other fungal infections.

**Adult Dosage.** PO (microsize) 0.5–1 g/day in a single or 2–4 divided doses; (ultramicrosize) 330–660 mg/day in 1–2 divided doses. Therapy usually must be continued for at least 3 weeks; infections of the palms or soles require 4–8 weeks of therapy; nail infections usually require 6–12 months of therapy. Instruct patients to take the drug with meals to enhance absorption, avoid prolonged sun exposure, and avoid alcohol.

**Pediatric Dosage.** (Microsize) 11 mg/kg/day; (ultramicrosize) 7.3 mg/kg/day, given as for adults.

**Dosage Forms.** (Microsize) Cap 250 mg; Tab 250, 500 mg; Susp 25 mg/mL; (ultramicrosize) Tab 125, 165, 250, 330 mg.

**Adverse Reactions.** Adverse reactions include occasional nausea and vomiting. Photosensitivity reactions, peripheral neuritis, and leukopenia are rare. The drug can exacerbate acute intermittent porphyria.

**ITRACONAZOLE**

**Pharmacology.** Itraconazole is a synthetic triazole antifungal agent that is more active than ketoconazole or fluconazole against certain fungi, notably Aspergillus spp. It also has activity against Coccidioides, Cryptococcus, Candida, Histoplasma, Blastomyces, and Sporotrichosis spp. Itraconazole inhibits fungal cytochrome P450-dependent enzymes. This inhibition blocks ergosterol biosynthesis, creating disturbances in membrane function and membrane-bound enzymes and affecting fungal cell growth and viability.

**Administration and Adult Dosage.** PO for systemic fungal infections 200–600 mg/day, depending on site and severity of infection. Give dosages over 200 mg/day in 2–3 divided doses. PO for vulvovaginal candidiasis 200 mg bid for 1 day or 200 mg/day for 7 days. PO for dermatomycoses 100 mg/day for 15 days or 200 mg/day for 7 days. PO for pityriasis versicolor 200 mg/day for 7 days. PO for plantar tinea pedis and palmar tinea manuum 100 mg/day for 30 days or 200 mg bid for 7 days. PO for onychomycosis 200 mg once daily for 3 months. IV for blastomycosis, histoplasmosis or aspergillosis 200 mg bid for 4 doses, then 200 mg/day.

**Special Populations.** **Pediatric Dosage.** Safety and efficacy not established.

**Geriatric Dosage.** Same as adult dosage.

**Other Conditions.** Dosage reduction in patients with hepatic impairment might be necessary, but guidelines are not established. No dosage adjustment is necessary in renal impairment. However, the manufacturer recommends that the injection not be used in patients with Clcr <30 mL/min.
Dosage Forms. Cap 100 mg; Soln 10 mg/mL; Inj 10 mg/mL.

Patient Instructions. Take this drug with food to ensure maximal absorption. Do not take with medications that decrease stomach acid (eg, antacids, H₂-blockers, omeprazole). Report symptoms of fatigue, loss of appetite, nausea, vomiting, yellowing of the skin, dark urine, or pale stools.

Missed Doses. Take this drug at regular intervals. If you miss a dose of this medicine, take it as soon as you remember. If it is almost time for your next dose, take that dose only and go back to your regular dosage schedule. Leave at least 12 hours between doses. Do not double the dose or take extra.

Pharmacokinetics. Serum Levels. Levels <5 mg/L (<7 μmol/L) are associated with treatment failure in Aspergillus infections.\textsuperscript{50}

Fate. Relative oral bioavailability of the capsules compared with an oral solution is >70%.\textsuperscript{43} The solubility of itraconazole is aided by an acidic environment, and food increases absorption. Peak serum concentration occurs in 4–5 hr; peak concentration is 20 μg/L (28 nmol/L) after a single 100 mg oral dose during fasting, increasing to 180 μg/L (0.26 μmol/L) when taken with food.\textsuperscript{48} The drug is >99% protein bound, primarily to albumin, with only 0.2% available as free drug.\textsuperscript{48} It is highly lipid soluble, and concentrations are much higher in tissues than in serum. Itraconazole is metabolized in the liver and exhibits dose-dependent elimination.\textsuperscript{43} One metabolite, hydroxyitraconazole, has antifungal activity, and serum concentrations are double those of itraconazole at steady state.

\( t_{1/2} \) 24–42 hr; possibly longer with large daily dosages.\textsuperscript{43}

Adverse Reactions. Itraconazole is generally well tolerated with long-term use. It has a negative inotropic effect and can worsen CHF. Occasional rash, pruritus, nausea, vomiting, abdominal discomfort, headache, dizziness, decreased libido, and hypertension occur. Mild transient elevations of transaminases occur frequently. Hepatotoxicity is rare, but deaths have occurred. There are no apparent adverse effects on testicular or adrenal steroidogenesis.\textsuperscript{43,48}

Contraindications. Coadministration with astemizole, cisapride, oral midazolam, pimozide, quinidine, doxetilide, triazolam or HMG-CoA reductase inhibitors metabolized by CYP3A4.

Precautions. Pregnancy; lactation. Treatment of onychomycosis in patients with ventricular dysfunction (eg, CHF).

Drug Interactions. Itraconazole inhibits CYP3A3/4 and inhibits metabolism of certain drugs such as cyclosporine and warfarin. (See Contraindications.) Warfarin dosage reduction might be necessary during concurrent use. Cyclosporine dosage might need to be reduced by 50% with itraconazole dosages over 100 mg/day. Avoid concurrent carbamazepine, phenytoin, or rifampin because they can dramatically reduce the serum itraconazole concentration.\textsuperscript{50,51}

Parameters to Monitor. Closely monitor prothrombin time in patients on concurrent warfarin and cyclosporine levels in patients taking these drugs. Monitor liver function tests in patients with pre-existing hepatic impairment. Monitoring serum drug concentrations can be helpful if poor absorption or increased metabolism of itraconazole is suspected.
KETOCONAZOLE

Pharmacology. Ketoconazole is an imidazole antifungal agent that exerts its antifungal effects through inhibition of the synthesis of ergosterol (a fungal cell wall component) by inhibiting fungal cytochrome P450. It is used primarily for mucocutaneous fungal infections, including candidiasis, and in tinea versicolor unresponsive to topical therapy. It is used to treat blastomycosis, histoplasmosis, and paracoccidioidiomycosis in immunocompetent patients. It appears to suppress rather than eliminate coccidioidomycosis. Because of its poor CSF penetration, ketoconazole is not recommended for fungal infections of the CNS. Because of its effects on steroid synthesis (see Adverse Reactions), the drug has been used in prostatic cancer and Cushing syndrome.

Administration and Adult Dosage. PO 200–400 mg daily or bid, depending on site and severity of infection. Top apply once daily or bid for dermatophytoses, superficial mycoses, or seborrheic dermatitis. Top for dandruff apply shampoo twice weekly for 4 weeks.

Special Populations. Pediatric Dosage. PO (<2 yr) not established; (>2 yr) 3.3–6.6 mg/kg/day in 1 or 2 divided doses. The drug is bioavailable when tablets are crushed and mixed in applesauce or juice. Top apply once daily.

Geriatric Dosage. Same as adult dosage.

Other Conditions. Limited data suggest that dosage adjustment is unnecessary in patients with hepatic impairment; however, definitive studies are needed. No adjustment is necessary in renal dysfunction.

Dosage Forms. Tab 200 mg; Crm 2%; Shampoo 1, 2%.

Patient Instructions. This drug may be taken with meals if stomach upset occurs, but do not take with medications that decrease stomach acid (eg, antacids, H2 blockers, omeprazole). Report symptoms of fatigue, loss of appetite, yellowing of the skin, dark urine, or pale stools. Taking this drug with an acidic beverage (eg, a cola drink) can increase the absorption substantially. In patients receiving the drug in 0.1 N HCl to promote absorption, the solution should be sipped through a straw to avoid damaging the teeth.

Missed Doses. Take this drug at regular intervals. If you miss a dose of this medicine, take it as soon as you remember. If it is almost time for your next dose, take that dose only and go back to your regular dosage schedule. Leave at least 12 hours between doses. Do not double the dose or take extra.

Pharmacokinetics. Fate. Bioavailability is about 75% and is dose dependent. An acidic environment is necessary for dissolution and absorption. Bioavailability appears to be decreased by 20–40% when the drug is administered with food and is even more markedly reduced if gastric pH is elevated. Poor absorption can occur in AIDS patients because of achlorhydria and other pathologic changes in the GI tract. Peak serum levels of 3.4 ± 0.3 mg/L (6.4 ± 0.6 μmol/L) are attained after a 200 mg dose taken with a meal. The drug is 93–96% plasma protein bound. Vd is estimated to be 0.36 ± 0.1 L/kg with a single dose, increasing to 2.4 ± 1.6 L/kg during long-term therapy; Cl is estimated to be 0.5 ± 0.25 L/hr/kg during long-
term therapy. Ketoconazole is extensively metabolized by the liver to inactive metabolites, with only 2–4% of a dose excreted unchanged in urine.\textsuperscript{39,52,53} \( t_{1/2} \) 8.7 ± 0.2 hr after a single dose, decreasing to 3.3 ± 1 hr during long-term therapy.\textsuperscript{39,52}

**Adverse Reactions.** Generally well tolerated, with the most frequent side effects being nausea, vomiting, pruritus, and abdominal discomfort. Hepatotoxicity, including massive hepatic necrosis, occurs occasionally, but mild elevations of transaminases occur frequently. Gynecomastia occurs, probably caused by ketoconazole-induced suppression of testosterone synthesis. Ketoconazole also blocks cortisol production; however, clinically apparent hypoadrenalism occurs rarely. Irritation, pruritus, and stinging can occur with topical use.

**Contraindications.** Co-administration with astemizole or cisapride.

**Precautions.** Pregnancy; lactation.

**Drug Interactions.** Ketoconazole inhibits human CYP3A4 and inhibits metabolism of certain drugs such as cyclosporine, methylprednisolone, and warfarin. (See Contraindications.) Warfarin dosage reduction may be necessary during concurrent use. H\(_2\)-receptor antagonists, antacids, and probably proton-pump inhibitors (eg, omeprazole, lanosprazole) might reduce ketoconazole oral absorption.

**Parameters to Monitor.** Monitor liver function tests before starting therapy and often during therapy. Closely monitor prothrombin time in patients on concurrent warfarin and cyclosporine levels in patients taking this drug.

**Notes.** Aclorhydric patients may be given the drug with glutamic acid hydrochloride or 0.1 N HCl (using a drinking straw) to increase absorption.\textsuperscript{53} An acidic drink (eg, a cola) also may be used to increase ketoconazole absorption by about 65% in achlorhydria.\textsuperscript{54}

**MICONAZOLE**

**MICONAZOLE NITRATE**

**Pharmacology.** Miconazole is an imidazole antifungal agent available in topical preparations and as a solubilized IV preparation in a polyethoxylated castor oil (Cremophor EL).\textsuperscript{39} (See Topical Antifungals Comparison Chart.)

**Adult Dosage.** IV 1.2–3.6 g/day in 3 divided doses, diluted in at least 200 mL of D5W or NS and infused over 30–60 min. Top for tinea infections apply bid. Vag Tab for vulvovaginal candidiasis 100 mg at bedtime for 7 days, or 200 mg hs for 3 days. Vag Crm for vulvovaginal candidiasis 5 g hs for 7 days.

**Pediatric Dosage.** IV (<1 yr) 15–30 mg/kg/day; (1–12 yr) 20–40 mg/kg/day. Do not exceed 15 mg/kg/dose. Top same as adult dosage; Vag Crm, Tab (<12 yr) safety and efficacy not established; (≥12 yr) same as adult dosage.

**Dosage Forms.** Inj 10 mg/mL; Top Crm, Top Spray, Top Pwdr, Vag Crm, 2%; Vag Supp 100, 200 mg. Combination Packages (Monistat Dual-Pak, M-Zole 3 Combination Pak) Vag Supp 200 mg (#3) and Vag Crm 2%; (Monistat 7 Combination Pak) Vag Supp 100 mg (#7) and Vag Crm 2%.

**Adverse Reactions.** Phlebitis, pruritus, nausea, vomiting, fever, chills, and rash are frequent side effects of IV miconazole.
Notes. Because of the serious toxicity (e.g., cardiorespiratory arrest, hyponatremia) of the parenteral preparation (most likely caused by the vehicle) and data challenging the clinical effectiveness of this agent, restrict parenteral use to treating fungal infections known to be resistant to amphotericin B (e.g., Scadosporium apiospermum). Vaginal and topical effects are similar to those of clotrimazole.

Pharmacology. Nystatin is a polyene antifungal agent very similar to amphotericin B but too toxic for parenteral use. Oral absorption is negligible, and there is no absorption through intact skin or mucous membranes.39 (See Topical Antifungals Comparison Chart.)

Adult Dosage. PO for oral candidiasis (Susp) 400,000–600,000 units qid (as a “swish and swallow”); (troches) 200,000–400,000 units 4–5 times/day. Treat for at least 48 hr after oral symptoms have cleared and cultures have returned to normal. Immunocompromised patients require longer therapy (e.g., 10–14 days). The vaginal tablet has been successfully used orally in place of the oral suspension; its slow dissolution allows prolonged contact time. PO for GI candidiasis 500,000–1,000,000 units tid. Vag for candidiasis 100,000 units daily or bid for 2 weeks.

Pediatric Dosage. PO for candidiasis (newborns) 100,000 units qid; (older infants and children) 200,000–400,000 units qid. Top same as adult dosage.

Dosage Forms. PO Tab 500,000 units; PO Troche 200,000 units; Susp 100,000 units/mL; Top Crm, Oint, Pwdr 100,000 units/g; Vag Tab 100,000 units.


Notes. Nyotran (Investigational-Aronex) is an injectable liposomal formulation of nystatin being studied for candidemia, cryptococcal meningitis and aspergillosis.

Terbinafine is a synthetic allylamine antifungal agent that exerts its activity by inhibiting fungal ergosterol synthesis through inhibition of squalene epoxidase. Terbinafine is active orally and topically. It has demonstrated activity against dermatophyte infections but is less active thanazole antifungal against yeast species.39,55 (See Topical Antifungals Comparison Chart.)

Adult Dosage. PO 250 mg once daily for 6 weeks for onychomycosis of fingernails or for 12 weeks for onychomycosis of the toenails. Reduce dosage in severe hepatic or renal dysfunction. Top for tinea corporis or cruris, or cutaneous candidiasis apply cream bid for 1 week; Top for tinea pedis apply solution or spray bid for 1 week, cream may require therapy up to 4 weeks, especially for plantar infections; Top for tinea versicolor apply solution or spray bid for 1 week.

Pediatric Dosage. Safety and efficacy not established <12 yr.

Dosage Forms. Crm 1%; Top Spray 1%; Top Soln 1%; Tab 250 mg.

Pharmacokinetics. Terbinafine is 70–80% orally absorbed regardless of the presence of food. Peak concentrations after 250 and 500 mg oral doses are 0.9 mg/L (3.1 μmol/L) and 2 mg/L (6.9 μmol/L), respectively, within 2 hr. Terbinafine is highly lipophilic and is widely distributed with a Vd of 13.5 L/kg. It is extensively
metabolized to inactive metabolites, and its elimination half-life is 11–16 hr; however, an additional elimination phase of 200–400 hr may reflect the gradual release of terbinafine from adipose tissue.

**Adverse Reactions.** Frequent adverse reactions during oral therapy are dyspepsia, abdominal pain, diarrhea, skin reactions, malaise, lethargy, and taste disturbance. Hepatic failure has been reported rarely with the treatment of onychomycoses. Avoid in patients with liver disease.

**Drug Interactions.** Terbinafine can inhibit CYP2D6 and increase levels of drugs metabolized by this route. Its clearance is increased 100% by rifampin and decreased 33% by cimetidine.

**Parameters to Monitor.** Baseline AST and ALT; repeat if symptoms of hepatotoxicity occur.

**VORICONAZOLE** *(Investigational-Pfizer) Vfend*

**Pharmacology.** Voriconazole is an azole antifungal that is a derivative of fluconazole, with the same mechanism of action. It has superior activity against *Candida albicans*, *C. krusei* and *C. glabrata*. Activity against *Aspergillus* sp. is equivalent to intraconazole. Activity also extends to *Pseudallescheria boydii* and *Scedosporium aspergillum*.

**Adult Dosage.** PO or IV 50–400 mg/day has been used investigationally.

**Pediatric Dosage.** PO or IV Little data, but 7–10 mg/kg/day has been used.

**Dosage Forms.** Not yet available.

**Pharmacokinetics.** Oral bioavailability is 90%. Steady-state plasma levels are 2.1–4.8 mg/L with an oral dosage of 200 mg bid. The drug is 51–67% plasma protein bound; $V_d$ is 2 L/kg. It is metabolized by the liver, primarily by CYP2C9 and 3A4. Elimination half-life is about 6 hr, but the drug can be detected in urine and feces for several days after prolonged therapy. Less than 5% of unchanged drug appears in urine.

**Adverse Reactions.** Reversible mild to moderate dose-related visual disturbances occur frequently. Elevations in hepatic enzymes are also frequent. One case of photosensitivity has been reported. Voriconazole may interact with drugs that affect or are metabolized by CYP2C9 and 3A4, but more data are needed.
# Topical Antifungals Comparison Chart

<table>
<thead>
<tr>
<th>CLASS AND DRUG</th>
<th>DOSAGE FORMS</th>
<th>ADULT DOSAGE**</th>
<th>COMMENTS</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>ALLYLAMINES AND BENZYLAMINES</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Butenafine HCl</td>
<td>Top Crm 1%</td>
<td>Top (tinea pedis) apply daily for 1–4 weeks.</td>
<td>A benzylamine similar to the allylamines.</td>
</tr>
<tr>
<td>Mentax</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Naftifine HCl</td>
<td>Top Crm 1%</td>
<td>Top (tinea) apply bid for 4 weeks.</td>
<td>First agent of allylamine class; response is faster than with imidazoles.</td>
</tr>
<tr>
<td>Naftin</td>
<td>Top Gel 1%</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Terbinafine HCl</td>
<td>Top Crm 1%</td>
<td>Top (tinea cruris or corporis) apply daily–bid for 1–4 weeks; (tinea pedis) apply bid for up to 4 weeks.</td>
<td>Allylamine; 10–100 times more potent than naftifine. Response is more rapid than imidazoles, and it has excellent penetration in tinea pedis.</td>
</tr>
<tr>
<td>Lamisil</td>
<td>Top Crm 1%</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Top Soln 1%</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Top Spray 1%</td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>IMIDAZOLES</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Butoconazole Nitrate</td>
<td>Vag Crm 2%</td>
<td>Vag (nonpregnant) 2% crm hs for 3–6 days; or (Gynazole-1) 2% crm 1 applicatorful once (pregnant, 2nd or 3rd trimester) 2% crm hs for 6 days.</td>
<td>Spectrum similar to other imidazoles.</td>
</tr>
<tr>
<td>Femstat</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Gynazole-1</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Clotrimazole</td>
<td>Top Crm 1%</td>
<td>Top (Candida, tinea) apply bid.</td>
<td>Useful for 1st trimester Trichomonas vaginitis, but less effective than metronidazole.</td>
</tr>
<tr>
<td>Lotrimin</td>
<td>Top Lot 1%</td>
<td>Vag 100 mg supp or 1% crm hs for 7 days; 500 mg supp hs once;</td>
<td></td>
</tr>
<tr>
<td>Mycelex</td>
<td>Top Soln 1%</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Vag Tab 100, 200, 500 mg</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Vag Crm 1%</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Econazole Nitrate</td>
<td>Top Crm 1%</td>
<td>Top (Candida) apply bid; (tinea) apply once daily.</td>
<td>Activity similar to other imidazoles.</td>
</tr>
<tr>
<td>Spectazole</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

*(continued)*
<table>
<thead>
<tr>
<th>CLASS AND DRUG</th>
<th>DOSAGE FORMS</th>
<th>ADULT DOSAGE</th>
<th>COMMENTS</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Ketoconazole</strong></td>
<td>Top Crm 2%</td>
<td>Top (Candida, tinea) apply daily for 2–6 weeks; (seborrhea) apply bid for 4 weeks; (shampoo) twice weekly for 4 weeks, then prn.</td>
<td></td>
</tr>
<tr>
<td>Nizoral</td>
<td>Shampoo 1, 2%</td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Miconazole Nitrate</strong></td>
<td>Top Crm 2%</td>
<td>Top apply bid.</td>
<td>Possibly less effective than some newer topical imidazoles.</td>
</tr>
<tr>
<td>Micatin,</td>
<td>Top Oint 2%</td>
<td>Vag 100 mg supp or 2% crm hs for 7 days or 200 mg supp hs for 3 days.</td>
<td></td>
</tr>
<tr>
<td>Monistat</td>
<td>Vag Crm 2%</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Vag Supp 100, 200 mg</td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Oxiconazole Nitrate</strong></td>
<td>Top Crm 1%</td>
<td>Top (tinea) apply daily–bid for 2–4 weeks.</td>
<td>Similar to other imidazoles; superior to tolnaftate in dermatomycoses.</td>
</tr>
<tr>
<td>Oxistat</td>
<td>Top Lot 1%</td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Sulconazole Nitrate</strong></td>
<td>Top Crm 1%</td>
<td>Top (tinea) apply daily–bid for 3–4 weeks.</td>
<td>Similar to other imidazoles, but superior to miconazole in dermatomycoses.</td>
</tr>
<tr>
<td>Exelderm</td>
<td>Top Soln 1%</td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Terconazole</strong></td>
<td>Vag Crm 0.4, 0.8%</td>
<td>Vag 0.4% crm hs for 7 days; 0.8% crm or supp hs for 3 days.</td>
<td>Similar to other imidazoles, but superior to miconazole in vaginal candidiasis.</td>
</tr>
<tr>
<td>Terazol</td>
<td>Vag Supp 80 mg</td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Tioconazole</strong></td>
<td>Vag Oint 6.5%</td>
<td>Vag hs once.</td>
<td>Possibly more effective than older imidazoles; appears effective in vaginal trichomoniass.</td>
</tr>
<tr>
<td>Vagistat-1</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

**POLYENES**

<table>
<thead>
<tr>
<th>CLASS AND DRUG</th>
<th>DOSAGE FORMS</th>
<th>ADULT DOSAGE</th>
<th>COMMENTS</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Amphotericin B</strong></td>
<td>Top Crm 3%</td>
<td>Top (Candida) apply bid–qid for 1–4 weeks.</td>
<td>Inconvenient application schedule.</td>
</tr>
<tr>
<td>Fungizone</td>
<td>Top Lot 3%</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

(continued)
### TOPOCAL ANTIFUNGALS COMPARISON CHART (continued)

<table>
<thead>
<tr>
<th>CLASS AND DRUG</th>
<th>DOSAGE FORMS</th>
<th>ADULT DOSAGE Comments</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Nystatin</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Mycostatin</td>
<td>Top Crm 100,000 units/g</td>
<td>Top (Candida) apply bid–tid. Similar to amphotericin B.</td>
</tr>
<tr>
<td>Nilstat</td>
<td>Top Oint 100,000 units/g</td>
<td>Vag 1 tab hs for 14 days.</td>
</tr>
<tr>
<td></td>
<td>Top Pwdr 100,000 units/g</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Vag Tab 100,000 units.</td>
<td></td>
</tr>
</tbody>
</table>

### MISCELLANEOUS

<table>
<thead>
<tr>
<th><strong>Ciclopirox Olamine</strong></th>
<th>Top Crm 1%</th>
<th>Top Crm, Lot (Candida, tinea) apply bid. A hydroxypyridone. More effective than clotrimazole for tinea versicolor. Nail lacquer is inexpensive, but has poor efficacy rate.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Loprox</td>
<td>Top Lot 1%</td>
<td>Top Nail Lacquer apply daily. Communicated</td>
</tr>
<tr>
<td>Penlac</td>
<td>Nail Lacquer 8%</td>
<td></td>
</tr>
<tr>
<td><strong>Haloprogin</strong></td>
<td>Top Crm 1%</td>
<td>Top (tinea) apply for 2–4 weeks. Equivalent to tolnaftate.</td>
</tr>
<tr>
<td></td>
<td>Soln 1%</td>
<td></td>
</tr>
<tr>
<td><strong>Tolnaftate</strong></td>
<td>Top Crm 1%</td>
<td>Top (tinea) apply for 2–6 weeks. A thiocarbamate. Possibly slightly less effective than imidazoles in dermatomycoses.</td>
</tr>
<tr>
<td></td>
<td>Top Gel 1%</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Top Soln 1%</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Pwdr 1%</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Spray Liquid 1%</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Spray Pwdr 1%</td>
<td></td>
</tr>
</tbody>
</table>

*The dosage for vaginal creams for candidal infections is one applicatorful at the interval shown. Tinea pedis should be treated at the maximum dosage (usually bid) for the longest time mentioned, usually 4 weeks.*

*From references 39 and product information.*
Antimycobacterial Drugs

CLOFAZIMINE  Lamprène

Pharmacology. Clofazimine is a lipophilic rhimophenazine dye approved for treating leprosy and used in atypical *Mycobacterium* infections, discoid lupus erythematosus, and pyoderma gangrenosum.57,58

**Adult Dosage.** PO for leprosy, *Mycobacterium avium* complex infections, and discoid lupus erythematosus 100 mg/day with food; dosages up to 200 mg/day are used for erythema nodosum leprosum. **PO for pyoderma gangrenosum** 300–400 mg/day have induced remission, but the manufacturer states that dosages >200 mg/day are not recommended.

**Pediatric Dosage.** PO for leprosy 1 mg/kg/day; **PO for *M. avium* complex** 1–2 mg/kg/day.

**Dosage Forms.** Cap 50 mg.

**Pharmacokinetics.** The drug is about 50% bioavailable. A peak serum concentration of 0.5–2 mg/L (1–4 μmol/L) 2 hr after an oral 100 to 200 mg dose is proposed as evidence of adequate absorption. Clofazimine accumulates in fatty tissues and the reticuloendothelial system and is eliminated with a half-life of about 70 days.

**Adverse Reactions.** Bodily secretions, skin, conjunctivae, cornea, urine, and feces can turn red to brownish black; an orange–pink skin discoloration is common and can take months to years to disappear after stopping the drug. Dose-related GI pain, nausea, vomiting, and diarrhea can occur because of crystalline deposits in GI tissue. Eosinophilic enteritis and splenic infarction occur rarely at dosages >100 mg/day. (See also Second-Line Antituberculosis Agents Comparison Chart.)

ETHAMBUTOL  Myambutol

**Pharmacology.** Ethambutol is a tuberculostatic agent that is only active against mycobacteria, including *Mycobacterium avium* complex. It does not directly enhance short course (6–9 months) regimens of isoniazid, rifampin, and pyrazinamide. Ethambutol is recommended to be included as part of a 4-drug initial regimen if there is a possibility of drug resistance and should be continued for 12 months if isoniazid resistance is demonstrated. Ethambutol is also used in combination with clarithromycin to treat disseminated *M. avium intracellulare* (MAI) infection in patients with AIDS.58–62

**Adult Dosage.** **PO for treatment of active tuberculosis** 15–25 mg/kg/day as a single dose given in combination with isoniazid and/or rifampin and/or pyrazinamide. **PO for MAI** 15 mg/kg/day, to a maximum of 1 g/day as a single dose in combination with clarithromycin or azithromycin.

**Pediatric Dosage.** Same as adult dosage.

**Dosage Forms.** Tab 100, 400 mg.
Pharmacokinetics. Ethambutol is about 80% absorbed from the GI tract with complex disposition characteristics. A peak serum concentration of 2–6 mg/L (8–25 μmol/L) 2 hr after an oral 15–25 mg/kg dose is proposed as evidence of adequate absorption. Its half-life is 4–6 hr, increasing to 32 hr in severe renal impairment. Approximately 80% is excreted unchanged in urine.

Adverse Reactions. Adverse reactions are rare with the recommended dosage of 15–25 mg/kg/day. Optic neuritis (manifested as blurred vision, color blindness, and restricted visual fields) occurs rarely with dosages of 15 mg/kg/day and is usually reversible with prompt drug discontinuation. Hyperuricemia can occur because of impairment of uric acid excretion.

Pharmacology. Isoniazid (INH) is a synthetic hydrazine derivative of isonicotinic acid that inhibits the synthesis of mycolic acid, a component of the mycobacterial cell wall; it probably has other actions. Its activity is limited to mycobacteria; it is tuberculostatic or tuberculocidal depending on concentration and reproductive rate of the organism. Resistance is uncommon in preventive therapy but can develop rapidly if used alone in active tuberculosis. Primary resistance is becoming increasingly common in certain communities and has occurred in a variety of institutional settings (eg, hospitals, prisons). These settings are characterized by a high prevalence of HIV infection.

Administration and Adult Dosage. PO for treatment of latent tuberculosis infection 5 mg/kg/day (usually 300 mg) as a single daily dose, to a maximum of 300 mg/day, given as a single agent for 6–9 months. Alternatively, give INH 15 mg/kg/dose (up to 900 mg) twice weekly by directly observed therapy (DOT) for 6–9 months. PO for treatment of active tuberculosis same dosage as above combined with rifampin 600 mg/day and pyrazinamide 15–30 mg/kg/day for 8 weeks, followed by 16 weeks of INH and rifampin. Alternatively, give the doses of INH, rifampin, ethambutol, and pyrazinamide for 2 weeks, followed by INH 15 mg/kg (to a maximum of 900 mg), rifampin 600 mg, ethambutol 50 mg/kg (to a maximum of 2.5 g), and pyrazinamide 50–70 mg/kg (to a maximum of 4 g) in 2 or 3 divided doses twice weekly for a total of 6 weeks by directly observed therapy (DOT), then continue INH and rifampin twice weekly for 16 weeks by DOT. In 3-times-a-week regimens, ethambutol dosage is 25–30 mg/kg/day (to a maximum of 2.5 g), with INH, rifampin, and pyrazinamide at the same doses as in the twice-weekly regimen, but continued for 6 months by DOT. If pyrazinamide cannot be taken, a 9-month course may be administered in which INH in the above dosage is combined with rifampin 600 mg/day. IM or IV (rarely used) same as oral dosage.

Special Populations. Pediatric Dosage. PO for treatment of latent tuberculosis infection 10–20 mg/kg/day as a single dose, to a maximum of 300 mg/day, given as a single agent for 6–9 months. Alternatively, give INH 20–40 mg/kg/dose (up to 900 mg) twice weekly by directly observed therapy (DOT) for 6–9 months. PO for treatment of active tuberculosis same dosage as above, but combine with rifampin 10–20 mg/kg (to a maximum of 600 mg), and pyrazinamide
15–30 mg/kg/day (to a maximum of 2 g) in 2 or 3 divided doses for 8 weeks followed by 16 weeks of INH and rifampin. Alternatively, give the daily doses of INH, rifampin, ethambutol, and pyrazinamide for 2 weeks, followed by INH 20–40 mg/kg (to a maximum of 900 mg), rifampin 10–20 mg/kg (to a maximum of 600 mg), ethambutol 50 mg/kg (to a maximum of 2.5 g), and pyrazinamide 50–70 mg/kg (to a maximum of 4 g) in 2 or 3 divided doses twice weekly for a total of 6 weeks by DOT, then continue INH and rifampin twice weekly for 16 weeks by DOT. In 3-times-a-week regimens, pyrazinamide dosage is 50–70 mg/kg/day (to a maximum of 3 g) in 2–3 divided doses. If pyrazinamide cannot be taken, a 9-month course may be administered in which INH in the above dosage is combined with rifampin 10–20 mg/kg (to a maximum of 600 mg).

**Geriatric Dosage.** Same as adult dosage.

**Other Conditions.** Acetylator phenotype has not been evaluated as a parameter for dosage individualization; however, some sources recommend a dosage of 150–200 mg/day in slow acetylators with renal impairment. In individuals with HIV infection being treated for tuberculosis, treatment regimens are not altered but should continue for a total of 9 months and at least 6 months beyond culture conversion.

**Dosage Forms.** Tab 50, 100, 300 mg; Syrup 10 mg/mL; Cap 150 mg with rifampin 300 mg (Rifamate); Tab 50 mg with rifampin 120 mg and pyrazinamide 300 mg (Rifater); Inj 100 mg/mL.

**Patient Instructions.** Report any burning, tingling, or numbness in the extremities; unusual malaise; fever; dark urine; or yellowing of the skin or eyes.

**Missed Doses.** Take this drug at regular intervals. If you miss a dose of this medicine, take it as soon as you remember. If it is almost time for your next dose, take that dose only and go back to your regular dosage schedule. Leave at least 12 hours between doses. Do not double the dose or take extra.

**Pharmacokinetics.**

**Serum Levels.** A peak serum level of 3–5 mg/L (22–36 µmol/L) 2 hr postdose is proposed as evidence of adequate absorption. Rapid and nearly complete oral absorption with peak serum concentrations of 1–5 mg/L (7–36 µmol/L) 1 hr after a 5 mg/kg dose. Widely distributed in body tissues including the CSF of normal patients and those with meningitis. Vd is 0.67 ± 0.15 L/kg; Cl is 0.22 ± 0.07 L/hr/kg in slow acetylators and 0.44 ± 0.12 L/hr/kg in rapid acetylators. Eliminated primarily by acetylation in the liver to inactive metabolites that are excreted in the urine. Specific pattern of elimination depends on acetylator phenotype of the individual.

**t½.** (Rapid acetylators) 1.1 ± 0.1 hr, (slow acetylators) 2.1 ± 1.1 hr. Increased to 4 hr with renal impairment and 6.7 hr with liver disease.

**Adverse Reactions.** Pyridoxine-responsive peripheral neuropathy can occur, especially in alcoholics, diabetics, patients with renal failure, malnourished patients, and slow acetylators, and with dosages >5 mg/kg/day. Subclinical hepatitis is frequent (10–20%) and characterized by usually asymptomatic elevations of AST and ALT, which can return to normal despite continued therapy; it might be more
frequent with combined INH–rifampin therapy. Clinical hepatitis is rare in those <20 yr, but is strikingly related to age (rising to 2–3% in 50–65 yr-old patients). Rare cases of massive liver atrophy resulting in death usually appear in association with alcoholism or pre-existing liver disease; most severe cases occur within the first 6 months. With acute overdosage (usually 6–10 g), INH can produce severe CNS toxicity including coma and seizures as well as hypotension, acidosis, and occasionally death.

**Contraindications.** Acute or chronic liver disease; previous INH-associated hepatitis.

**Precautions.** Pregnancy; lactation. Use with caution in daily users of alcohol, elderly patients, and those with a slow acetylator phenotype.

**Drug Interactions.** INH can inhibit the metabolism of carbamazepine and phenytoin, increasing the risk of toxicity, particularly of phenytoin in slow acetylators. Mental changes can result from effects of INH and disulfiram on metabolism of adrenergic neurotransmitters; avoid the use of disulfiram in patients who must take INH. Aluminum-containing antacids can interfere with INH absorption. Rifampin can increase the metabolism of INH to hepatotoxic metabolites.

**Parameters to Monitor.** Question for prodromal signs of hepatitis (eg, fever, malaise) and signs of peripheral neuropathy (eg, burning, tingling, numbness) monthly during therapy. Baseline and monthly AST and ALT are recommended only in high-risk groups (those >35 yr, daily alcohol users, and those with a history of liver dysfunction), although they are not predictive of clinical hepatitis.

**Notes.** It is generally recommended that all patients receive INH for treatment of latent tuberculosis infection who have had positive reactions to intermediate-strength purified protein derivative (PPD, 5 tuberculin units) and who (1) are household contacts of patients with active tuberculosis; (2) converted their PPD to positive within the past 12–24 months; (3) have radiologic evidence of inactive tuberculosis or a history of inadequately treated active tuberculosis; (4) are foreign-born persons (and their families) from high-prevalence areas who have entered the United States within the past 2 years; (5) are persons with known or suspected HIV infection; (6) are persons with medical or iatrogenic conditions that increase the risk of tuberculosis—silicosis, gastrectomy, jejunoileal bypass, weight of 10% or more below ideal, chronic renal failure, diabetes mellitus, corticosteroid or other immunosuppressive therapy, hematopoietic malignancy, other malignancy, and other conditions in which immunosuppression results from the disease or its treatment. Most sources suggest that the use of INH prophylaxis in patients >35 yr should be further restricted because of the increased risk of fatal hepatotoxicity, although this is controversial.

To prevent peripheral neuropathy, give pyridoxine in a dosage of 50 mg/day to adults receiving large dosages of INH (10 mg/kg/day or more) and those who are predisposed to peripheral neuritis (eg, diabetics, HIV-infected, alcoholics). Pyridoxine IV in a dosage equal to the estimated amount of INH ingested is recommended for acute INH overdose.

Add ethambutol or streptomycin to the initial treatment regimen until drug susceptibility studies are available, or unless there is little possibility of drug resis-
tance (ie, there is <4% primary resistance to INH in the patient’s community, and
the patient has had no previous treatment with antituberculosis medications, is not
from a country with a high prevalence of drug resistance, and has no known expo-
sure to a drug-resistant case).60,69

**PYRAZINAMIDE**

**Pharmacology.** Pyrazinamide is a synthetic analogue of niacinamide that is only
active against mycobacteria. The mode of action is unknown. The drug is most ac-
tive at acid pH and is active against intracellular organisms. Resistance develops
rapidly when used alone, but no cross-resistance with isoniazid is observed.60–63

**Adult Dosage.** PO for treatment of latent tuberculosis infection 15–
20 mg/kg/day (to a maximum of 2 g) in combination with rifampin 10 mg/kg/day
(to a maximum of 600 mg) as a single daily dose for 2 months. Alternatively, give
pyrazinamide 50 mg/kg/dose (to a maximum of 4 g) in combination with rifampin
10 mg/kg/dose (to a maximum of 600 mg) twice weekly for a total of 2–3 months
by DOT. (See Treatment of Latent Tuberculosis Infection Comparison Chart.) PO
for treatment of active tuberculosis (see Isoniazid Dosage).

**Dosage Forms.** Tab 500 mg; Tab 300 mg with isoniazid 50 mg and rifampin 120
mg (Rifater).

**Pharmacokinetics.** The drug is well absorbed from the GI tract with serum con-
centrations of 40–50 mg/L (0.3–0.4 mmol/L) achieved about 2 hr after a 1 g dose.
A peak serum concentration of 20–60 mg/L (163–488 H9262 mol/L) 2 hr after an oral
1–2 g dose is proposed as evidence of adequate absorption. The parent compound
and several metabolites are excreted in urine.

**Adverse Reactions.** Frequent hyperuricemia, probably caused by prevention of
uric acid excretion by one of the metabolites, and occasional dose-dependent hep-
atotoxicity occur. As many as 1–5% of patients taking regimens including isoni-
azid, rifampin, and pyrazinamide develop laboratory evidence of hepatic damage.

**RIFABUTIN**

**Pharmacology.** Rifabutin is a rifamycin similar to rifampin chemically and in an-
tibacterial spectrum. Rifabutin is more active against mycobacteria than rifampin,
including some rifampin-resistant strains of *Mycobacterium tuberculosis* and
atypical mycobacteria, and is particularly active against MAI.70–73

**Adult Dosage.** PO for prophylaxis of MAI infections in patients with ad-
vanced HIV infection 300 mg/day. PO for treatment of active tuberculosis 300
mg/day as a single daily dose in combination with at least one other antitubercular
agent. (See Notes.)

**Dosage Forms.** Cap 150 mg.

**Pharmacokinetics.** Well absorbed orally, but rifabutin has a low bioavailability
of 12–20% because of first-pass metabolism. Rifabutin is widely distributed in
the body and is concentrated intracellularly to a greater extent than rifampin. It is 71 ±
2% plasma protein bound and has an estimated *Vd* of 45 ± 17 L/kg and Cl of 0.69
± 0.32 L/hr/kg. The drug is hepatically metabolized to a number of compounds,
with about 10% excreted unchanged in urine. It induces its own metabolism; its terminal half-life after long-term use is 45 ± 16 hr.

**Adverse Reactions.** The most frequent adverse reactions are rash, taste alterations, anorexia, nausea, insomnia, nervous system disorders (facial paralysis, twitching, and peripheral neuritis), leukopenia, and hyperbilirubinemia. Uveitis has occurred with dosages >300 mg/day.

**Drug Interactions.** Rifabutin induces the metabolism of drugs metabolized via CYP3A4; although the clinical importance of this effect is not clear, it appears to be less than that of rifampin.

**Notes.** Rifabutin can be substituted for rifampin in antituberculosis regimens.

**Pharmacology.** Rifampin is a synthetic rifamycin B derivative that inhibits the action of DNA-dependent RNA polymerase. It is highly active against mycobacteria, most Gram-positive bacteria, and some Gram-negative bacteria, most notably *Neisseria meningitidis*. It is also used to enhance bactericidal activity of other antistaphylococcal agents in refractory or chronic infections. Antagonism with vancomycin is observed in vitro but is probably not clinically relevant. Primary resistance is uncommon, but resistance can develop rapidly if used alone.

**Administration and Adult Dosage.** PO for treatment of latent tuberculosis infection 10 mg/kg/day (to a maximum of 600 mg) as a single daily dose for a total of 4 months. Alternatively, rifampin can be combined with pyrazinamide. (See pyrazinamide dosage and Treatment of Latent Tuberculosis Infection Comparison Chart.) **PO or IV (rarely used) for treatment of tuberculosis** 600 mg/day as a single daily dose in combination with at least one other antitubercular agent. (See Isoniazid.) **PO for prophylaxis of meningococcal meningitis** 600 mg/day for 4 days or 600 mg bid for 2 days. **PO for staphylococcal infection** 600 mg/day as a single dose in combination with another antistaphylococcal agent.

**Special Populations. Pediatric Dosage.** PO for treatment of tuberculosis (>5 yr) 10–20 mg/kg/day as a single daily dose, to a maximum of 600 mg/day, in combination with at least one other antitubercular agent. **PO for prophylaxis of meningococcal meningitis** (<1 month) 5 mg/kg bid for 2 days; (1 month–12 yr) 10 mg/kg/day, to a maximum of 600 mg/day for 4 days, or 10 mg/kg bid, to a maximum of 600 mg bid for 2 days.

**Geriatric Dosage.** Same as adult dosage.

**Other Conditions.** Accumulation is expected in patients with hepatic dysfunction or biliary obstruction, but dosage guidelines are not available. No dosage adjustment is necessary in patients with impaired renal function.

**Dosage Forms.** **Cap** 150, 300 mg; **Cap** 300 mg with isoniazid 150 mg (Rifamate); **Tab** 120 mg with isoniazid 50 mg and pyrazinamide 300 mg (Rifater); **Inj** 600 mg.

**Patient Instructions.** Take this medication with a full glass of water on an empty stomach (1 hour before or 2 hours after meals) for best absorption. It is important
to take this medication regularly as directed because inconsistent use might increase its toxicity. This drug can cause harmless red–orange discoloration of sweat, tears (it can permanently discolor soft contact lenses), saliva, feces, and urine.

**Missed Doses.** Take this drug at regular intervals. If you miss a dose of this medicine, take it as soon as you remember. If it is almost time for your next dose, take that dose only and go back to your regular dosage schedule. Leave at least 12 hours between doses. Do not double the dose or take extra.

**Pharmacokinetics. Serum Levels.** A peak serum level of 8–24 mg/L (10–29 μmol/L) 2 hr after a 600–750 mg oral dose is proposed as evidence of adequate absorption.\(^5\)

**Fate.** 100% absorbed orally, with a 600 mg dose producing a peak serum concentration of approximately 10 mg/L (12 μmol/L) 1–3 hr after administration. Food delays absorption but does not affect overall bioavailability. First-pass hepatic extraction is substantial but saturated with doses >300–450 mg; thus, larger doses produce disproportionate increases in serum levels. Widely distributed throughout the body; however, useful amounts appear in the CSF only in the presence of inflamed meninges. About 80% plasma protein bound; \(V_d\) is 0.97 ± 0.36 L/kg; Cl is 0.21 ± 0.1 L/hr/kg. Eliminated primarily by deacetylation in the liver to a partially active metabolite that is extensively enterohepatically recirculated, producing very high biliary concentrations. About 50–60% of a dose is eventually excreted in the feces. Urinary excretion is variable and appears to increase with the dose. At usual dosages, 12–15% is excreted unchanged in the urine.\(^7\)

**\(t_{1/2}\).** 3.5 ± 0.8 hr. Half-life increases with higher doses but can become shorter over the first few weeks of treatment. It is not changed by renal impairment but is increased unpredictably by liver disease or biliary obstruction.\(^7\)

**Adverse Reactions.** Adverse reactions are more frequent and severe with intermittent, high-dose administration. GI symptoms are frequent. Acute, reversible renal failure, characterized as tubular damage with interstitial nephritis, sometimes appearing with concomitant hepatic failure has been reported rarely, especially in association with intermittent administration.\(^6\)–\(^6\) Asymptomatic elevation of liver enzymes occurs frequently, whereas clinical hepatitis is rare but more common with pre-existing liver disease or alcoholism; the effect of isoniazid coadministration on the frequency of hepatitis is unclear.\(^6\) Competition with bile for biliary excretion can produce jaundice, especially with pre-existing liver disease. Intermittent therapy is also associated with thrombocytopenia and a flu-like syndrome (ie, fever, joint pain, muscle cramps).

**Contraindications.** Hypersensitivity to any rifamycin derivative.

**Precautions.** Pregnancy; lactation. Use with caution in daily users of alcohol, those with pre-existing liver disease, and those with a history of drug-associated hepatic damage (especially from antituberculurs).

**Drug Interactions.** Rifampin accelerates the metabolism of many drugs such as oral contraceptives, corticosteroids, cyclosporine, enalapril, HIV protease inhibitors, propranolol, methadone, metoprolol, mexiletine, phenytoin, quinidine, theophylline, tolbutamide, oral verapamil, warfarin, and zidovudine because of
potent inducing effects on CYP3A.\textsuperscript{76} The dosage of these drugs may need to be increased during concurrent use. Rifampin can increase the metabolism of isoniazid to hepatotoxic metabolites.

**Parameters to Monitor.** Question for prodromal signs of hepatitis (eg, fever, malaise). Baseline and monthly AST and ALT have been recommended, especially for patients with factors predisposing to hepatotoxicity (eg, alcoholism, pre-existing liver disease), although they are not predictive of clinical hepatitis in the absence of symptoms.

**Notes.** Rifampin is a useful drug for tuberculosis but should be used only in combination regimens because of rapid emergence of resistant mutants of *Mycobacterium tuberculosis* when it is used alone. The recent emergence of multiple drug resistance among strains of *M. tuberculosis* in patients with AIDS includes high-level rifampin resistance. The routine use of rifampin in methicillin-resistant *Staphylococcus aureus* (MRSA) endocarditis is not recommended except after failure of conventional therapy and possibly with renal, myocardial, splenic, or cerebral abscess. If rifampin is added to vancomycin for treatment of MRSA, add a third drug (eg, gentamicin) to reduce the likelihood of resistance development. In nonendocarditis infections caused by MRSA, do not use rifampin unless there is inadequate response to vancomycin alone.

**RIFAPENTINE**

**Pharmacology.** Rifapentine is a rifamycin, similar to rifampin and rifabutin. It is used in the treatment of pulmonary tuberculosis and is similar in efficacy to daily rifampin, although relapse rates can be greater with rifapentine.\textsuperscript{77}

**Adult Dosage.** PO for *tuberculosis* 600 mg twice weekly for 2 months with at least 72 hr between doses, then once weekly for 4 months. It should be used in conjunction with other antitubercular drugs in both phases. The drug may be taken with food to decrease nausea, vomiting, or GI upset.

**Dosage Forms.** Tab 150 mg.

**Pharmacokinetics.** Peak serum levels occur 5–6 hr after an oral dose. Food increases the bioavailability and peak serum level. Rifapentine is 93% bound to serum albumin. *V$_d$* is estimated to be 70 ± 1 L/kg. *Cl* is 2.5 ± 0.14 L/hr/kg in males and 1.7 ± 0.41 L/hr/kg in females. The drug is hydrolyzed by an esterase to the active metabolite 25-desacetyl rifapentine. Half-lives of the drug and metabolite are each about 13 hr.

**Adverse Reactions.** The most frequent side effects in combination regimens were neutropenia, leukopenia, increased liver enzymes, dyspepsia, and anorexia, although these appeared to be less frequent than in equivalent rifampin-containing regimens. Pyuria and hematuria occurred more frequently with rifapentine than with rifampin. Obtain baseline liver enzymes, bilirubin, CBC, and platelet count before starting therapy. Routine laboratory monitoring during therapy is not necessary unless clinically indicated.

**Contraindications.** Hypersensitivity to any rifamycin.
Drug Interactions. Rifapentine induces CYP2C8/9 and 3A4 and can increase the metabolism of drugs metabolized by these isozymes. Rifapentine decreases indinavir peak by 55% and AUC by 70%. Use with great caution in conjunction with protease inhibitors and other drugs metabolized by CYP2C8/9 or 3A4.

<table>
<thead>
<tr>
<th>DRUGS</th>
<th>DURATION (MONTHS)</th>
<th>DOSAGE INTERVAL</th>
<th>HIV- RATING (EVIDENCE)</th>
<th>HIV+ RATING (EVIDENCE)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Isoniazid</td>
<td>9</td>
<td>Daily</td>
<td>A (II)</td>
<td>A (II)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Twice weekly</td>
<td>B (II)</td>
<td>B (II)</td>
</tr>
<tr>
<td>Isoniazid</td>
<td>6</td>
<td>Daily</td>
<td>B (I)</td>
<td>C (I)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Twice weekly</td>
<td>B (II)</td>
<td>C (I)</td>
</tr>
<tr>
<td>Rifampin-pyrazinamide</td>
<td>2</td>
<td>Daily</td>
<td>B (II)</td>
<td>A (I)</td>
</tr>
<tr>
<td></td>
<td>2–3</td>
<td>Twice weekly</td>
<td>C (II)</td>
<td>C (I)</td>
</tr>
<tr>
<td>Rifampin</td>
<td>4</td>
<td>Daily</td>
<td>B (II)</td>
<td>B (III)</td>
</tr>
</tbody>
</table>

Rating: A = preferred; B = acceptable alternative; C = offer when A and B cannot be given. Evidence: I = randomized clinical trial data; II = data from clinical trials that are not randomized or were conducted in other populations; III = expert opinion.

From reference 65.
### SECOND-LINE ANTITUBERCULOSIS AGENTS COMPARISON CHART

<table>
<thead>
<tr>
<th>DRUG</th>
<th>DOSAGE FORMS</th>
<th>ADULT DOSAGE</th>
<th>PEDIATRIC DOSAGE</th>
<th>SERUM LEVELS (MG/L)</th>
<th>HALF-LIFE</th>
<th>MAJOR ADVERSE EFFECTS</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Aminosalicylic Acid Salts</strong>&lt;sup&gt;a&lt;/sup&gt;</td>
<td>Tab 500 mg</td>
<td>PO 8–12 g/day in 2–4 divided doses (as the acid).</td>
<td>150–300 mg/kg/day in 3–4 divided doses, to a maximum of 12 g/day.</td>
<td>20–60&lt;sup&gt;b&lt;/sup&gt; (4 g)</td>
<td>1 hr</td>
<td>—</td>
</tr>
<tr>
<td><strong>Streptomycin Sulfate</strong></td>
<td>Gran 4 g.</td>
<td>Same as streptomycin.</td>
<td>Same as streptomycin.</td>
<td>2.5 hr</td>
<td></td>
<td>Nephrotoxicity; ototoxicity.</td>
</tr>
<tr>
<td><strong>Capreomycin</strong></td>
<td>Inj 1 g.</td>
<td>Same as streptomycin.</td>
<td>Same as streptomycin.</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Lamprene</strong></td>
<td>Cap 50 mg.</td>
<td>PO 100–200 mg/day.</td>
<td>Not well established.</td>
<td>0.5–2 (100–200 mg)</td>
<td>70 days</td>
<td>—</td>
</tr>
<tr>
<td><strong>Cycloserine</strong></td>
<td>Cap 250 mg.</td>
<td>PO 15–20 mg/kg/day (usually 500 mg) in 2 divided doses, to a maximum of 1 g/day.</td>
<td>PO 10–15 mg/kg/day to a maximum of 1 g/day.</td>
<td>20–35 (250–500 mg)</td>
<td>10 hr</td>
<td></td>
</tr>
<tr>
<td><strong>Ethionamide</strong></td>
<td>Tab 250 mg.</td>
<td>PO 15–20 mg/kg/day (usually 500–750 mg) as a single daily dose, to a maximum of 1 g/day.</td>
<td>PO 15–20 mg/kg/day to a maximum of 1 g/day.</td>
<td>1–5 (250–500 mg)</td>
<td>3 hr</td>
<td>—</td>
</tr>
</tbody>
</table>

<sup>a</sup> Data from references. See text for details.

<sup>b</sup> Values represent normal serum levels.
<table>
<thead>
<tr>
<th>DRUG</th>
<th>DOSE FORMS</th>
<th>ADULT DOSAGE</th>
<th>PEDIATRIC DOSAGE</th>
<th>SERUM LEVELS&lt;sup&gt;b&lt;/sup&gt; (MG/L)</th>
<th>HALF-LIFE</th>
<th>MAJOR ADVERSE EFFECTS</th>
</tr>
</thead>
<tbody>
<tr>
<td>Kanamycin</td>
<td>Inj 37.5, 250, 333, 500 mg/mL</td>
<td>Same as streptomycin.</td>
<td>Same as streptomycin.</td>
<td>Same as streptomycin.</td>
<td>2–3 hr</td>
<td>80–90 hr Nephrotoxicity; ototoxicity.</td>
</tr>
<tr>
<td>Sulfate</td>
<td>Kantrex</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Rifabutin</td>
<td>Cap 150 mg.</td>
<td>PO 300 mg/day as a single dose.</td>
<td>Not well established. (1 yr) 15–25 mg/kg/day; (2–10 yr) 4–19 mg/kg/day; (14–16 yr) 2.8–5.4 mg/kg/day,</td>
<td>—</td>
<td>45 ± 16 hr</td>
<td>—</td>
</tr>
<tr>
<td>Mycobutin</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Streptomycin</td>
<td>Inj 400 mg/mL.</td>
<td>IM 12–15 mg/kg/day to a maximum of 1 g, or 22–25 mg/kg to a maximum of 1.5 g 2–3 times/week.</td>
<td>IM 20–40 mg/kg/day to a maximum of 1 g, or 25–30 mg/kg to a maximum of 1.5 g 2–3 times/week.</td>
<td>35–40 (12–15 mg/kg) 65–80 (22–25 mg/kg)</td>
<td>2–3 hr</td>
<td>↑ Vestibular ototoxicity.</td>
</tr>
<tr>
<td>Sulfate</td>
<td>Various</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

<sup>a</sup>Use only in combination with other effective antituberculars.

<sup>b</sup>Peak serum level 1 hr (parenteral) or 2 hr (oral) after the adult dose in parentheses that is evidence of adequate absorption.

<sup>c</sup>Sodium salt contains 73% aminosalicylic acid; increase dosage accordingly. Sodium content is 4.7 mEq/g.

<sup>d</sup>Peak serum level 6 hr after a dose of Paser granules.

*Adapted from references 58, 60, 63, and 66.*
**Class Instructions.** Pinworms. Purgation, enemas, or special dietary restrictions are unnecessary with this drug, which may be taken with food or beverages. To avoid reinfection with pinworms, wash the perianal area thoroughly each morning. Change and wash nightclothes, undergarments, and bedclothes daily. Wash hands and under fingernails thoroughly after bowel movements and before eating. Treat all family members simultaneously and clean bedroom and bathroom floors thoroughly at the end of the course of treatment. To demonstrate a cure, no eggs must be found in the anal area at least 5 weeks after the end of treatment.

**Pharmacology.** Albendazole is a benzimidazole drug related to mebendazole and has a similar mechanism of action; however, it has a broader range of activity than mebendazole.

**Administration and Adult Dosage.** PO for hydatid cyst 400 mg bid for 1–6 months; PO for cysticercosis 400 mg bid for 8–30 days, repeat prn; PO for Clonorchis sinensis 10 mg/kg/day for 7 days; PO for cutaneous larva migrans 400 mg/day for 3 days; PO for capillarisis 400 g/day for 10 days; PO for ascariasis, eosinophilic enterocolitis, hookworm, trichostrongylus, or trichuriasis 400 mg once; PO for pinworms 400 mg once, then repeat in 2 weeks; PO for microsporidiosis 400 mg bid (ocular infections require the addition of fumagillin); PO for trichinosis 400 mg bid for 8–14 days; PO for visceral larva migrans (toxocariasis) 400 mg bid for 5 days.\(^78\)

**Special Populations.** Pediatric Dosage. Safety and efficacy not established. PO for hydatid cyst 15 mg/kg/day to a maximum of 800 mg/day in 2 divided doses for 8–30 days, repeat prn; PO for cysticercosis 15 mg/kg/day to a maximum of 800 mg/day in 2 divided doses for 8–30 days, repeat prn; PO for Clonorchis sinensis 10 mg/kg/day for 7 days; PO for cutaneous larva migrans 400 mg/day for 3 days; PO for capillarisis 400 g/day for 10 days; PO for ascariasis, eosinophilic enterocolitis, hookworm, trichostrongylus, or trichuriasis 400 mg once; PO for eosinophilic enterocolitis 400 mg once; PO for pinworms 400 mg once, then repeat in 2 weeks; PO for trichinosis 400 mg bid for 8–14 days; PO for visceral larva migrans (toxocariasis) 400 mg bid for 5 days.\(^78\)

**Geriatric Dosage.** Same as adult dosage.

**Dosage Forms.** Tab 200 mg.

**Patient Instructions.** Take this drug with a fatty meal to increase absorption and improve effectiveness.

**Missed Doses.** Take this drug at regular intervals. If you miss a dose of this medicine, take it as soon as you remember. If it is almost time for your next dose, take that dose only and go back to your regular dosage schedule. Do not double the dose or take extra.

**Pharmacokinetics.** Fate. Absorption is poor but enhanced by fat. Oral bioavailability of unchanged albendazole is negligible because of first-pass metabolism to
albendazole sulfoxide, the active form of the drug. The sulfoxide has a peak serum level 2–3 hr after a dose. CNS concentrations are 40% of serum levels; concentration in echinococcal cysts is about 25% of serum levels. The absorbed drug is excreted primarily in urine as metabolites.\textsuperscript{79} 

\[ t_{1/2}. \text{(Albendazole sulfoxide)} 10–15 \text{ hr}. \text{\textsuperscript{79}} \]

**Adverse Reactions.** Occasionally, diarrhea, abdominal pain, and migration of roundworms through the mouth and nose occur. Rarely, leukopenia, alopecia, or increased transaminases occur.\textsuperscript{78}

**Precautions.** Pregnancy; liver dysfunction.

**Drug Interactions.** Concurrent dexamethasone increases serum levels by 50%.\textsuperscript{79}

**Parameters to Monitor.** Monitor hepatic transaminases and WBC count during prolonged therapy.

**IVERMECTIN**

**Pharmacology.** Ivermectin is a semisynthetic anthelmintic that binds to glutamate-gated chloride channels in invertebrate nerve and muscle cells, leading to increase cellular permeability, hyperpolarization of nerve cells, paralysis, and death.

**Administration and Adult Dosage.** PO for strongyloidiasis 200 \( \mu \)g/kg/day for 1–2 days; PO for onchocerciasis 150 \( \mu \)g/kg once, repeat in 3–12 months until asymptomatic; PO for Mansonella streptocerca 150 \( \mu \)g/kg once; PO for pediculosis (head or pubic lice) or scabies 200 \( \mu \)g/kg once; PO for cutaneous larva migrans 200 \( \mu \)g/kg/day for 1–2 days.\textsuperscript{78} (See Notes.)

**Special Populations.** **Pediatric Dosage.** (<15 kg) safety and efficacy not established; PO for above infestations same as adult dosage.

**Geriatric Dosage.** Same as adult dosage.

**Dosage Forms.** Tab 6 mg.

**Pharmacokinetics.** \textit{Fate.} Ivermectin is absorbed orally. It does not enter the CNS. Most of the drug is metabolized hepatically, and the drug and metabolites are excreted in the feces. Less than 1% is excreted unchanged in urine. Its half-life is about 16 hr.

**Adverse Reactions.** Fairly well tolerated, with abdominal and chest pain, dizziness, pruritus, rash, urticaria, diarrhea, nausea, and vomiting occurring frequently. In treating onchocerciasis, inflammation caused by dead and dying larvae can cause more severe and frequent cutaneous reactions, fever, lymph node swelling and tenderness, edema, and arthralgia; ocular effects include limbitis and punctate opacity.

**Precautions.** Pregnancy.

**Notes.** Ivermectin is the drug of choice for strongyloidiasis and onchocerciasis and an alternative for the other infestations listed.

**MEBENDAZOLE**

**Pharmacology.** Mebendazole is active against many intestinal roundworms. It binds to helminth tubulin and inhibits glucose uptake in the parasite, with no effect on blood glucose concentrations in the host.\textsuperscript{79}
Administration and Adult Dosage. PO for pinworms 100 mg once, repeat in 2 weeks; PO for ascariasis or hookworms 100 mg bid for 3 days or 500 mg once; PO for capillariasis 200 mg bid for 20 days; PO for eosinophilic enterocolitis 100 mg bid for 3 days; PO for roundworms, or whipworms 100 mg bid for 3 days.78

Special Populations. Pediatric Dosage. (<2 yr) Safety and efficacy not established. PO for pinworms, ascariasis, capillariasis, roundworms, whipworms, eosinophilic enterocolitis, or hookworms same as adult dosage.78

Geriatric Dosage. Same as adult dosage.

Dosage Forms. Chew Tab 100 mg.

Patient Instructions. (See Pinworms Class Instructions.) Chew tablets before swallowing.

Pharmacokinetics. Fate. Poorly absorbed orally. Almost all eliminated unchanged in the feces, but up to 10% can be recovered in the urine 48 hr after a dose, primarily as the decarboxylated metabolite.79

Adverse Reactions. Occasional abdominal pain and diarrhea in cases of massive infestation and expulsion of worms. Occasionally, migration of roundworms through the mouth and nose occurs. Rarely, leukopenia, agranulocytosis, and hypospermia have been reported.78

Precautions. Pregnancy.

Drug Interactions. Carbamazepine and hydantoins can reduce mebendazole serum levels.

Parameters to Monitor. When treating whipworm, take a stool sample for egg count 3 weeks after treatment to detect frequent (about 30%) persistent infestation requiring retreatment.

Notes. Mebendazole is the agent of choice for whipworm, producing about a 70% cure rate with a single treatment; the cure rate is 90–100% with roundworms, hookworms, and pinworms. Particularly useful in mixed infestations.78,79

PERMETHRIN Acticin, Elimite, Nix, Various

Pharmacology. Permethrin is a pyrethroid that acts on arthropod nerve cell membranes to cause delayed polarization and paralysis. It is active against lice (including unhatched eggs) and mites (eg, scabies).

Administration and Adult Dosage. Top for head lice apply 1% cream rinse to hair one time after washing hair. Leave on for no longer than 10 min and rinse with water. If live lice are seen after >7 days, reapply as above. Top for pubic lice although not FDA-approved, use of topical 5% permethrin has been used (see also Ivermectin);78 repeat application at 10 days has been recommended.80,81 Top for scabies thoroughly massage 5% cream into the skin from the head to the soles of the feet. Remove by showering or bathing after 8–14 hr.

Special Populations. Pediatric Dosage. Top for lice (<2 months) safety and efficacy not established; (>2 months) same as adult dosage. Top for scabies (<2
months) safety and efficacy not established. Neonates have been treated, but re-
move cream after 6 hr;82 (>2 months) same as adult dosage.

**Geriatric Dosage.** Same as adult dosage.

**Dosage Forms.** Liquid (creme rinse) 1% (Nix, various); Crm 5% (Acticin, Elimite).

**Patient Instructions.** Lice. Wash hair and towel dry. Apply enough creme rinse
to saturate hair and scalp, especially behind the ears and on the nape of the neck.
Use the comb provided with the product to remove nits. Wash all pillow cases, pa-
jamas, and towels in hot, soapy water and dry using the hot cycle of a dryer for at
least 20 minutes. Clothing and bedding that cannot be washed should be sealed in
a plastic bag for 2 weeks or dry cleaned. Soak combs in hot water for 5–10 min-
utes. If infestation of the eyebrows or eyelashes occurs, consult your health care
provider. Scabies. Itching, mild burning, or stinging can occur after application.
Itching usually resolves by 4 weeks. If irritation persists, consult your health care
provider.

**Pharmacokinetics.** **Fate.** Usually <2% absorbed after topical application. Per-
methrin is rapidly metabolized by ester hydrolysis to inactive metabolites which
are excreted in urine.83

**Adverse Reactions.** Adverse reactions are mild and occur only occasionally with
the treatment of lice. With the 5% cream for treatment of scabies, mild, transient
burning, stinging, or tingling occurs in about 10% of patients. Itching, edema, and
erythema are often symptoms of scabies and can be exacerbated temporarily by
treatment with permethrin. Intolerable burning and stinging can occur in patients
with AIDS and scabies.83 Itching and skin irritation can persist after successful
treatment because of local allergic reactions to the dead mites. Allergic reactions
are rare and might be caused by the formaldehyde preservative.

**Contraindications.** Documented allergy to any pyrethroid or vehicle component.

**Precautions.** Pregnancy, although animal studies indicate no teratogenicity. Dur-
ing lactation, discontinue nursing temporarily during treatment with 5% cream;
use of 1% creme rinse poses little risk during breastfeeding. Avoid contact with
eyes and mucous membranes.

**Drug Interactions.** None known.

**Parameters to Monitor.** Observe for parasites 7–10 days after treatment of lice
infestation or 14 days after treatment of scabies.

**Notes.** Permethrin is the drug of choice for pediculosis and scabies. Malathion
0.5% lotion (Ovide) is an alternative for pediculosis. Synergized pyrethrins
(pyrethrins and piperonyl butoxide) have efficacies similar to that of permethrin
for head lice but are not as persistent as permethrin and require a repeat treatment
after 1 week.78 In scabies, permethrin is safer and more effective than lindane and
more effective and easier to use than crotamiton.81,83 Lindane is not recom-
}{
**PRAZIQUANTEL**

**Pharmacology.** Praziquantel causes a loss of intracellular calcium, resulting in paralysis and dislodgement of worms from sites of attachment. In higher dosages, it damages the parasite’s surface membrane, allowing the host’s immune response to destroy the worm.84

**Administration and Adult Dosage.** **PO for schistosomiasis** (*Schistosoma haematobium* or *mansoni*) 40 mg/kg in 2 divided doses the same day, but heavy infestations require 60 mg/kg in 3 divided doses at 4–6 hr intervals;78,84 (*Schistosoma japonicum* or *mekongi*) 60 mg/kg in 3 doses at 4–6 hr intervals. **PO for flukes** (eg, clonorchiasis, opisthorchiasis) 25 mg/kg tid for 1 day; (paragonimiasis) 25 mg/kg tid for 2 days.78,84 **PO for tapeworms** (beef, dog, fish, pork) 5–10 mg/kg once; (dwarf tapeworm) 25 mg/kg once. **PO for neurocysticercosis** 50–100 mg/kg/day in 3 doses for 30 days.78 (See Notes.)

**Special Populations.** **Pediatric Dosage.** (<4 yr) safety and efficacy not established. **PO for above infestations** same as adult dosage.78

**Geriatric Dosage.** Same as adult dosage.

**Dosage Forms.** Tab 600 mg.

**Patient Instructions.** Take with liquid during meals but do not chew tablets. This drug can cause dizziness or drowsiness. Use caution when driving, operating machinery, or performing other tasks requiring mental alertness.

**Pharmacokinetics.** **Fate.** The drug is 80% absorbed orally, but undergoes extensive first-pass metabolism. CSF concentrations are 14–20% of serum levels. The drug is metabolized and metabolites are excreted primarily in urine. $t_1/2$ 1.1 ± 0.3 hr.

**Adverse Reactions.** Side effects are usually mild. Dizziness, headache, and malaise occur frequently after large doses. Occasionally, abdominal discomfort, fever, sweating, and eosinophilia occur. Drowsiness or fatigue might occur because of a structural similarity to benzodiazepines. Pruritus and rash occur rarely.78 In patients treated for cysticercosis, an inflammatory response, presumably caused by dead and dying organisms, occurs that is manifested by headache, seizures, and increased intracranial pressure.

**Contraindications.** Ocular cysticercosis.

**Precautions.** Pregnancy; liver disease; avoid breastfeeding for 72 hr after the last dose.

**Drug Interactions.** Drugs that induce CYP3A4 (eg, dexamethasone, carbamazepine, phenobarbital, phenytoin) can increase clearance, decrease bioavailability, and cause treatment failure; drugs that inhibit CYP3A4 (eg, cimetidine, ketoconazole, erythromycin) decrease clearance, increase serum levels, and lengthen half-life.79,84

**Parameters to Monitor.** Observe for CNS toxicity when treating cysticercosis.

**Notes.** Concomitant corticosteroid therapy is recommended for patients treated for neurocysticercosis.
Pharmacology. Pyrantel is a depolarizing neuromuscular blocker that produces spastic paralysis of the parasite with no similar effects on the host after oral use. It also inhibits acetylcholinesterases.79

Administration and Adult Dosage. PO for pinworms and roundworms 11 mg/kg, to a maximum of 1 g in a single dose; for pinworms repeat in 2 weeks; PO for moniliformis 11 mg/kg 3 times at 2-week intervals; PO for hookworms and eosinophilic enterocolitis 11 mg/kg/day, to a maximum of 1 g for 3 days.78

Doses are expressed as base equivalent.

Special Populations. Pediatric Dosages. (<2 yr) Safety and efficacy not established. PO for above infestations same as adult dosage.78

Geriatric Dosage. Same as adult dosage.

Dosage Forms. Cap 180 mg (62.5 mg as pyrantel base); Liq 50 mg/mL (as pamoate base); Susp 50 mg/mL (as pamoate base).

Patient Instructions. (See Pinworms Class Instructions.)

Pharmacokinetics. Fate. Slight oral absorption. Over 50% is excreted unchanged in feces, and less than 15% of the dose is excreted as parent drug and metabolites in the urine.79

Adverse Reactions. Occasional nausea, vomiting, headaches, dizziness, rash, and transient AST elevations.78

Contraindications. Liver disease.

Precautions. Avoid during pregnancy.

Drug Interactions. Piperazine and pyrantel might be mutually antagonistic in ascariasis.

Notes. Except for pinworms, for which it is virtually 100% effective, and moniliformis, pyrantel is an alternative to other drugs.78

Antiviral Drugs

Class Instructions: HIV Drugs. Underdosage, noncompliance, or partial compliance with drug regimens for these drugs might result in development of a resistant strain(s) of HIV that will not be susceptible to treatment. Do not stop taking this medication unless told to do so by your health care provider. This drug should be used in combination with other anti-HIV medications. Protease inhibitors do not cure or prevent HIV infection. It is possible for a person taking this medication to transmit the virus to another person. Opportunistic infections and other complications associated with HIV infection can continue to develop while you take this medication. Protease inhibitors and nonnucleoside reverse transcriptase inhibitors have a potential for serious interactions with a large number of commonly prescribed drug products. Always check with your health care provider before starting any new medication.

Missed Doses. Missing doses can result in the development of resistance that can lead to treatment failure. If you forget a dose, take it as soon as you remember. If
it is almost time for your next scheduled dose (within 4 hours), skip the missed
dose. Do not double your dose.

ACYCLOVIR

Pharmacology. Acyclovir is an acyclic nucleoside analogue of deoxyguanosine
that is selectively phosphorylated by the virus-encoded thymidine kinase to its
monophosphate form. Cellular enzymes then convert the monophosphate to the
active antiviral acyclovir triphosphate, which inhibits viral DNA synthesis by in-
corporation into viral DNA, resulting in chain termination. Acyclovir has potent
activity against herpes simplex virus (HSV) I and II and herpes zoster virus
(varicella-zoster virus [VZV]). Activity against cytomegalovirus, which lacks a
specific virus-encoded thymidine kinase, is limited, but resistance can be overcome
with high serum concentrations in some patient populations. Acyclovir inhibits
Epstein-Barr virus but has not been found clinically useful. Human herpes virus 6
is resistant. Valacyclovir is the L-valyl ester of acyclovir, which undergoes exten-
sive first-pass hydrolysis to yield high serum acyclovir concentrations.

Administration and Adult Dosage. IV for severe localized HSV infection (acyc-
lovir) 5 mg/kg q 8 hr for 5 days for nonimmunocompromised patients or 7–
10 days for immunocompromised patients; IV for VZV (chickenpox) infection in
immunocompromised patients 10 mg/kg q 8 hr for 7–10 days; IV for HSV en-
cephalitis 10 mg/kg q 8 hr for 10–14 days. Dilute to 50–250 mL and infuse over
at least 60 min; avoid bolus IV, SC, or IM injections. Maintain minimum urine
output of 500 mL/24 hr for each gram of acyclovir administered. PO for primary
or recurrent genital HSV infection (acyclovir) 200 mg 5 times/day for 10 days;
(valacyclovir, immunocompetent patients) 500 mg bid for 5 days. PO for preven-
tion of recurrent genital HSV infection (acyclovir) 400 mg bid or 200 mg 3–
5 times/day; (valacyclovir, immunocompetent patients) 1 g/day or 500 mg bid;
PO for active VZV (chickenpox) or herpes zoster (acyclovir) 800 mg q 4 hr
5 times/day for 5 days (chickenpox) or 7–10 days (zoster). PO for herpes zoster
in immunocompetent patients (valacyclovir) 1 g q 8 hr for 7 days. Top for ini-
tial genital HSV infection and limited non–life-threatening mucocutaneous
HSV infections in immunocompromised patients (acyclovir) 0.5-inch ribbon to
cover 4-square-inch affected skin area q 3 hr 6 times/day for 7 days.

Special Populations. Pediatric Dosage. (All dosages apply to acyclovir.) IV for
HSV infection (neonates) 10 mg/kg q 8 hr for 10–14 days; (13 months–11 yr)
750 mg/m²/day in 3 divided doses for 7 days. IV for VZV (chickenpox) in
immunocompromised children (13 months–11 yr) 1500 mg/m²/day in 3 divided
doses. IV for HSV encephalitis (6 months–11 yr) 1500 mg/m²/day in 3 divided
doses for 10 days. Infuse over at least 60 min; avoid bolus IV, SC, or IM injec-
tions. PO for VZV (chickenpox) (acyclovir) (>2 yr and <40 kg) 20 mg/kg/dose,
to a maximum of 800 mg q 6 hr for 5 days; (>40 kg) same as adult dosage; (vala-
cylovir) safety and efficacy not established.

Geriatric Dosage. Same as adult dosage, adjusted for renal function.
**Other Conditions.** (Acyclovir) in obesity, base dosage on IBW. In renal insufficiency, reduce parenteral and oral dosage: (Clcr 25–50 mL/min) usual dose q 12 hr; (Clcr 10–25 mL/min) usual dose q 24 hr; (Clcr 0–10 mL/min) 50% of the usual dose q 24 hr. For patients on hemodialysis, give the usual daily dosage after dialysis. (Valacyclovir) in renal insufficiency, reduce the dosage: (Clcr 30–49 mL/min) 1 g q 12 hr; (Clcr 10–29 mL/min) 1 g q 24 hr; (Clcr <10 mL/min) 500 mg q 24 hr.

**Dosage Forms.** (Acyclovir) Cap 200; Tab 400, 800 mg; Inj 500 mg, 1 g; Inj 50 mg/mL; Oint 5%; Susp 40 mg/mL. (Valacyclovir) Tab 500 mg.

**Patient Instructions.** Use a finger cot or latex glove when applying acyclovir ointment. The ointment might cause transient burning or stinging.

**Missed Doses.** Take this (oral) drug at regular intervals. If you miss a dose of this medicine, take it as soon as you remember. If it is almost time for your next dose, take that dose only and go back to your regular dosage schedule. Leave at least 4 hours between doses. Do not double the dose or take extra.

**Pharmacokinetics. Fate.** Oral bioavailability of acyclovir is estimated to be 15–30%; valacyclovir is well absorbed, with a bioavailability of 54%. Valacyclovir is extensively converted to acyclovir after oral administration. After 200–600 mg of acyclovir orally, mean peak steady-state levels are 0.56–1.3 mg/L (2.5–5.9 μmol/L); levels after IV doses of 2.5–15 mg/kg are 5–24 mg/L (23–105 μmol/L). After an oral dose of 1 g of valacyclovir, mean peak steady-state acyclovir level is 5–6 mg/L (22–27 μmol/L). CSF acyclovir concentrations are 25–70% of simultaneous serum level. Decay is biphasic, with $V_d$ of 0.69 ± 0.19 L/kg; $Cl$ is 0.21 ± 0.03 L/hr/kg with normal renal function; 86–92% is excreted unchanged in urine; the remainder is metabolized to 9-carboxymethoxymethylguanine. Renal clearance is 75–80% of total clearance and is markedly reduced by concomitant probenecid.

$t_\alpha$. (Acyclovir) $\alpha$ phase 0.34 hr, $\beta$ phase 2.9 ± 0.8 hr in adult patients, increasing to nearly 20 hr in end-stage renal disease; 5.7 hr on dialysis; about 4 hr in neonates.

**Adverse Reactions.** (Acyclovir) nephrotoxicity, thought to be caused by precipitation of acyclovir crystals in the nephron, occurs in about 10% of patients if the drug is given by bolus (<10 min) injection. Phlebitis at injection site occurs frequently with IV infusion because of the high pH (9–11) of the product. Other reported side effects are CNS toxicity (eg, headache, lethargy, tremulousness, delirium, seizures), nausea, vomiting, and skin rash. CNS toxicity occurs primarily in patients with underlying neurologic disease or end-stage renal disease, or with cancer chemotherapy and irradiation to the CNS, and might not be primarily caused by the drug. Topical application to herpes lesions can be painful. (Valacyclovir) adverse reactions appear comparable to acyclovir. Nausea, vomiting, diarrhea, abdominal pain, and headache have been reported frequently with valacyclovir use. Thrombotic thrombocytopenic purpura/hemolytic-uremic syndrome has been reported in patients with advanced HIV disease and in bone marrow and renal transplant patients. This phenomenon has not been reported in immunocompetent patients.

**Contraindications.** (Valacyclovir) allergy to the drug or to acyclovir.
Precautions. Use caution in renal impairment, dehydration, or pre-existing neurologic disorders. Valacyclovir not indicated in immunocompromised patients.

Drug Interactions. Zidovudine and acyclovir can result in drowsiness and lethargy. Probenecid can increase oral bioavailability and half-life of acyclovir.

Parameters to Monitor. Monitor renal function and injection site for signs of phlebitis daily. Carefully monitor patients with underlying neurologic diseases for evidence of neurotoxicity. (See Adverse Reactions.)

Notes. Acyclovir-resistant strains of virus that are deficient in thymidine kinase have been isolated from patients after treatment. Although thought to be less virulent than sensitive strains, HSV strains resistant to acyclovir have been described in AIDS patients.86

Pharmacology. Cidofovir (HPMPC) is a nucleotide analogue with potent in vitro and in vivo activities against cytomegalovirus (CMV) and other herpes viruses. Cidofovir contains a phosphonate group that enables it to bypass initial virus-dependent phosphorylation. Cellular enzymes convert cidofovir to cidofovir diphosphate, the active intracellular metabolite.92–94

Adult Dosage. IV induction for CMV retinitis 5 mg/kg infused over 1 hr once weekly for 2 weeks, then IV maintenance 5 mg/kg once every other week. Reduce dosage to 3 mg/kg if Crs increases by 0.3–0.4 mg/dL above baseline or if >2+ proteinuria occurs. It is essential to give the following with cidofovir: 2 g probenecid PO 3 hr before administration and 1 L of NS IV over 1 hr just before administration. If tolerated, give another liter of NS with or after cidofovir administration; finally, give PO 1 g probenecid 2 hr and 8 hr after the end of cidofovir infusion. Cidofovir is also being investigated as an intravitreal injection for CMV retinitis.

Dosage Forms. Inj 75 mg/mL.

Pharmacokinetics. Peak serum cidofovir concentration averages 26.1 ± 3.2 mg/L (83 ± 10 µmol/L) after a 5 mg/kg IV infusion with concomitant probenecid and hydration. Cidofovir is not appreciably bound to plasma proteins; Vd averages 0.5 L/kg. Cidofovir is excreted almost entirely unchanged in the urine. The elimination half-life of cidofovir is 3–6 hr when administered with probenecid. Cidofovir diphosphate has a prolonged intracellular half-life, with a range of 17–65 hr, which allows infrequent administration schedules of once weekly to once every other week.

Adverse Reactions. Nephrotoxicity is the most frequent adverse reaction, and high-dose probenecid (see Adult Dosage) must be used with administration of cidofovir. Probenecid decreases uptake of cidofovir in proximal renal tubular cells, decreasing the risk of nephrotoxicity. Other frequent adverse reactions are proteinuria, elevated Crs, nausea, vomiting, fever, asthenia, neutropenia, rash, headache, diarrhea, alopecia, anemia, and abdominal pain. Ocular hypotony and decreased intraocular pressure have been reported occasionally. Nausea, vomiting, fever, rash, and chills are frequent reactions reported with probenecid.
Pharmacology. Didanosine (dideoxyinosine [ddI]) is a purine nucleoside that undergoes complex metabolism in vivo to dideoxyadenosine (ddA), which ultimately undergoes metabolism to an active triphosphorylated form (ddATP). Incorporation of ddATP into viral DNA leads to chain termination, and ddATP is a competitive inhibitor of HIV reverse transcriptase, which further contributes to the interference of HIV replication.95,96

Administration and Adult Dosage. PO for HIV infection (tablets or solution) (≥60 kg) 200 mg (as 2 tablets) q 12 hr, or 400 mg/day (as 2 tablets), or 250 mg (as powder) q 12 hr; (<60 kg) 125 mg (as 2 tablets) q 12 hr, or 250 mg/day (as 2 tablets), or 167 mg (as powder) q 12 hr. Take each dose as 2 whole (not partial) tablets to provide adequate buffering. PO for HIV infection (EC capsules) same dosage as above, but as a single daily dose.

Special Populations. Pediatric Dosage. PO for HIV infection 120 mg/m² bid.

Geriatric Dosage. Same as adult dosage, but not studied in this population.

Other Conditions. Consider dosage reduction in patients with renal or hepatic impairment. (Tablets or solution) Clcr 30–59 mL/min: (≥60 kg) 200 mg/day (as 2 tablets or EC capsules) or 100 mg (as 2 tablets) bid; (<60 kg) 150 mg/day (as 2 tablets), 75 mg (as 2 tablets) bid or 125 mg /day (as EC capsules). Clcr 10–29 mL/min: (≥60 kg) 150 mg/day (as 2 tablets) or 125 mg/day (as EC capsules); (<60 kg) 100 mg/day (as 2 tablets). Clcr <10 mL/min: (≥60 kg) 100 mg/day (as 2 tablets); (<60 kg) 75 mg/day (as 2 tablets)—EC capsules not recommended. Didanosine is removed by hemodialysis, but the quantity removed is low and supplemental doses are not recommended.97

Dosage Forms. Chew/Dispersible Tab 25, 50, 100, 150, 200 mg; EC Cap 125, 200, 250, 400 mg; Pwdr for Oral Soln 100, 167, 250 mg; Pwdr for Oral Soln (pediatric) 2, 4 g.

Patient Instructions. (See HIV Drugs Class Instructions.) Didanosine must be taken on an empty stomach 1 hour before or 2 hours after a meal. It is essential that the 2-tablet dose be taken each time to avoid destruction of the drug by stomach acid. For children >1 year, use the 2-tablet dose; for those <1 year of age, use the 1-tablet dose. Tablets may be chewed and swallowed or dissolved in at least 30 mL of water and swallowed immediately. Do not swallow the tablets whole. Reconstituted solution may be stored for up to 30 days when refrigerated. Shake solution thoroughly before administering each dose. Do not crush or chew EC capsule.

Missed Doses. Take this drug at regular intervals. If you miss a dose of this medicine, take it as soon as you remember. If it is almost time for your next dose, take that dose only and go back to your regular dosage schedule. Leave at least 12 hours between doses. Do not double the dose or take extra.

Pharmacokinetics. Fate. Didanosine is rapidly degraded at acidic pH. Appreciable interpatient variability and dose-dependent characteristics affect didanosine absorption. Oral bioavailability of the buffered powder for oral solution is 33 ± 11%.98,99 The chewable/dispersible buffered tablets are 20–25% more bioavailable than the buffered powder for solution. The peak serum concentration is 1.1 ±
0.7 mg/L (4.7 ± 2.9 μmol/L) after a 375 mg oral dose of buffered powder for solution. Protein binding is less than 5%. CSF concentration 1 hr after infusion of didanosine averages 21% of the concurrent serum concentration. V_dβ is 1 ± 0.7 L/kg; Cl is 1 ± 0.08 L/hr/kg. Up to 60% of dose is excreted unchanged in the urine; the remainder is extensively metabolized to ddATP, hypoxanthine, and uric acid.97,100

\[ t_{1/2} = 1.75 ± 0.99 \text{ hr}; \]

**Adverse Reactions.** Pancreatitis has occurred at a frequency of 5–9% in clinical trials at or below current recommended dosages and can be fatal. Peripheral neuropathy occurs in 16–34% of patients, with 12% requiring dosage reduction. Diarrhea has been reported with the buffered powder for oral solution at a frequency of 34%. In children, pancreatitis and peripheral retinal depigmentation have occurred frequently, although the latter has not been associated with visual impairment.102 Peripheral neuropathy has not occurred in children.

**Precautions.** Avoid didanosine tablets in patients with phenylketonuria because these contain phenylalanine. Didanosine has been associated with hyperuricemia; use caution in patients with a history of gout or baseline hyperuricemia; avoid in individuals with a history of pancreatitis.

**Drug Interactions.** Administration with fluoroquinolones can reduce fluoroquinolone serum levels because of buffers in formulation. Avoid concurrent administration with dapsone, indinavir, itraconazole, ketoconazole, or other medications requiring an acidic environment for absorption because of buffers in didanosine formulation. Ganciclovir and trimethoprim-sulfamethoxazole appear to increase didanosine’s bioavailability, but the clinical importance is unknown. Use with alcohol, high-dose trimethoprim-sulfamethoxazole, or other pancreatitis-associated drugs can increase the risk of pancreatitis.103,104

**Parameters to Monitor.** Obtain serum amylase, lipase, and triglycerides monthly. Symptoms of abdominal pain, nausea, and vomiting can indicate pancreatitis. Symptoms of distal numbness, tingling, or pain in the feet or hands can indicate neuropathy and might necessitate dosage modification. Monitor clinical signs, symptoms, and laboratory markers for progression of HIV disease to help decide regimen changes in antiretroviral therapy. Baseline CD4 and HIV-1 RNA polymerase chain reaction viral load tests are useful to measure clinical benefit of therapy. Repeat tests after 1 month and q 3–4 months thereafter have been suggested to monitor benefit of antiretroviral therapy.

**Notes.** As with other nucleoside reverse transcriptase inhibitors, drug-resistant HIV-1 isolates emerge with long-term didanosine therapy (≥12 months).105 (See Antiviral Drugs for HIV Infection Comparison Chart.)

**FAMCICLOVIR**

**PENCICLOVIR**

**Pharmacology.** Famiclovir is the diacetyl, 6-deoxy ester of the antiviral guanosine analogue penciclovir. Famiclovir is absorbed rapidly and converted to penciclovir in the intestinal wall and liver. Viral thymidine kinase converts penci-
clovir to its monophosphate form. Cellular enzymes then convert the monophosphate to the active antiviral penciclovir triphosphate. The triphosphate inhibits viral DNA synthesis by incorporation into viral DNA, resulting in termination of the chain. Penciclovir has potent activity against HSV I and II and herpes zoster virus (varicella-zoster). Penciclovir also has some activity against Epstein-Barr virus and CMV but has not demonstrated clinical usefulness against infections with these agents.106–109

**Adult Dosage.** PO for herpes zoster (famciclovir) 500 mg q 8 hr for 7 days. In renal insufficiency, reduce the dosage as follows: Clcr 40–59 mL/min, 500 mg q 12 hr; Clcr 20–39 mL/min, 500 mg q 24 hr; Clcr <20 mL/min, 250 mg q 48 hr; with hemodialysis, 250 mg after each dialysis. PO for recurrent genital HSV infection (famciclovir) 125 mg bid for 5 days. In renal insufficiency, reduce the dosage as follows: Clcr 20–39 mL/min, 125 mg q 24 hr; Clcr <20 mL/min, 125 mg q 48 hr; with hemodialysis, 125 mg after each dialysis. Top for herpes labialis (penciclovir) apply to lesions q 2 hr while awake for 4 days, starting as early as possible at the beginning of an outbreak.

**Dosage Forms.** Crm (penciclovir) 1%; Tab (famciclovir) 125, 250, 500 mg.

**Pharmacokinetics.** Topical penciclovir is virtually unabsorbed. The absolute bioavailability of penciclovir is 77% after a 500 mg oral dose of famciclovir. Peak serum concentrations are 0.84 ± 0.22 (3.3 ± 0.9 μmol/L) and 3.34 ± 0.58 mg/L (13 ± 2.3 μmol/L) 45 min after 125 and 500 mg oral doses of famciclovir, respectively. Penciclovir is <20% protein bound, and the Vd is approximately 1 L/kg. Penciclovir is eliminated primarily by renal excretion. The elimination half-life is approximately 2 hr with normal renal function, increasing to over 13 hr in patients with severely impaired renal function.

**Adverse Reactions.** Nausea, vomiting, diarrhea, and headache occur frequently with famciclovir. Pruritus, paresthesias, and fatigue occur occasionally. Penciclovir causes mild erythema occasionally.

**Drug Interactions.** Cimetidine might enhance the bioavailability of famciclovir and its conversion to penciclovir.

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**FOSCARNET SODIUM**

**Pharmacology.** Foscarnet sodium (phosphonoformic acid [PFA]) is a pyrophosphate analogue. Foscarnet actively inhibits viral DNA polymerases in its parent form and does not require phosphorylation for optimal antiviral activity. It has antiviral activity against HSV I and II, human CMV, Epstein-Barr virus, hepatitis B virus, varicella-zoster virus, and some retroviruses including HIV. Foscarnet sodium inhibits DNA synthesis in CMV and other herpes viruses by inhibiting viral DNA polymerase.110

**Administration and Adult Dosage.** IV induction for CMV retinitis in AIDS patients 60 mg/kg q 8 hr or 90 mg/kg q 12 hr for 14–21 days.111 IV maintenance for CMV retinitis in AIDS patients 90–120 mg/kg/day in 1 dose. IV for acyclovir-resistant herpes virus infections 40 mg/kg q 8 hr or 60 mg/kg q 12 hr until clinical resolution.111 IV for acyclovir-resistant varicella-zoster infections
in immunocompromised patients 40 mg/kg q 8 hr or 60 mg/kg q 12 hr for 10–21 days or until clinical resolution.87,112

**Special Populations. Pediatric Dosage.** Safety and efficacy not established.

**Geriatric Dosage.** Same as adult dosage but adjusted for renal function.

**Other Conditions.** Reduce dosage in renal impairment. (See product information.)

**Dosage Forms.** Inj 24 mg/mL.

**Patient Instructions.** Foscarnet is not a cure for CMV retinitis, and progression of disease might continue during or after treatment. Regular eye examinations are important to monitor for disease progression. Report symptoms of tingling around the mouth or numbness in extremities, which might indicate a need for temporary discontinuation of foscarnet.

**Missed Doses.** Take this drug at regular intervals. If you miss a dose of this medicine, take it as soon as you remember. If it is almost time for your next dose, take that dose only and go back to your regular dosage schedule. Leave at least 12 hours between doses. Do not double the dose or take extra.

**Pharmacokinetics. Fate.** After twice-daily infusion of 90 mg/kg over 2 hr, peak serum levels are 98 ± 27 mg/L (577 ± 161 μmol/L) and troughs are 6.4 ± 8.3 mg/L (38 ± 49 μmol/L).111 Plasma protein binding is 14–17%. CSF concentrations are 35–103% of simultaneous serum levels. V_{ds} is 0.3–0.7 L/kg; Cl is 0.13 ± 0.05 L/hr/kg. Foscarnet is not metabolized and is 70–90% excreted unchanged in the urine.111

\[ t_{1/2} = 1.4 ± 0.6 \text{ hr}, \beta \text{ phase } 6.8 ± 5 \text{ hr in patients receiving continuous or intermittent infusions. A terminal half-life of 36–196 hr might represent release of the drug from binding sites in bone.}^{111} \]

**Adverse Reactions.** Abnormal renal function, including decreased Cl_{cre} and acute renal failure, occurs in about one-third of patients. Electrolyte abnormalities such as hypocalcemia, hypophosphatemia, hyperphosphatemia, hypokalemia, and hypomagnesemia occur in 6–16% of patients. Seizures have been reported in 10% of patients and might be related to electrolyte abnormalities or underlying disease. Other adverse reactions frequently reported are fever 65%, nausea 47%, anemia 33%, diarrhea 30%, vomiting 26%, headache 26%, and granulocytopenia 14%. Local irritation, inflammation, and pain might occur at the injection site with peripheral administration at a frequency of 1–5%.111,113,114

**Precautions.** Use with extreme caution in patients with renal impairment of nephrotoxic drugs, pre-existing cytopenias, pre-existing electrolyte abnormalities, or underlying neurologic disorders.

**Drug Interactions.** Concurrent use of nephrotoxic drugs such as aminoglycosides or radiologic contrast media can increase risk and severity of nephrotoxicity. IV pentamidine can increase the risk of hypocalcemia; avoid this combination, if possible, although inhaled pentamidine does not seem to be a risk factor.104

**Parameters to Monitor.** Monitor Cl_{cre} 2 or 3 times/week during induction therapy and weekly during maintenance therapy. Monitor serum calcium, magnesium, potassium, and phosphorus at the same frequency as Cl_{cre}. Symptoms of perioral tingling, numbness in extremities, or other paresthesias might indicate electrolyte
abnormalities and require more frequent monitoring and a need to obtain ionized calcium levels.

GANCICLOVIR  
Valganciclovir

Pharmacology. Ganciclovir (DHPG) is a synthetic acyclic nucleoside analogue of guanine. Antiviral activity is a result of its conversion to the triphosphate form, which functions as an inhibitor of and faulty substrate for viral DNA polymerase. Ganciclovir has antiviral activity against HSV I and II, human CMV, Epstein-Barr virus, and varicella-zoster virus. Valganciclovir is the valine ester prodrug that is hydrolyzed to ganciclovir after oral administration.

Administration and Adult Dosage. Take all oral doses with food. IV for CMV retinitis (induction) 5 mg/kg q 12 hr for 14–21 days, then (maintenance) 5 mg/kg once daily for 7 days/week or 6 mg/kg once daily for 5 days/week. PO for CMV retinitis (induction) (valganciclovir) 900 mg q 12 hr for 21 days, then (maintenance) 900 mg once daily. Induction may be repeated for patients who experience disease progression. IV for prevention of CMV disease in transplant recipients 5 mg/kg q 12 hr for 7–14 days, followed by 5 mg/kg once daily for 7 days/week or 6 mg/kg once daily for 5 days/week. Duration depends on duration and degree of immunosuppression. Dilute IV dose in 100 mL NS or D5W and infuse over 60 min. PO for CMV retinitis (maintenance after IV induction) (ganciclovir) 1 g q 8 hr or (valganciclovir) 900 mg once daily. PO for prophylaxis of CMV disease (ganciclovir) 1 g q 8 hr indefinitely; (valganciclovir) 900 mg once daily.

Special Populations. Pediatric Dosage. The adult dosage in mg/kg has been used.

Geriatric Dosage. Same as adult dosage adjusted for renal function.

Other Conditions. In renal insufficiency (Ganciclovir). Parenteral induction: (Cl cr 50–69 mL/min) 2.5 mg/kg q 12 hr; (Cl cr 25–49 mL/min) 2.5 mg/kg q 24 hr; (Cl cr <25 mL/min) 1.25 mg/kg q 24 hr; (hemodialysis) 1.25 mg/kg 3 times/week. On hemodialysis days, give dose after hemodialysis. Parenteral maintenance: (Cl cr 50–69 mL/min) 2.5 mg/kg q 24 hr; (Cl cr 25–49 mL/min) 1.25 mg/kg q 24 hr; (Cl cr 10–24 mL/min) 0.625 mg/kg q 24 hr; (hemodialysis) 0.625 mg/kg 3 times/week after hemodialysis. Oral maintenance: (Cl cr 50–69 mL/min) 1.5 g once daily or 500 mg tid; (Cl cr 25–49 mL/min) 1 g/day in 1 or 2 doses; (Cl cr 10–24 mL/min) 500 mg/day; (Cl cr <10 mL/min) 500 mg 3 times/week after hemodialysis. (Valganciclovir) Oral induction: (Cl cr 40–59 mL/min) 450 mg q 12 hr; (Cl cr 25–39 mL/min) 450 mg/day; (Cl cr 10–24 mL/min) 450 mg q 48 hr; (hemodialysis) use ganciclovir. Maintenance: (Cl cr 40–59 mL/min) 450 mg/day; (Cl cr 25–39 mL/min) 450 mg q 48 hr; (Cl cr 10–24 mL/min) 450 mg twice weekly.

Dosage Forms. (Ganciclovir) Cap 250, 500 mg; Inj 500 mg; Ocular Implant 4.5 mg (nominal release). (Valganciclovir) Tab 450 mg.

Patient Instructions. This drug is not a cure for CMV retinitis, and progression might continue during or after treatment. Concurrent use with zidovudine can result in severe reduction in white blood cell count; therefore, report any signs or symptoms of infection, such as fever, chills, or sweats. Take oral ganciclovir or valganciclovir with food.
Missed Doses. Take this drug at regular intervals. If you miss a dose of this medicine, take it as soon as you remember. If it is almost time for your next dose, take that dose only and go back to your regular dosage schedule. Leave at least 4 hours between doses. Do not double the dose or take extra.

Pharmacokinetics. Fate. Ganciclovir is absorbed poorly from the GI tract; oral bioavailability is 6% when taken with food (about 20% greater than when taken on an empty stomach). Average peak serum concentration of 0.34 ± 0.13 mg/L (1.3 ± 0.5 μmol/L) occurs 1–2 hr after a single 1 g oral dose. Valganciclovir bioavailability is 61%. Mean peak and trough steady-state levels after IV doses of 5 mg/kg q 12 hr in patients with normal renal function are 5.3 ± 2.8 mg/L (21 ± 11 μmol/L) and 1.1 ± 0.4 mg/L (4.3 ± 1.5 μmol/L), respectively. Ganciclovir is 1–2% plasma protein bound; CSF concentration is 24–67% of simultaneous serum level. Vc is 0.26 ± 0.08 L/kg; Vd is 1.17 ± 0.54 L/kg; Cl is 0.25 ± 0.13 L/hr/kg with normal renal function. The drug is 90–99% excreted unchanged in the urine. Hemodialysis reduces serum levels by 53 ± 12%. Renal excretion occurs principally via glomerular filtration, although limited renal tubular secretion also can occur.114,115

t1/2α phase 0.76 ± 0.67 hr; t1/2β phase 3.6 ± 1.4 hr in adult patients, increasing to 11.5 ± 3.9 hr in renal insufficiency.114,115

Adverse Reactions. Granulocytopenia (ANC <1000/μL) occurs in 13–67% of patients and is the most frequent dose-limiting adverse effect.114 Thrombocytopenia (platelets <50,000/μL) occurs in 20% of patients. CNS toxicity (headache, lethargy, dizziness, confusion, seizure, coma) has been reported at a frequency of 5–17%. Phlebitis, inflammation, and pain at the site of IV infusion occur frequently because of the high pH of the solution. Anemia, fever, rash, and abnormal liver function tests occur in about 2% of patients.114,116

Contraindications. Hypersensitivity to acyclovir or ganciclovir.

Precautions. Use with caution in renal impairment, pre-existing cytopenias, or concurrent myelosuppressive drug therapy.

Drug Interactions. Didanosine AUC can be increased when given within 2 hr of ganciclovir. Probenecid decreases the renal excretion of ganciclovir. Use extreme caution in combination with zidovudine because of additive myelosuppression. Concurrent nephrotoxic drugs can increase the nephrotoxicity of ganciclovir. Concurrent cytotoxic drugs increase the toxicity of ganciclovir. Seizures have been reported with concurrent use of ganciclovir and imipenem-cilastatin.

Parameters to Monitor. Monitor CBC and platelet counts twice weekly during induction treatment and at least weekly during maintenance. Monitor renal function at least q 2 weeks. Check injection site for phlebitis and infection daily.

Notes. Ganciclovir-resistant CMV strains have been isolated from patients during treatment.117 Disease progression caused by these strains has been observed and might require changing therapy to an alternative antiviral (eg, foscarnet).
Pharmacology. Indinavir is an HIV protease inhibitor with a mechanism of action similar to that of saquinavir. (See Antiviral Drugs for HIV Infection Comparison Chart.)

**Adult Dosage.** PO for HIV infection 800 mg q 8 hr. Take each dose on an empty stomach with water or other fat-free liquid or with light, fat-free foods (eg, toast, jelly, skim milk, coffee). PO for HIV infection with ritonavir 400 mg q 12 hr with ritonavir 400 mg q 12 hr, or 800 mg q 12 hr with ritonavir 200 mg q 12 hr. In mild to moderate hepatic insufficiency caused by cirrhosis, the dosage is 600 mg q 8 hr. The combination can be taken with food.

**Dosage Forms.** Cap 200, 333, 400 mg.

Pharmacokinetics. Indinavir is rapidly absorbed in the fasting state. Administration of indinavir with a meal high in calories, fat, or protein decreases oral absorption by about 75%. When indinavir is combined with ritonavir, food does not decrease bioavailability of indinavir. Absolute bioavailability not been determined in humans, but fasting bioavailability is 14–70% in animals. Indinavir is 60% bound to human plasma proteins. It is primarily metabolized by CYP3A4 and <20% is excreted unchanged in the urine; half-life is 1.8 ± 0.4 hr.

**Adverse Reactions.** Frequent adverse reactions are nausea, vomiting, abdominal pain, diarrhea, headache, asthenia, insomnia, taste perversion, transient elevations of hepatic transaminases, asymptomatic hyperbilirubinemia, and nephrolithiasis. Dizziness, somnolence, anorexia, malaise, and dry mouth occur occasionally. Nephrolithiasis occurred in 4% of patients in clinical trials and can be managed with hydration and temporary drug discontinuation. Patients should drink at least 1.5 L/day of liquids to ensure adequate hydration while taking indinavir.

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**LAMIVUDINE**

Pharmacology. Lamivudine (3TC) is a synthetic pyrimidine nucleoside active against HIV-1, HIV-2, and hepatitis B. Lamivudine is metabolized intracellularly to lamivudine triphosphate and acts as a chain terminator of viral DNA and a competitive inhibitor of HIV reverse transcriptase. Lamivudine alone to treat HIV infection leads to rapid emergence of high-level resistance; therefore, it is used in combination with zidovudine. Resistance to zidovudine is markedly delayed when the drug is used with lamivudine, and the combination results in greater and more sustained elevations in CD4 cell counts than zidovudine monotherapy. (See Antiviral Drugs for HIV Infection Comparison Chart.)

**Adult Dosage.** PO for HIV infection 150 mg bid. PO for chronic hepatitis B 100 mg/day. Reduce dosage in renal impairment. For HIV co-infection with hepatitis B use HIV dosage with appropriate combination antiretroviral therapy.

**Pediatric Dosage.** PO (3 months–12 yr) 4 mg/kg, to a maximum of 150 mg bid with zidovudine.

**Dosage Forms.** Tab 100, 150 mg; Soln 5, 10 mg/mL.
Pharmacokinetics. Oral bioavailability is 82%. \( V_d \) is 1.3 L/kg; elimination half-life is 2.5 hr. Excretion is primarily by the renal route, with 68–71% of drug excreted unchanged in urine.

Adverse Reactions. The most frequently reported adverse effects have been headache, fatigue, nausea, insomnia, neuropathy, and musculoskeletal pain.

NELFINAVIR MESYLATE

Pharmacology. Nelfinavir mesylate is an antiviral that inhibits HIV-1 and HIV-2 proteases by binding to the active enzymatic site, preventing cleavage of polyprotein precursors. This cleavage is essential for maturation of infectious virus, and its inhibition results in the formation of immature, noninfectious HIV particles. (See Antiviral Drugs for HIV Infection Comparison Chart.)

Administration and Adult Dosage. PO for HIV disease in combination with nucleoside analogues 750 mg tid or 1250 mg bid.¹²¹

Special Populations. Pediatric Dosage. PO for HIV disease in combination with nucleoside analogues (<2 yr) safety and efficacy not established; (2–13 yr) 20–30 mg/kg tid.

Geriatric Dosage. Not studied but expected to be the same as adult dosage.

Dosage Forms. Tab 250 mg; Pwdr 50 mg nelfinavir base/level scoopful (1 g).

Patient Instructions. (See HIV Drugs Class Instructions.) Each dose must be taken orally with a light snack or meal to increase the amount of the drug absorbed. If you are taking an oral contraceptive, you should use an alternate or additional contraceptive measure. Store nelfinavir in a dry place at room temperature. New onset diabetes mellitus, exacerbation of pre-existing diabetes mellitus, and hyperglycemia have been reported in HIV-infected patients receiving protease inhibitors. Some patients require initiation or dosage adjustments of insulin or oral hypoglycemic agents. Diabetic ketoacidosis has also occurred. Hyperglycemia persisted in some cases after drug discontinuation.

Pharmacokinetics. Fate. Bioavailability is unknown in humans, but animal data suggest an oral bioavailability of 20–80%. Nelfinavir absorption is increased 2- to 3-fold when administered with food. Peak serum concentrations occur 2–4 hr after a dose. After multiple oral doses of 750 mg tid, peak serum concentrations average 3–4 mg/L (5.3–7 \( \mu \)mol/L) and trough concentrations average 1–3 mg/L (1.8–5.3 \( \mu \)mol/L). Plasma protein binding is >98%. Nelfinavir is metabolized by cytochrome P450 enzymes, primarily CYP3A4 and to a minor extent by CYP2C9, 2C19, and 2D6. The major oxidative metabolite has in vitro antiviral activity comparable to the parent drug. Less than 2% of nelfinavir is excreted unchanged in urine.

\( t_{1/2} \) 3.5–5 hr.

Adverse Reactions. Diarrhea, abdominal pain or discomfort, flatulence, nausea, rash, and difficulty swallowing tablets are frequent. Diarrhea often resolves spontaneously 1–2 weeks after initiation of therapy. Antidiarrheal medications are often beneficial in alleviating or minimizing symptoms. Oral calcium carbonate 500-1000 mg once or twice daily has decreased prevalence of nelfinavir-associ-
ated diarrhea in some patients. Occasional reactions include asthenia, headache, and fatigue.

**Contraindications.** (See Drug Interactions.)

**Precautions.** Do not use nelfinavir as monotherapy. Appropriate use is with other antiretroviral therapy to reduce potential for developing drug resistance. Nelfinavir powder for oral solution contains 11.2 mg phenylalanine/g of powder and should be used cautiously in patients with phenylketonuria.

**Drug Interactions.** Nelfinavir is an inhibitor of CYP3A and can cause increased serum concentrations of drugs primarily metabolized by CYP3A. It is also a substrate for CYP3A, and nelfinavir concentrations can be affected by the induction or inhibition of CYP3A by other drugs. Do not co-administer nelfinavir with rifampin because it decreases nelfinavir’s steady-state AUC by 82%. Co-administration with rifabutin reduces nelfinavir’s AUC by 32% and increases rifabutin’s AUC by 207%. If administered together, the manufacturer recommends reducing the rifabutin dosage by 50%, although alternatives should be considered. Avoid other drugs (eg, carbamazepine, phenobarbital, phenytoin) that strongly induce CYP3A4 because they can substantially reduce nelfinavir serum concentrations. Avoid co-administration with astemizole or cisapride because of possible prolonged QT intervals and serious cardiovascular adverse events. Co-administration with ethinyl estradiol/norethindrone resulted in a 47% decrease in ethinyl estradiol serum concentration and an 18% decrease in norethindrone serum concentration. Alternative contraceptives need to be used while receiving nelfinavir therapy. Co-administration with indinavir results in an 83% increase in nelfinavir AUC and a 51% increase in indinavir AUC. Co-administration with ritonavir results in a 152% increase in nelfinavir AUC and minimal change in ritonavir AUC. Various protease inhibitor combinations are under study, but safety and efficacy of these combinations have not been established.

**Parameters to Monitor.** Monitor clinical signs, symptoms, and laboratory markers for progression of HIV disease to help decide regimen changes in antiretroviral therapy. Baseline CD4+ and HIV-1 RNA polymerase chain reaction viral load tests are standard of practice markers to measure clinical benefit of therapy. Monitor adherence to the drug regimen throughout treatment course to help in assessment of effectiveness. Repeat tests after 1 month and q 3–4 months to monitor benefit of antiretroviral therapy.

**NEVIRAPINE**

**Pharmacology.** Nevirapine is a dipyridodiazepinone nonnucleoside HIV-1 reverse transcriptase inhibitor. Nevirapine and other reverse transcriptase inhibitors are not active against HIV-2 reverse transcriptase. The inhibition by nevirapine is noncompetitive, and the binding site is located near but not directly at the catalytic amino acid residues, which might provide nevirapine activity against HIV-1 mutants that are resistant to nucleoside reverse transcriptase inhibitors. Nevirapine provides added benefit (eg, increased CD4 count, decreased viral load) in combination with zidovudine and didanosine.122-125 (See Antiviral Drugs for HIV Infection Comparison Chart.)
Adult Dosage. PO for HIV 200 mg/day for 2 weeks, followed by 200 mg bid or 400 mg once daily.

Pediatric Dosage. PO (<13 yr) 120 mg/m²/day for 2 weeks, then bid.

Dosage Forms. Tab 200 mg; Susp 10 mg/mL.

Pharmacokinetics. Oral absorption is not affected by food or antacids; bioavailability is 90%. The median time to peak concentration is 4 hr after a 400 mg dose with average peak concentrations after the first dose of 3.4 ± 1 mg/L. Peak and trough concentrations average 7.2 ± 1.4 mg/L (27 ± 5 μmol/L) and 4 ± 1.2 mg/L (15 ± 4 μmol/L), respectively, after 14 days of therapy. The average elimination half-life is 45 hr in the initial 2-week period and decreases to 25–30 hr thereafter because of metabolic autoinduction mediated by the cytochrome P450 system. Less than 3% of the dose is excreted renally.

Adverse Reactions. A mild to moderate rash occurs in up to 48% of patients. Rash can be associated with liver function test elevations and a low frequency of clinical hepatitis. Severe, occasionally fatal, hepatotoxicity has occurred in those using nevirapine in postexposure prophylactic regimens with various other antiretrovirals. This use is not recommended, but the single-dose use to prevent HIV transmission appears to be safe. The risk of developing rash is highest within 2 weeks of drug initiation or dosage escalation to 400 mg/day and is reduced by following the recommended dosage escalation schedule. Other occasional adverse reactions are arthralgia, fatigue, fever, myalgia, and somnolence.

Parameters to Monitor. Monitor liver function closely for at least the first 12 weeks of therapy and periodically thereafter.

RITONAVIR Norvir

Pharmacology. Ritonavir is an HIV protease inhibitor with a mechanism of action similar to saquinavir. (See Antiviral Drugs for HIV Infection Comparison Chart.)

Adult Dosage. PO for treatment of HIV 600 mg q 12 hr with food in combination with nucleoside analogues. Ritonavir might be better tolerated initially if the dosage is initiated at 300 mg q 12 hr and increased to 600 mg q 12 hr over 10–14 days. If the 600 mg q 12-hr dosage is not reached after 2 weeks of therapy, discontinue therapy because the risk of developing viral resistance to ritonavir or cross-resistance to other protease inhibitors is increased with lower dosages. PO in protease inhibitor combination treatment of HIV (see Antiviral Drugs for HIV Infection Comparison Chart).

Dosage Forms. Cap 100 mg; Soln 80 mg/mL. Capsules must be refrigerated.

Pharmacokinetics. Ritonavir is rapidly absorbed and increased by approximately 15% with food. Absolute bioavailability has not been determined in humans, but bioavailability is 30–70% in animals. Ritonavir is 98–99% protein bound, primarily to albumin and α1-acid glycoprotein. After a 600 mg oral dose taken with food, a peak serum concentration of 11.2 ± 3.6 mg/L (15.5 ± 5 μmol/L) occurs at 3.3 ± 2.2 hr and the trough is 3 ± 2.1 mg/L (4.2 ± 2.9 μmol/L). Serum concentrations
can decrease over time because of autoinduction of the CYP3A and CYP2D isoenzymes responsible for metabolism of ritonavir.

**Adverse Reactions.** Nausea, vomiting, diarrhea, asthenia, anorexia, abdominal pain, taste perversion, perioral paresthesia, peripheral paresthesia, headache, insomnia, and elevated serum triglyceride concentrations occur frequently. Occasionally, elevations of hepatic transaminases and CPK occur.

**Drug Interactions.** Ritonavir is a potent inhibitor of several cytochrome P450 enzymes (CYP2C9, 2C19, 2D6, and 3A3/4) and can produce large increases in serum concentrations of highly metabolized drugs. Consult the product information for contraindicated drugs and carefully review the patient’s medication list for interactions before starting this therapy.

**Pharmacology.** Saquinavir is a synthetic peptide-like substrate analogue that inhibits HIV protease. Inhibition of HIV protease prevents the cleavage of polyprotein precursors, which is essential for maturation of infectious virus.\(^{128,129}\) Saquinavir mesylate is formulated in a hard gelatin capsule. Saquinavir has been reformulated into a soft gelatin capsule that combines saquinavir base in an oil-like substance that allows microdispersion upon contact with gastric fluids enhancing oral bioavailability. (See Antiviral Drugs for HIV Infection Comparison Chart.)

**Administration and Adult Dosage.** PO for advanced HIV disease in combination with other nucleoside analogues (saquinavir mesylate) 600 mg q 8 hr (FDA-approved regimen but achieves inadequate serum concentrations to suppress HIV); (saquinavir) 1200 mg q 8 hr.

**Special Populations.** Pediatric Dosage. (<16 yr) safety and efficacy not established.

**Geriatric Dosage.** Not studied but expected to be same as adult dosage.

**Dosage Forms.** (Saquinavir) Cap 200 mg; (saquinavir mesylate) Cap 200 mg.

**Patient Instructions.** (See HIV Drugs Class Instructions.) Saquinavir mesylate (Invirase) must be taken within 2 hours after a full meal to achieve adequate concentrations of drug to inhibit viral replication. Saquinavir (Fortovase) is better absorbed and requires a snack or some food to help increase the amount of medication getting into the blood. Store saquinavir in the refrigerator. New onset diabetes mellitus, worsening of pre-existing diabetes mellitus, and hyperglycemia have been reported in HIV-infected patients receiving protease inhibitors. Some patients require initiation or dosage adjustments of insulin or oral hypoglycemic agents. Diabetic ketoacidosis also has occurred. Hyperglycemia persists in some cases after drug discontinuation.

**Pharmacokinetics.** Fate. Oral absorption of saquinavir mesylate is erratic and the drug undergoes extensive first-pass metabolism. Approximately 30% of a 600 mg dose is absorbed when given within 2 hr after food; absolute bioavailability averages 4%. Saquinavir bioavailability relative to saquinavir mesylate is
Saquinavir is 98% plasma protein bound; concentrations in the CSF are negligible. Saquinavir undergoes metabolism primarily by CYP3A4; Cl is 1.14 L/hr/kg.\(^{123}\)

\(t^{1/2} = 12\) hr.

**Adverse Reactions.** Abdominal discomfort or pain, diarrhea, anorexia, and nausea occur frequently. Occasional adverse reactions include asthenia, rash, elevations of transaminases, and headache. Rare reactions include ataxia, confusion, hemolytic anemia, thrombophlebitis, attempted suicide, seizures, and exacerbation of chronic liver disease.

**Contraindications.** (See Drug Interactions.)

**Precautions.** Do not use saquinavir as monotherapy because of the greater potential for developing resistance.

**Drug Interactions.** Do not administer saquinavir with rifampin because steady-state AUC of saquinavir decreases by 80%. Administration with rifabutin reduces saquinavir plasma concentrations by 40% and alternatives to this combination should be considered. Avoid other drugs that strongly induce CYP3A4 because they can substantially decrease saquinavir serum concentrations. Avoid coadministration with astemizole or cisapride because of possible prolonged QT intervals and serious cardiovascular adverse events. Concurrent ketoconazole and possibly other inhibitors of CYP3A4 can increase the bioavailability and half-life of saquinavir. (See Cytochrome P450 Enzyme Interactions.) Ingesting grapefruit juice with saquinavir has been suggested to increase the bioavailability of saquinavir by inhibition of CYP3A4. However, the grapefruit juice must be concentrated, taken with every dose of saquinavir, and contain flavinoids to have any benefit. This method is not likely to be palatable to most patients because of gastric irritation and appears unnecessary with the soft gelatin capsule formulation of saquinavir.

**Parameters to Monitor.** (See Nelfinavir.)

**Notes.** Saquinavir (Fortovase) should be refrigerated; once brought to room temperature (\(\leq 25^\circ\)C), use it within 3 months. Fortovase has a dosage of 1200 mg tid to achieve saquinavir plasma concentrations sufficient to inhibit the replication of HIV. The hard gelatin capsule formulation dosage of 600 mg tid does not consistently achieve adequate saquinavir plasma concentrations. The use of ritonavir 400 mg bid and saquinavir 400 mg bid in combination has been used to improve concentrations of saquinavir and tolerance of ritonavir.

**STAVUDINE**

**Pharmacology.** Stavudine (d4T) is a synthetic pyrimidine nucleoside reverse transcriptase inhibitor that is structurally similar to zidovudine and has been shown to inhibit HIV replication in vitro. Stavudine is phosphorylated by cellular enzymes to stavudine triphosphate, which acts as a competitive inhibitor of HIV reverse transcriptase and an alternative nucleoside substrate, which leads to premature elongation of viral DNA.\(^{130,131}\) (See Antiviral Drugs for HIV Infection Comparison Chart.)
**ZIDOVUDINE**

**Pharmacology.** Zidovudine is a thymidine analogue that inhibits HIV replication. It is converted to the active monophosphate form by thymidine kinase and ultimately to zidovudine triphosphate by intracellular enzymes. This form exerts its activity at viral DNA polymerase (reverse transcriptase) by competing with other cellular deoxynucleosides and by acting as a chain terminator of DNA synthesis. (See Antiviral Drugs for HIV Infection Comparison Chart.)

**Administration and Adult Dosage.** PO for HIV infection with 300 mg bid or 200 mg tid. PO for maternal–fetal HIV transmission (maternal) 300 mg bid, begun after the 14th week of pregnancy and continued throughout the pregnancy, then IV during labor 2 mg/kg over 1 hr, followed by a continuous infusion of 1 mg/kg/hr until delivery. (See also Pediatric Dosage.) PO for combination therapy with zalcitabine 200 mg q 8 hr with zalcitabine 0.75 mg q 8 hr. PO for post-exposure prophylaxis 1–1.5 g/day in 4 or 5 divided doses has been used, but the effectiveness of this regimen is not confirmed in humans and informed consent should be obtained. IV for patients unable to take oral medication 1–2 mg/kg q 4 hr infused over 1 hr, only until oral therapy can be initiated.

**Special Populations. Pediatric Dosage.** PO for prevention of maternal HIV transmission 2 mg/kg/dose q 6 hr for first 6 weeks of life, beginning 8–12 hr after birth. IV for prevention of maternal HIV transmission if unable to receive PO 1.5 mg/kg/dose q 6 hr until oral therapy can be initiated. PO for HIV infection (0–2 weeks) 2 mg/kg/dose q 6 hr; (2–4 weeks) 3 mg/kg/dose q 6 hr; (4 weeks–13 yr) 180 mg/m²/dose (to a maximum of 200 mg) q 6 hr; (over 13 yr) 100 mg q 4 hr 5 times/day.

**Geriatric Dosage.** Same as adult dosage but adjust for age-related reduction in renal function.

**Other Conditions.** Reduce dosage by 50% in patients with Clcr <25 mL/min and 75% in those with cirrhosis.
**Dosage Forms.** Cap 100 mg; Tab 300 mg; Syrup 10 mg/mL; Inj 10 mg/mL. (See Notes.)

**Patient Instructions.** (See HIV Drugs Class Instructions.) This drug is not a cure for HIV disease. Opportunistic infections and other complications associated with HIV infection can continue to develop. This drug may be taken with food to decrease abdominal discomfort or nausea. It is important to have blood counts followed closely during therapy to monitor for decreases in blood cell counts.

**Pharmacokinetics.**

**Serum Levels.** Not established; intracellular concentrations of zidovudine triphosphate might correlate with therapeutic benefit, but in vivo data are not available.

**Fate.** Zidovudine (ZDV) undergoes marked presystemic metabolism. Oral bioavailability is 60–70%, possibly reduced with high-fat meals. Peak serum levels are approximately 1.2 mg/L (4.5 μmol/L) after a 250 mg oral dose. Protein binding is less than 25%. CSF concentrations are 24% of serum in children receiving a continuous infusion of the drug. $V_{dss}$ is 1.6 ± 0.6 L/kg; $Cl$ is 1.3 ± 0.3 L/hr/kg in adults and 36.4 ± 11.5 L/hr/m² in children. ZDV is rapidly metabolized to the inactive ether glucuronide (GZDV). GZDV formation is reduced, and zidovudine AUC and half-life are increased in patients with cirrhosis. About 60% of an oral dose is excreted as GZDV in urine. GZDV excretion is reduced in patients with renal dysfunction; hemodialysis removes GZDV but not ZDV.¹⁰⁷,¹³⁴,¹³⁵ $t_{1/2}$: (Adults) 1.1 ± 0.2 hr; 2.1 hr in uremia; 2.4 hr in cirrhosis.⁹⁷ (Children) 1.5 ± 0.6 hr.

**Adverse Reactions.** Severe anemia and granulocytopenia occur frequently and might necessitate blood transfusions; epoetin might help alleviate anemia in patients with low serum erythropoietin levels. Other frequent adverse reactions associated with zidovudine in placebo-controlled trials include abdominal discomfort, nausea, vomiting, insomnia, myalgias, and headaches. Adverse reactions that occasionally occur with long-term use (>12 weeks) are myopathy and nail pigmentation.¹⁰⁰

**Contraindications.** Life-threatening allergy to the drug or its components.

**Precautions.** Pregnancy; lactation. Use with caution in liver disease or hepatomegaly, especially in obese women.

**Drug Interactions.** Several drugs decrease the glucuronidation of zidovudine, including atovaquone, methadone, probenecid, valproic acid, and possibly fluconazole; rifampin increases zidovudine glucuronidation; however, the clinical importance of these interactions is not established.¹⁰⁴ Initial studies showed that prolonged administration of acetaminophen was associated with increased hematologic toxicity from zidovudine, but further study does not support this finding.¹³⁶

**Parameters to Monitor.** Hemoglobin, hematocrit, MCV, and WBC for hematologic toxicity. Monitor clinical signs, symptoms, and laboratory markers for progression of HIV disease to help decide regimen changes in antiretroviral therapy. Baseline CD4 and HIV-1 RNA polymerase chain reaction viral load tests are useful to measure clinical benefit of therapy. Repeat tests after 1 month and q 3–4 months thereafter have been suggested to monitor benefit of antiretroviral therapy.
**Notes.** Viral resistance to zidovudine has occurred in vitro with isolates recovered from patients and is associated with prolonged zidovudine use and more advanced disease; correlation between viral resistance in vitro and progression of disease has not been established. Studies with lamivudine (3TC) suggest that the combination can delay or prevent HIV-1 viral resistance to zidovudine. Aztec (Verex) is an SR dosage form in late-stage testing.

**ANTIRETROVIRAL THERAPY FOR HIV**

The use of protease inhibitors and/or nonnucleoside reverse transcriptase inhibitors in combination with nucleoside reverse transcriptase inhibitors has dramatically changed the treatment of HIV infection. Regimens containing a protease inhibitor or nonnucleoside reverse transcriptase inhibitor have enhanced the ability to inhibit replication of HIV, affecting immunologic and viral markers, delaying progression of disease, and improving survival. Many formidable hurdles stand in the way of effective treatment, including patient adherence to dosage regimens, adverse effects, and drug–drug interactions. These hurdles interfere with quality of life and control of the viral burden and also contribute to the emergence of resistance. It is essential for health care providers and patients to appreciate the complexity of antiretroviral medication regimens to achieve harmony between goals of antiretroviral therapy and optimal patient care. General principals of treatment that guide contemporary treatment decisions are outlined below:

- Viral load monitoring is essential to guide decision making.
- Attaining and maintaining an undetectable HIV RNA in blood (which can indirectly reflect lymph concentrations) is the goal of therapy.
- Introduce effective antiretroviral therapy before extensive immune system damage has occurred.
- Three-drug combination therapy, is the regimen most likely to achieve the goal of an undetectable HIV RNA level and provide a durable response.
- Compliance with the treatment regimen is critical to success and must be considered in initiating and choosing regimens.
- Change most or all drugs in a failing regimen simultaneously; use antiretroviral drug resistance testing to guide new antiretroviral regimen decisions.

For further information and clarification on appropriate uses of antiretroviral therapy, see U.S. Public Health Service guidelines for the use of antiretroviral agents in pediatric HIV infection and HIV-infected adults and adolescents (references 137 and 138).
<table>
<thead>
<tr>
<th>HIV NUCLEOSIDE REVERSE TRANSCRIPTASE INHIBITORS</th>
<th>ADULT DOSAGE</th>
<th>PEDIATRIC DOSAGE</th>
<th>EFFECT ON CYP450 ISOZYMES</th>
<th>ADVERSE REACTIONS</th>
<th>COMMENTS</th>
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</thead>
<tbody>
<tr>
<td><strong>Abacavir</strong></td>
<td>300 mg PO bid.</td>
<td>300 mg PO (3 months–16 yr)</td>
<td>No effect. Rash, asthenia, pancreatitis, diarrhea, headache, nausea, pruritus</td>
<td>No effect.</td>
<td>Patients who have a hypersensitivity reaction must not take the drug; it could be fatal.</td>
</tr>
<tr>
<td><strong>Zidovudine</strong></td>
<td>50 mg PO bid.</td>
<td>50 mg PO (infants ≤ 2 kg)</td>
<td>No effect.</td>
<td>No effect.</td>
<td></td>
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<tr>
<td><strong>Didanosine</strong></td>
<td>300 mg PO bid.</td>
<td>300 mg PO (infants ≤ 2 kg)</td>
<td>No effect.</td>
<td>No effect.</td>
<td></td>
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<tr>
<td><strong>Zalcitabine</strong></td>
<td>50 mg PO bid.</td>
<td>50 mg PO (infants ≤ 2 kg)</td>
<td>No effect.</td>
<td>No effect.</td>
<td></td>
</tr>
<tr>
<td>DRUG</td>
<td>DOSAGE FORMS</td>
<td>ADULT DOSAGE</td>
<td>PEDIATRIC DOSAGE</td>
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<tr>
<td>Lamivudine</td>
<td>Tab 100, 150 mg PO 150 mg bid.</td>
<td>PO (3 months–12 yr) 4 mg/kg q 12 hr, to a maximum of 150 mg q 12 hr.</td>
<td>No effect.</td>
<td>Nausea, headache, fatigue, rash, anorexia; generally well tolerated, but pancreatitis is a risk in pediatric population, but not in adults.</td>
<td>Reduce dosage in renal impairment: Cl\textsubscript{u} 30–49 mL/min, 150 mg/day; Cl\textsubscript{u} 15–29 mL/min, 150 mg once, then 100 mg/day; Cl\textsubscript{u} 5–14 mL/min, 150 mg once, then 50 mg/day; Cl\textsubscript{u} &lt;5 mL/min, 50 mg once, then 25 mg/day.</td>
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<tr>
<td>Stavudine</td>
<td>Cap 15, 20, 30, 40 mg PO (≤60 kg) 30 mg bid; PO (&gt;60 kg) 40 mg bid.</td>
<td>PO (≤30 kg) 1 mg/kg q 12 hr.</td>
<td>No effect.</td>
<td>Neuropathy, headache, nausea, asthenia, insomnia, elevated hepatic enzymes.</td>
<td>Reduce dosage in renal impairment: Cl\textsubscript{u} 26–50 mL/min, reduce dosage by 50% and give q 12 hr; Cl\textsubscript{u} 10–25 mL/min, reduce dosage by 50% and give q 24 hr.</td>
</tr>
<tr>
<td>Zalcitabine</td>
<td>Tab 0.375, 0.75 mg PO 0.75 mg tid.</td>
<td>Not established.</td>
<td>No effect.</td>
<td>Neuropathy, oral and esophageal ulceration, elevated hepatic enzymes, pancreatitis, rash, pruritus.</td>
<td>Reduce dosage in renal impairment: Cl\textsubscript{u} 10–40 mL/min, same dose q 12 hr; Cl\textsubscript{u} &lt;10 mL/min, same dose q 24 hr.</td>
</tr>
</tbody>
</table>

(continued)
### ANTIVIRAL DRUGS FOR HIV INFECTION COMPARISON CHART (continued)

<table>
<thead>
<tr>
<th>DRUG</th>
<th>DOSAGE FORMS</th>
<th>ADULT DOSAGE</th>
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</thead>
</table>
| Zidovudine Retrovir | Cap 100 mg Tab 300 mg Syrup 10 mg/mL Inj 10 mg/mL. | PO 200 mg tid or 300 mg bid. (See monograph for other indications.) | PO (neonates) 2 mg/kg q 6 hr; IV (neonates) 1.5 mg/kg q 6 hr (infants and children) 80 mg/m² q 6 hr | No effect. Bone marrow suppression (anemia, neutropenia), nausea, abdominal pain, elevated hepatic enzymes, headache, malaise, elevated CPK, myopathy, nail discoloration. | Reduce dosage in renal impairment: Cl<sub>r</sub> ≤25 mL/min, reduce recommended dosage by 50%.

| Zidovudine, Lamivudine and Abacavir Trizivir | Tab 300 mg zidovudine plus 150 mg lamivudine plus 300 mg abacavir. | PO 1 tablet bid. Not established | No effect. (See individual agents.) | Contraindicated in renal impairment. |

<p>| HIV NUCLEOTIDE REVERSE TRANSCRIPTASE INHIBITORS | | | | |
| Tenofovir DF Disoproxil Fumarate (Investigational Gilead) | Tab 300 mg. | PO 300 mg/day Not established. | No effect. Nausea | Well tolerated in early trials. Activity may be enhanced by concomitant lamivudine. |</p>
<table>
<thead>
<tr>
<th>DRUG</th>
<th>NONNUCLEOSIDE REVERSE TRANSCRIPTASE INHIBITORS</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Delavirdine</strong></td>
<td>Rescriptor (Tab 100, 200 mg.) PO 400 mg tid. or 600 mg bid. Not established. Inhibits CYP2C9, CYP2C19, CYP3A4.</td>
</tr>
<tr>
<td><strong>Efavirenz</strong></td>
<td>Sustiva (Cap 50, 100, 200 mg.) PO 600 mg qd. (10–14 kg) 200 mg/day; (15–19 kg) 250 mg/day; (20–24 kg) 300 mg/day; (25–32.5 kg) 350 mg/day; (32.5–39 kg) 400 mg/day; (≥40 kg) 600 mg/day. Induces CYP3A4 Inhibits CYP2C9, CYP2C19, CYP3A4.</td>
</tr>
<tr>
<td><strong>Nevirapine</strong></td>
<td>Viramune (Tab 200 mg) PO 200 mg/day for 14 days, then 400 mg/day in 1 or 2 doses. (See comments.) PO initiate with 120 mg/m² once daily for 14 days, then increase to full dosage of 120–200 mg/m² q 12 hr. Induces CYP3A4.</td>
</tr>
</tbody>
</table>
**ANTIVIRAL DRUGS FOR HIV INFECTION COMPARISON CHART (continued)**

<table>
<thead>
<tr>
<th>DRUG</th>
<th>DOSAGE FORMS</th>
<th>ADULT DOSAGE</th>
<th>PEDIATRIC DOSAGE</th>
<th>EFFECT ON CYP450 ISOZYMES</th>
<th>ADVERSE REACTIONS</th>
<th>COMMENTS</th>
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<tbody>
<tr>
<td><strong>PROTEASE INHIBITORS</strong></td>
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<tr>
<td>Amprenavir</td>
<td>Cap 50, 150 mg</td>
<td>PO 1.2 g bid.</td>
<td>PO 20–22.5 mg/kg bid</td>
<td>Inhibits CYP3A4, CYP2C19, CYP2E1</td>
<td>Rash (frequent), nausea, vomiting, diarrhea, flatulence, perioral paresthesias, triglycerides, LFTs, and glucose.</td>
<td>Rash usually occurs within 9 days and resolves in 1 week after discontinuation; Stevens-Johnson syndrome has occurred.</td>
</tr>
<tr>
<td>Agenerase</td>
<td>Soln 15 mg/mL</td>
<td>Combination: PO 600 mg plus ritonavir PO 100 mg bid.</td>
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<tr>
<td>Indinavir Mesylate</td>
<td>Cap 200, 333, 400 mg.</td>
<td>PO 800 mg q 8 hr.</td>
<td>PO 500 mg/m&lt;sup&gt;2&lt;/sup&gt; q 8 hr (under study in clinical trials).</td>
<td>Inhibits CYP3A4</td>
<td>Nausea, headache, abdominal pain, hyperbilirubinemia, insomnia, dizziness, nephrolithiasis.</td>
<td>Administer on an empty stomach 1 hr before or 2 hr after a meal (or can take with a light meal). Adequate hydration is required to minimize risk of nephrolithiasis.</td>
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<tr>
<td>Crixivan</td>
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<tr>
<td>Lopinavir and Ritonavir Kaletra</td>
<td>Cap 133.3 mg lopinavir plus 33.3 mg ritonavir.</td>
<td>PO 3 caps bid.</td>
<td>Not established</td>
<td>(See ritonavir.)</td>
<td>(See ritonavir.)</td>
<td>Generally well tolerated because of lowered ritonavir dosage. Refrigerate but may keep at room temperature for 30 days.</td>
</tr>
<tr>
<td>DRUG</td>
<td>DOSAGE FORMS</td>
<td>ADULT DOSAGE</td>
<td>PEDIATRIC DOSAGE</td>
<td>EFFECT ON CYP450 ISOZYMES</td>
<td>ADVERSE REACTIONS</td>
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<tr>
<td>Nelfinavir</td>
<td>Tab 250 mg</td>
<td>PO 750 mg tid or 1250 mg bid.</td>
<td>PO 20–30 mg/kg tid.</td>
<td>Inhibits CYP3A4</td>
<td>Diarrhea, nausea, dysphagia, rash.</td>
<td>Administer with food or light snack to increase absorption 2- to 3-fold.</td>
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<tr>
<td>Mesylate</td>
<td>Pwdr 50 mg</td>
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<tr>
<td>Viracept</td>
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<tr>
<td>Ritonavir</td>
<td>Capb 100 mg</td>
<td>PO 600 mg q 12 hr.</td>
<td>PO 400 mg/m² q 12 hr.</td>
<td>Inhibits CYP3A4</td>
<td>Nausea, vomiting, diarrhea, headache, circumsoral and extremity paresthesias, asthenia, taste perversion, elevated serum triglycerides, hepatic transaminases, CPK, uric acid.</td>
<td>Titrate dosage from 300 mg q 12 hr to 600 mg q 12 hr over 10–14 days to reduce adverse events.</td>
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<td>Norvir</td>
<td>Soln8 80 mg/mL</td>
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<td>Combination:</td>
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<tr>
<td>Saquinavir</td>
<td>Cap 200 mg.</td>
<td>PO 1.2 g tid.</td>
<td>Not established.</td>
<td>Inhibits CYP3A4</td>
<td>Abdominal cramping, nausea, diarrhea, headache.</td>
<td>Bioavailability relative to Invirase formulation is 331%. Refrigerate capsules, but may keep at room temperature for 90 days.</td>
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<td>Fortovase</td>
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<td>Combination:</td>
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<tr>
<td>Saquinavir</td>
<td>Cap 200 mg.</td>
<td>PO 600–1800 mg tid.</td>
<td>Not established.</td>
<td>Inhibits CYP3A4</td>
<td>Nausea, headache, elevated hepatic transaminases.</td>
<td>Bioavailability is 4% and erratic; use Fortovase if tolerated.</td>
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<tr>
<td>Mesylate</td>
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<tr>
<td>Invirase</td>
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</table>

- Saquinavir Mesylate Invirase

- Under study in clinical trials.
- Ritonavir capsules must be kept refrigerated. Ritonavir solution must be stored in the original container.
- Adult dosage escalation for ritonavir: days 1–2, 300 mg PO bid; days 3–5, 400 mg PO bid; days 6–13, 500 mg PO bid; day 14, 600 mg PO bid.
- Pediatric dosage escalation for ritonavir: Initiate therapy at 250 mg/m² q 12 hr and increase stepwise to full dosage over 5 days as tolerated.

From references 137–139 and product information.
**OSELTIMIVIR PHOSPHATE**

**Pharmacology.** Oseltamivir phosphate is the ethyl ester prodrug of oseltamivir carboxylate, which is a selective inhibitor of the enzyme neuraminidase. (See Zanamavir.)

**Administration and Adult Dosage.** PO for treatment of influenza virus A or B (start within 48 hr of onset of symptoms) 75 mg bid for 5 days.

**Special Populations.**
- **Pediatric Dosage.** (<18 yr) Safety and efficacy not established.
- **Geriatric Dosage.** Same as adult dosage.
- **Other Conditions.** In renal insufficiency (Cl\(_r\) 10–30 mL/min) reduce dose to 75 mg/day for 5 days. There is no dosage information for Cl\(_r\) <10 mL/min.

**Dosage Forms.** Cap 75 mg.

**Patient Instructions.** Begin treatment with oseltamivir within 2 days of initial flu symptoms. Oseltamivir is not a substitute for influenza vaccination.

**Missed Doses.** Take this drug at regular intervals. If you miss a dose of this medicine, take it as soon as you remember. If it is almost time for your next dose (within 2 hours), take that dose only and go back to your regular dosage schedule. Leave at least 12 hours between doses. Do not double the dose or take extra.

**Pharmacokinetics.**

- **Fate.** Oseltamivir phosphate is extensively absorbed after oral ingestion and converted by hepatic esterases to the active oseltamivir carboxylate. Food does not affect overall systemic exposure to the oseltamivir carboxylate. Oral bioavailability of oseltamivir carboxylate is >75% after a 75 mg dose. The peak serum concentration is 348 ± 63 μg/L within 2–3 hr after a 75 mg dose. Protein binding of oseltamivir carboxylate is approximately 3%. V\(_d\) is estimated to be 0.35 ± 0.02 L/kg. Oseltamivir is eliminated (>99%) by renal excretion.\(^{140,141}\)

\( t_{1/2} \) 7.5 ± 0.7 hr.\(^{141}\)

**Adverse Reactions.** Nausea and vomiting are the most frequent adverse events, occurring in about 10% of patients. Bronchitis, insomnia, and vertigo occur occasionally.\(^ {141,143} \)

**Drug Interactions.** Oseltamivir is not a substrate and does not affect cytochrome P450 isoenzymes. There are no known drug interactions.

**Parameters to Monitor.** Progression of influenza symptoms.

**Notes.** There are no data to support the safety or efficacy in patients who begin oseltamivir after 48 hr of influenza symptom onset. Patients should continue to receive an annual influenza vaccination according to guidelines on immunization practices.

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**ZANAMIVIR**

**Pharmacology.** Zanamivir is an inhibitor of the enzyme neuraminidase (sialidase), which is essential for the replication of type A and B influenza viruses. Neuraminidase catalyzes the viral cleavage of terminal sialic acid (N-acetylneuraminic acid) and this action allows release of budded virus from infected cells.
such that virons do not aggregate at the cell surface or with each other, allowing
viral spread to occur within the host.\textsuperscript{140,143,144}

\textbf{Administration and Adult Dosage.} \textit{Inhal for influenza virus A or B (start
within 48 hr of onset of symptoms)} 10 mg (2 inhalations) bid for 5 days. Give
the first dose under the supervision of an informed healthcare professional to ob-
serve correct use of the inhalation device.

\textbf{Special Populations.} \textbf{Pediatric Dosage.} (<7 yr) Safety and efficacy not estab-
lished; (≥7 yr) same as adult dosage.

\textbf{Geriatric Dosage.} Same as adult dosage.

\textbf{Dosage Forms.} Dry Pwdr Inhal 5 mg.

\textbf{Patient Instructions.} Read and follow carefully the accompanying Patient In-
structions for Use with each Diskhaler device. Take 2 doses on the first day of
treatment if they are given at least 2 hours apart. Take doses on days 2 through 5
approximately 12 hours apart and at the same time each day. To avoid the spread
of infection, do not use the inhaler for more than one person. Zanamivir is not a
substitute for influenza vaccination.

\textbf{Missed Doses.} Take this drug at regular intervals. If you miss a dose of this medi-
cine, take it as soon as you remember. If it is almost time for your next dose, take
that dose only and go back to your regular dosage schedule. Leave at least 12
hours between doses. Do not double the dose or take extra.

\textbf{Pharmacokinetics.} \textit{Fate.} (Inhal) The peak serum concentration is 39–54 \textmu g/L
within 1–2 hr after a 10 mg inhaled dose. Oral bioavailability of inhaled zanamivir
is 4–17%. Protein binding is less than 10%. Zanamivir is excreted unchanged in
the urine.\textsuperscript{144}

\[ t_{1/2} = 3.6 \pm 1.3 \text{ hr}. \]

\textbf{Adverse Reactions.} Nasal and throat discomfort, cough, headache have occurred
in 2–3% of patients. This prevalence is similar to placebo and might be related to
inhalation of the lactose vehicle. Bronchospasm has occurred occasionally in pa-
tients with asthma or COPD.\textsuperscript{143}

\textbf{Precautions.} Use with extreme caution in patients with underlying airway dis-
cases such as asthma or COPD because of the potential for causing bronchospasm.
Instruct patients who use inhaled bronchodilators concurrently with zanamivir to
use their bronchodilators before inhaling zanamivir.

\textbf{Drug Interactions.} Zanamivir is not a substrate and does not affect cytochrome
P450 isoenzymes. There are no known clinically relevant drug interactions.

\textbf{Parameters to Monitor.} Inhalation technique, progression of influenza symp-
toms.

\textbf{Notes.} There are no data to support the safety or efficacy in patients who begin
zanamivir treatment after 48 hr of influenza symptom onset. Patients should con-
tinue to receive an annual influenza vaccination.
Amoxicillin differs from ampicillin by the presence of a hydroxyl group on the amino side chain. It has activity essentially identical to ampicillin.\textsuperscript{145,146} (See Ampicillin and β-Lactams Comparison Chart.)

**Adult Dosage.** PO 250–500 mg q 8 hr or 500-875 mg bid, to a maximum of 4.5 g/day. PO for endocarditis prophylaxis 2 g 1 hr before dental or upper airway procedures.

**Pediatric Dosage.** PO 20–40 mg/kg/day in 3 equally divided doses q 8 hr. PO for endocarditis prophylaxis 50 mg/kg 1 hr before dental or upper airway procedures.

**Dosage Forms.** Cap 250, 500 mg; Chew Tab 125, 200, 250, 400 mg; Drp 50 mg/mL; Susp 25, 50 mg/mL; Tab 500, 875 mg.

**Pharmacokinetics.** Amoxicillin is completely absorbed, with about 85% bioavailability because of a small first-pass effect. Serum concentrations are greater than those after equal doses of ampicillin; postabsorptive pharmacokinetics are identical to those of ampicillin.

**Adverse Reactions.** Adverse effects are similar to those of ampicillin, although diarrhea and rashes are much less frequent with amoxicillin.

Amoxicillin does not substitute combinations of lower-dose tablets to make a higher dose because diarrhea is markedly increased. Tab (8 hr) 250 mg amoxicillin/125 mg clavulanic acid, 500 mg amoxicillin/125 mg clavulanic acid; (12 hr) 875 mg amoxicillin/125 mg clavulanic acid; Chew Tab (8 hr) 125 mg amoxicillin/125 mg clavulanic acid, 250 mg amoxicillin/62.5 mg clavulanic acid; (12 hr) 200 mg amoxicillin/28.5 mg clavulanic acid, 400 mg amoxicillin/57 mg clavulanic acid; Susp (8 hr) 25 mg amoxicillin/6.25 mg clavulanic acid/mL, 50 mg amoxicillin/12.5 mg clavulanic acid/mL; (12 hr) 40 mg amoxicillin/5.7 mg clavulanic acid/mL, 80 mg amoxicillin/11.4 mg clavulanic acid/mL.

**Pharmacokinetics.** Peak serum clavulanate concentration is 2.6 mg/L 40–60 min after an oral dose of 250 mg amoxicillin/125 mg clavulanate. Amoxicillin phar-
macokinetics are not affected by clavulanic acid. Clavulanic acid half-life is approximately 60 min.

**Adverse Reactions.** Adverse effects of this preparation include those of amoxicillin; however, diarrhea is more frequent with the combination and depends on the dosage of clavulanate. The 12-hr formulations reduce the frequency of diarrhea. Nausea and diarrhea is less frequent when this preparation is administered with food. (See β-Lactams Comparison Chart.)

**AMPICILLIN**

**Pharmacology.** Ampicillin has a similar mechanism of action and is comparable in activity to penicillin G against Gram-positive bacteria, but is more active than penicillin G against Gram-negative bacteria. (See β-Lactams Comparison Chart.)

**Adult Dosage.** PO 250–500 mg q 6 hr. IM or IV 500 mg–3 g q 4–6 hr to a maximum of 12 g/day. Give the same dose q 12 hr with a Clcr <20 mL/min. IV or IM for endocarditis prophylaxis 2 g within 30 min of procedure.

**Pediatric Dosage.** PO (<20 kg) 50–100 mg/kg/day in 2–4 divided doses; (>20 kg) 100–400 mg/kg/day in 4–6 divided doses. IV (neonates) 25–100 mg/kg/dose q 6–12 hr—higher dosages for meningitis; (<20 kg) 50–100 mg/kg/day in 2–4 divided doses; (>20 kg) 400 mg/kg/day in 4–6 divided doses. IV or IM for endocarditis prophylaxis 50 mg/kg within 30 min of procedure.

**Dosage Forms.** Cap 250, 500 mg; Susp 25, 50 mg/mL; Inj 125, 250, 500 mg, 1, 2, 10 g.

**Pharmacokinetics.** Oral forms are about 50% absorbed in the fasting state; food delays absorption. Plasma protein binding is low, and therapeutic concentrations are attained in most tissues and fluids including CSF (in the presence of inflammation). About 90% is excreted unchanged in urine. Half-life is 1.2 hr, 2 hr in neonates, increasing to 20 hr in anuric patients.

**Adverse Reactions.** Nausea and diarrhea occur frequently with oral therapy. Other reactions include frequent skin rash (more frequent in patients receiving allopurinol and very frequent in patients with Epstein-Barr virus infection [mononucleosis]). Most of these eruptions probably are not hypersensitivity reactions but immunologically mediated. They are generally dose related (higher frequency at higher dosages), are macular rather than urticarial, and disappear with continued administration of the drug.

**ANTISTAPHYLOCOCCAL PENICILLINS**

**Pharmacology.** Methicillin, nafoxalone, oxacillin, cloxacillin, and dicloxacillin are similar to other penicillins in their mechanism of action. However, these drugs are not hydrolyzed by staphylococcal penicillinases. Therefore, nearly all isolates of *Staphylococcus aureus* and some isolates of coagulase-negative staphylococci are susceptible to these drugs. Methicillin- (actually β-lactam-) resistant staphylococci have altered penicillin-binding proteins (transpeptidases). Although these drugs are used primarily in staphylococcal infection, they retain good activity against most streptococci, except enterococci.
Adult Dosage. Oral administration of nafcillin and oxacillin is not recommended because they are poorly absorbed. (See β-Lactams Comparison Chart.)

Pediatric Dosage. (See β-Lactams Comparison Chart.)

Dosage Forms. (See β-Lactams Comparison Chart.)

Pharmacokinetics. Only cloxacillin and dicloxacillin are adequately absorbed from the GI tract. Except for methicillin, these drugs are mostly hepatically eliminated by metabolism and biliary excretion.

Adverse Reactions. Interstitial nephritis is frequent with methicillin but occurs only rarely with the other drugs. Hepatic damage occurs rarely with oxacillin. Nafcillin has a propensity for local irritation at the IV infusion site and causes neutropenia more frequently than other antistaphylococcal penicillins.

AZTREONAM

Pharmacology. Aztreonam is a monobactam with activity similar to that of third-generation cephalosporins against most Gram-negative aerobic bacteria (including *P. aeruginosa*) but it is inactive against Gram-positive bacteria and anaerobes.150–152 (See β-Lactams Comparison Chart.)

Adult Dosage. IM or IV 500 mg–2 g q 6–12 hr, to a maximum of 8 g/day, depending on severity and site of infection. Reduce maintenance dosage by 50% with a Clcr of 10–30 mL/min and by 75% with a Clcr <10 mL/min. Give one-eighth of the initial dose after hemodialysis.

Pediatric Dosage. IV (<1 month) 30 mg/kg q 6–12 hr; (1 month–16 yr) 30 mg/kg q 6–8 hr. Dosages as high as 50 mg/kg q 6 hr have been used in children with cystic fibrosis or serious Gram-negative infections (eg, *P. aeruginosa*).

Dosage Forms. Inj 500 mg, 1, 2 g.

Pharmacokinetics. Peak serum concentrations of 164 and 255 mg/L occur after 30-min IV infusions of 1 and 2 g, respectively. With inflamed meninges, CSF concentrations are similar to those observed with comparable dosages of third-generation cephalosporins, but experience in treating meningitis is limited. The drug is 60% plasma protein bound and has a Vd of about 0.24 L/kg; 60–70% is excreted in urine unchanged. The half-life is 1.5–2 hr, increasing to 6 hr in renal failure and 3.2 hr in alcoholic cirrhosis.

Adverse Reactions. Adverse effects of aztreonam are minimal. Cross-allergenicity between aztreonam and other β-lactams is low, and aztreonam has been used safely in penicillin- or cephalosporin-allergic patients.

CEPHALOSPORINS

Pharmacology. Cephalosporin antibiotics have broad-spectrum activity against many Gram-positive and Gram-negative pathogens. These agents are generally considered to be bactericidal through binding to various penicillin-binding proteins in bacteria, which results in changes in cell wall structure and function. Members of this class are frequently subdivided into “generations” based on their antimicrobial activity (as well as order of introduction into clinical use).152–156
First-generation cephalosporins have activity against Gram-positive bacteria (eg, Staphylococcus sp.) and a limited, but important, number of species of aerobic Gram-negative bacilli (eg, Escherichia coli, Klebsiella sp., Proteus mirabilis). Haemophilus influenzae and most other aerobic Gram-negative bacilli often indigenous to hospitals (eg, Enterobacter, Pseudomonas spp.) are resistant to these drugs. Anaerobic bacteria isolated in the oropharynx are generally susceptible to these agents; however, anaerobes such as Bacteroides fragilis are resistant.152,156

The second-generation cephalosporins cefamandole, cefonicid, and cefuroxime differ from first-generation agents in their improved activity against H. influenzae and some strains of Enterobacter, Providencia, and Morganella spp.152,156 The oral second-generation cephalosporins cefuroxime axetil, cephoperazone, and loracarbef (a carbacephem) have similar but less potent activity.157–160 Cefoxitin, cefmetazole, and cefotetan (which are actually cephemycins) have increased activity against anaerobes, including B. fragilis;152,156,162 the other second-generation cephalosporins have poor activity against this organism.152

Third-generation cephalosporins are noteworthy for their marked potency against common Gram-negative organisms (eg, E. coli, Klebsiella pneumoniae) and their activity against Gram-negative bacilli resistant to older agents (eg, Serratia sp., P. aeruginosa). Although grouped together, some agents have better activity against certain organisms (eg, ceftazidime is better against P. aeruginosa), and poorer activity against others (eg, cefixime and ceftazidime are poorer against Staphylococcus aureus).152,153,155,156

Fourth-generation cephalosporins have a spectrum similar to that of third-generation drugs, plus activity against some Gram-negative strains that are resistant to the third-generation agents, such as Enterobacter sp. Their antianaerobic activity is poor. Resistance to cephalosporins is mediated by β-lactamase; reduction in outer cell wall membrane permeability, and alteration of the affinity of these agents for penicillin-binding proteins. Resistance among certain β-lactamase–producing organisms (eg, Enterobacter and Citrobacter spp.) to third-generation cephalosporins has increased in recent years such that these agents cannot be relied on to provide effective therapy.156

Administration and Adult Dosage. (See β-Lactams Comparison Chart.)

Special Populations. Pediatric Dosage. (See β-Lactams Comparison Chart.)

Geriatric Dosage. Same as adult dosage but adjust for age-related reduction in renal function.

Other Conditions. Most agents require dosage modification in renal dysfunction; exceptions are ceftriaxone and cefoperazone, which have biliary and renal or primarily biliary elimination, respectively.155 Dosage reduction of all agents is required in patients with concomitant hepatic and renal dysfunction. (See β-Lactams Comparison Chart.)

Pharmacokinetics. Some of the greatest differences between agents reside in their pharmacokinetic properties. Of note is the improved CSF penetration of certain later-generation agents over the first-generation agents. Therapeutic CSF concentrations are achieved with cefotaxime, ceftriaxone, and ceftazidime; these
agents have proven efficacy in the treatment of meningitis caused by susceptible organisms in adults and children. Adequate CSF concentrations of ceftriaxone also have been observed, although its use in the treatment of meningitis is less well established. Cefuroxime penetrates adequately into CSF but is less effective for meningitis than third-generation agents. No data are available on cefepime concentrations in the CNS, but it does cross the blood–brain barrier. (See β-Lactams Comparison Chart.)

Adverse Reactions. Most cephalosporins are generally well tolerated, although a few agents have unique adverse reactions. Hypersensitivity reactions can occur in approximately 10% of patients known to be allergic to penicillin; do not administer these agents to patients with histories of an immediate reaction to penicillin. Nausea and diarrhea occur with all agents; however, diarrhea is more common with ceftriaxone and cefoperazone because of high biliary excretion. Colitis caused by Clostridium difficile has been reported with all the cephalosporins but might be more common with ceftriaxone and cefoperazone. Nephrotoxicity is rare, particularly when used without other nephrotoxic agents. All agents with an N-methylthiotetrazole (NMTT) moiety in the 3 positions of the cephem nucleus (cefoperazone, cefamandole, cefotetan, and cefmetazole) can produce a disulfiram-like reaction in some patients with ingestion of alcohol-containing beverages. In addition, these agents might be associated to varying degrees with bleeding secondary to hypoprothrombinemia, which is corrected or prevented by vitamin K administration. Although controversial, the mechanism of this reaction appears to involve inhibition of enzymatic reactions requiring vitamin K in the activation of prothrombin precursors by NMTT. However, other factors (eg, malnutrition, liver disease) might be more important risk factors for bleeding than the NMTT-containing cephalosporins. Thus, cautious use (and perhaps even avoidance) of agents with the NMTT side chain is recommended in patients with poor oral intake and critical illness. Administration of vitamin K and monitoring of the prothrombin time are indicated with these agents, particularly when therapy is prolonged. Positive direct Coombs’ tests occur frequently but hemolysis is rare. Ceftriaxone has been associated with biliary pseudolithiasis (sludging), which can be asymptomatic or resemble acute cholecystitis. This adverse effect occurs most often with dosages of ≥2 g/day, especially in patients receiving prolonged therapy or those with impaired gallbladder emptying. The mechanism is ceftriaxone-calcium complex formation, and it is usually reversible with drug discontinuation. Neonates given ceftriaxone can develop kernicterus caused by displacement of bilirubin from plasma protein binding sites; its use in this population is best avoided. Development of resistance during treatment of infections caused by Enterobacter sp., Serratia spp., and P. aeruginosa has occurred with all these agents.

**Drug Interactions.** Avoid concomitant ingestion of alcohol or alcohol-containing products with agents containing the NMTT side chain. Probenecid reduces renal clearance and increases serum levels of most agents, except those that do not undergo renal tubular secretion (eg, ceftazidime, ceftriaxone).

**Parameters to Monitor.** Monitor prothrombin time 2–3 times/week with agents having an NMTT side chain, particularly when using large dosages; monitor bleeding time with high dosages of agents having an NMTT side chain. Obtain antimicrobial susceptibility tests for development of resistance in patients relapsing during therapy. Monitor renal function tests initially and periodically during high-dose regimens or when the drug is used concurrently with nephrotoxic agents. Monitor for diarrhea, particularly with ceftriaxone and cefoperazone; test stool specimen for *C. difficile* toxin if diarrhea persists or is associated with fever or abdominal pain.

**Cefazolin Sodium**

**Pharmacology.** Cefazolin is a first-generation cephalosporin with activity against most Gram-positive aerobic organisms except enterococci and some Gram-negative bacilli (eg, *Escherichia coli*, *Klebsiella* sp., *Proteus mirabilis*).

**Adult Dosage.** IM or IV for treatment 250 mg–2 g q 6–12 hr (usually 1–2 g q 8 hr), to a maximum of 12 g/day. Decrease dosage in renal impairment. (See /H9252-**Lactams Comparison Chart.**) IM or IV for surgical prophylaxis 1 g 30–60 min before surgery. IM or IV for endocarditis prophylaxis 1 g within 30 min before a dental or upper airway procedure.

**Pediatric Dosage.** IM or IV (≤1 month) 25 mg/kg/dose given q 8–12 hr; (>1 month) 50–100 mg/kg/day in 3 divided doses given q 8 hr, to a maximum of 6 g/day. IM or IV for endocarditis prophylaxis 25 mg/kg within 30 min before a dental or upper airway procedure.

**Dosage Forms.** Inj 250, 500 mg, 1, 5, 10, 20 g.

**Pharmacokinetics.** Cefazolin is 75–85% plasma protein bound and widely distributed throughout the body, with high concentrations in many tissues and cavities but subtherapeutic concentrations in the CSF. Virtually 100% is excreted unchanged in the urine via filtration and secretion; the half-life is about 1.8 hr, increasing to 30–40 hr in renal impairment.

**Adverse Reactions.** (See Cephalosporins.)

**Cefepime Hydrochloride**

**Pharmacology.** Cefepime is a fourth-generation cephalosporin with a broader spectrum of activity than other cephalosporins. Its activity is similar to that of ceftazidime against Gram-negative bacteria, including *P. aeruginosa*, but it is also active against some isolates resistant to third-generation cephalosporins (eg, *Enterobacter* sp.). Cefepime has greater potency against Gram-positive organisms (eg, staphylococci) than ceftazidime and is similar in activity to ceftriaxone. Its anaerobic activity is poor, particularly against *Bacteroides fragilis*.
Administration and Adult Dosage. IM or IV 500 mg–2 g q 12 hr; moderate to severe infections are treated with IV 1–2 g q 12 hr. Higher dosages may be required in pseudomonal infections.

Special Populations. Pediatric Dosage. IM or IV for empiric therapy of febrile neutropenia (2 months–16 yr) 50 mg/kg q 8 hr; IM or IV for pneumonia, uncomplicated UTI, skin and soft tissue infections (2 months–16 yr) 50 mg/kg q 12 hr.

Geriatric Dosage. Same as adult dosage, adjusting for age-related renal impairment.

Other Conditions. In patients with Clcr of 30–60 mL/min, the usual dose is given q 24 hr; with a Clcr of 10–29 mL/min, 50% of the usual dose is given q 24 hr; and with a Clcr <10 mL/min, 25% of the dose (but no less than 250 mg) is given q 24 hr.

Dosage Forms. Inj 500 mg, 1, 2 g.

Pharmacokinetics. Fate. After a 30-min IV infusion of 1 g, serum concentrations of 79 mg/L are achieved. It is about 20% plasma protein bound. Cefepime penetrates most tissues and fluids well; CSF concentrations are 3.3–6.7 mg/L after 50 mg/kg q 8 hr. About 85% of a dose is eliminated renally by glomerular filtration. Elderly patients have a slightly lower total clearance, which parallels Clcr. $t_{\text{1/2}}$ 2.3 hr.

Adverse Reactions. The most common adverse reactions are injection-site reactions, rash, positive direct Coombs’ test without hemolysis, decreased serum phosphorus, increased hepatic enzymes, eosinophilia, and abnormal PT and PTT. Encephalopathy has been reported in patients with renal impairment given unadjusted dosages. (See Cephalosporins monograph.)

Contraindications. Previous immediate hypersensitivity reaction to any ß-lactam.


Parameters to Monitor. Obtain renal and hepatic function tests, and PT and PTT periodically.

**CEFOTAXIME SODIUM**

Pharmacology. Cefotaxime is a third-generation cephalosporin with activity against Gram-negative organisms resistant to first- and second-generation cephalosporins (eg, indole-positive Proteus sp., Serratia spp.). Its desacetyl metabolite (DACM) has good activity and might be synergistic with cefotaxime against certain organisms. The activity of cefotaxime against P. aeruginosa is inferior to ceftazidime and against Staphylococcus aureus is inferior to cefazolin. Cefotaxime is more active than other cephalosporins (except ceftiraxone) against Streptococcus pneumoniae that are intermediate resistant to penicillin G.\(^{152,153,155,156,170}\)

Adult Dosage. IM or IV 250 mg–2 g q 6–12 hr (usually 1–2 g q 8–12 hr), to a maximum of 12 g/day. Reduce dosage by 50% in patients with a Clcr <20 mL/min.
**Pediatric Dosage.** IV (newborns up to 1 week of age) 50 mg/kg q 12 hr; (newborns 1–4 weeks) 50 mg/kg q 8 hr; (older infants and children) 50–200 mg/kg/day (200 mg/kg/day for meningitis) given in 3–4 divided doses q 6–8 hr.

**Dosage Forms.** Inj 500 mg, 1, 2, 10 g.

**Pharmacokinetics.** CSF concentrations range from 0.3 to 0.44 mg/L after a 1 g dose and in higher dosages cefotaxime is effective for treatment of meningitis. About 50% of a dose is excreted unchanged in urine and 50% metabolized to DACM. DACM is metabolized to inactive metabolites and excreted unchanged in urine.

**Adverse Reactions.** Cefotaxime is well tolerated, with coagulopathies only rarely reported. (See Cephalosporins.)

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**CEFOTETAN DISODIUM**

**Pharmacology.** Cefotetan is a cephemycin, structurally and pharmacologically similar to the cephalosporins, particularly second-generation agents, and it contains an N-methylhidotetrazole side chain. It has greater activity against enteric Gram-negative bacteria than first- and second-generation cephalosporins and superior activity against *Bacteroides fragilis* and other anaerobic bacteria (comparable to cefoxitin and cefmetazole). Gram-positive activity is less than that of cefazolin.152,161,162

**Adult Dosage.** IV or IM for treatment 500 mg–2 g q 12–24 hr (usually 1–2 g q 12 hr), to a maximum of 6 g/day; IV or IM for surgical prophylaxis 1–2 g 30–60 min before surgery, then 1–2 g q 12 hr for up to 24 hr postoperatively. Reconstitute the drug with 0.5% lidocaine for IM administration because IM injection is painful. Give usual dose q 24 hr with a Clcr of 10–30 mL/min, and q 48 hr in patients with a Clcr <10 mL/min.

**Pediatric Dosage.** Safety and efficacy not established. IV 40–60 mg/kg/day given in equally divided doses q 12 hr.

**Dosage Forms.** Inj 1, 2, 10 g.

**Pharmacokinetics.** Cefotetan is excreted primarily unchanged in urine, with an elimination half-life of 3.5 hr.

**Adverse Reactions.** (See Cephalosporins.)

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**CEFTAZIDIME**

**Pharmacology.** Ceftazidime is a third-generation cephalosporin with activity generally similar to that of cefotaxime, but having superior activity against *P. aeruginosa* and inferior activity against Gram-positive (particularly against *Staphylococcus aureus* and penicillin-resistant pneumococci) and anaerobic bacteria.152–156,170

**Adult Dosage.** IM or IV 500 mg–2 g q 8–12 hr; q 12-hr administration appears to be adequate in the elderly. Reduce dosage by 50% with a Clcr of 30–50 mL/min; with a Clcr of 15–30 mL/min, the maximum dosage is 1 g q 24 hr; with a Clcr <15 mL/min, the dosage is 500 mg q 24–48 hr.
**Pediatric Dosage.** IV (newborns) 30 mg/kg q 12 hr; (older infants and children) IM or IV 30–50 mg/kg q 8 hr, to a maximum of 6 g/day (225 mg/kg/day for treatment of meningitis).

**Dosage Forms.** Inj 500 mg, 1, 2, 6, 10 g. Conventional formulations of cefazidime release carbon dioxide during reconstitution; the lysine formulation (eg, Ceptaz) avoids this problem.

**Pharmacokinetics.** Ceftazidime is less than 20% plasma protein bound and 80–90% excreted unchanged in urine by filtration, with a half-life of 1.6 hr, which increases to 25–34 hr in renal failure.

**Adverse Reactions.** The drug is generally well tolerated. (See Cephalosporins.)

**CEFUROXIME SODIUM**

**CEFUROXIME AXETIL**

**Pharmacology.** Cefuroxime is a second-generation cephalosporin whose activity is greater than cefazolin but less than cefotaxime, against *Haemophilus influenzae*, including β-lactamase–producing strains. The activity of cefuroxime against *Staphylococcus aureus* is slightly less than that of cefazolin. Its activity against anaerobes is poor, similar to the first-generation cephalosporins.

**Adult Dosage.** IM or IV for treatment 750 mg–1.5 g q 8 hr (q 6 hr in serious infections); IM or IV for prophylaxis 1.5 g 1 hr before surgery; doses of IM or IV 750 mg may be given q 8 hr for up to 24 hr postoperatively (1.5 g q 12 hr to a total of 6 g for open heart surgery). Reduce parenteral dosage in renal impairment; with a Cl<sub>cr</sub> of 10–20 mL/min, give the usual dose q 12 hr; with a Cl<sub>cr</sub> <10 mL/min, give the usual dose q 24 hr. PO 125–500 mg q 12 hr.

**Pediatric Dosage.** IM or IV (newborns) 10–25 mg/kg q 12 hr; (older infants and children) 50–100 mg/kg/day, to a maximum of 250 mg/kg/day for meningitis in 3–4 divided doses. PO 15–20 mg/kg q 12 hr in children (40 mg/kg/day for otitis media); it may be given in applesauce.

**Dosage Forms.** Inj 750 mg, 1.5, 7.5 g; Susp 25, 50 mg/mL; Tab 125, 250, 500 mg. Do not interchange the tablets and suspension on a mg/kg basis. (See β-Lactams Comparison Chart.)

**Pharmacokinetics.** In adults, oral bioavailability appears to be lower with the suspension than with the tablets, and food increases the bioavailability of the tablets. After absorption of oral cefuroxime axetil, it is hydrolyzed in the bloodstream to cefuroxime. Cefuroxime’s pharmacokinetics are similar to cefazolin’s, but CSF concentrations are adequate for treatment of meningitis caused by certain organisms; however, the third-generation agents ceftriaxone and cefotaxime are superior in *H. influenzae* meningitis. Over 95% of the drug is excreted unchanged in the urine and the elimination half-life is 1.2 hr.

**Adverse Reactions.** The drug is generally well tolerated. (See Cephalosporins.)

**EXTENDED-SPECTRUM PENICILLINS**

**Pharmacology.** The carboxypenicillin ticarcillin and the acylureidopenicillins (mezlocillin and piperacillin) have the same mechanisms of action as other peni-
cillins but are more active against enteric Gram-negative bacteria and *Pseudomonas aeruginosa*. Ticarcillin is not active against *Klebsiella* sp., but the acylureido derivatives have activity and are generally more potent against susceptible isolates. The acylureidopenicillins also have activity comparable to those of ampicillin against enterococci. The combination of clavulanic acid plus ticarcillin is active against *Klebsiella* sp. as well as β-lactamase–producing staphylococci, *Haemophilus influenzae*, and *Bacteroides* sp. The combination of tazobactam plus piperacillin is similar to clavulanic acid plus ticarcillin. These two combination products are not appreciably more active against *P. aeruginosa* or *Enterobacter cloacae* than ticarcillin or piperacillin alone.145,146,148,172–174

**Adult Dosage.** (See β-Lactams Comparison Chart.)

**Pediatric Dosage.** (See β-Lactams Comparison Chart.)

**Dosage Forms.** (See β-Lactams Comparison Chart.)

**Pharmacokinetics.** Usual half-life is 1–1.5 hr, which is prolonged in anuria, although acylureido derivatives are partially metabolized and accumulate to a lesser extent. The acylureidopenicillins are also subject to capacity-limited elimination (ie, increasing dosage results in progressive saturation of elimination pathways, resulting in decreased clearance), which allows administration of higher doses at 6- to 8-hr intervals.

**Adverse Reactions.** Adverse effects are similar to those of other penicillins. Sodium content of the usual daily dosage of parenteral ticarcillin approaches the equivalent of 1 L of NS. Prolonged bleeding time can occur as a result of binding to platelets and prevention of platelet aggregation.

**IMIPENEM AND CILASTATIN SODIUM**

**Pharmacology.** Imipenem is a carbapenem with an extremely broad spectrum of activity against many aerobic and anaerobic Gram-positive and Gram-negative bacterial pathogens. The commercial preparation contains an equal amount of cilastatin, a renal dehydropeptidase inhibitor that has no antimicrobial activity but prevents imipenem’s metabolism by proximal tubular kidney cells, thus increasing urinary imipenem concentrations and possibly decreasing nephrotoxicity.152,174,175 (See Notes.)

**Administration and Adult Dosage.** IV 1–4 g/day in 3 or 4 divided doses (usually 500 mg q 6–8 hr). For severe, life-threatening infections, a dose of 1 g q 6 hr is recommended (not to exceed 50 mg/kg/day or 4 g/day, whichever is less).174 Infuse 250–500 mg doses over 20–30 min and 1 g doses over 40–60 min; reduce infusion rate if nausea and/or vomiting develops. IM 500–750 mg q 12 hr.

**Special Populations.** Pediatric Dosage. (<1 week) 25 mg/kg q 12 hr; (1–4 weeks) 25 mg/kg q 8 hr; (4 weeks–3 months) 25 mg/kg q 6 hr; (3 months–3 yr) 25 mg/kg q 6 hr; (>3 yr) 15 mg/kg q 6 hr.174,176

**Geriatric Dosage.** Same as adult dosage but adjust for age-related reduction in renal function.

**Other Conditions.** Reduce dosage with renal insufficiency as follows: Clcr 30–70 mL/min, give 75% of the usual dosage; Clcr 20–30 mL/min, give 50% of
the usual dosage; Clcr <20 mL/min, give 25% of the usual dosage. Give a supplemental dose after hemodialysis.\textsuperscript{174}

**Dosage Forms.** **Inj (IV)** 250 mg imipenem/250 mg cilastatin, 500 mg imipenem/500 mg cilastatin; **Inj (Susp, IM only)** 500 mg imipenem/500 mg cilastatin, 750 mg imipenem/750 mg cilastatin. (See Notes.)

**Pharmacokinetics.** **Fate.** Peak serum imipenem concentrations are 21–58 mg/L after a 30-min infusion of 500 mg and 1–84 mg/L after a 30-min infusion of 1 g; levels are <1 mg/L at 6 hr. CSF levels are 0.5–11 mg/L with inflamed meninges and appear to be adequate to treat meningitis, but experience in treating meningitis is limited and seizures can occur in such patients. Imipenem is 20% plasma protein bound; $V_d$ is 0.26 L/kg. Probenecid increases imipenem serum levels and prolongs its half-life. About 70% of imipenem is excreted unchanged in urine when given with cilastatin, with the remaining excreted as metabolite; cilastatin is excreted 90% unchanged in urine.\textsuperscript{174,175}

$\frac{1}{2}$. (Imipenem) 0.9 ± 0.1 hr; 3–4 hr in renal failure; (cilastatin) 0.8 ± 0.1 hr; 17 hr in renal failure.\textsuperscript{174,175}

**Adverse Reactions.** Nausea and vomiting occur in 1–2% of patients, sometimes associated with hypotension or diaphoresis, particularly with high doses and rapid infusion.\textsuperscript{174,175} Rashes occur occasionally, and cross-allergenicity with penicillins has been documented. Convulsions have occurred, primarily in the elderly, in those with underlying CNS disease, with overdosage in patients with renal failure, or with other predisposing factors.\textsuperscript{174,175,177,178}

**Precautions.** Use with caution in elderly patients or those with a history of seizures or who are otherwise predisposed. Adjust dosage carefully in renal impairment. Imipenem can cause immediate hypersensitivity reactions in patients with a history of anaphylaxis to penicillin.\textsuperscript{178}

**Drug Interactions.** Concomitant administration with probenecid produces higher and prolonged serum concentrations of imipenem and cilastatin. Imipenem has been shown in vitro to antagonize the activity of other $\beta$-lactams (eg, acylureido-penicillins, most cephalosporins) presumably via $\beta$-lactamase induction; although the clinical relevance is unclear, avoid co-administration.\textsuperscript{175} Co-administration of imipenem/cilastatin with ganciclovir has been associated with generalized seizures in a few patients; the mechanism of this interaction is unknown.

**Parameters to Monitor.** Obtain renal function tests periodically.

**Notes.** Used alone, emergence of resistance during treatment of *Pseudomonas aeruginosa* infections occurs frequently; however, cross-resistance to other classes (eg, aminoglycosides, cephalosporins) does not occur.\textsuperscript{174,175} Addition of an aminoglycoside might prevent development of resistance, but in vitro synergism occurs only infrequently.

Vials may be reconstituted into a suspension using 10 mL of the infusion solution and then diluted further by transferring the suspension into the infusion container; alternatively, the powder in the 120-mL vials can be diluted initially with 100 mL of solution. The initial dilution must be shaken well to ensure suspension/solution. Do not inject the suspension. The resulting solution ranges from
colorless to yellow. Reconstituted solutions are stable in dextrose-containing solutions for 4 hr at room temperature and 24 hr under refrigeration, and in normal saline for 10 hr at room temperature and 48 hr under refrigeration. With IM administration use 2 mL of lidocaine 1% injection to reconstitute a 500 mg vial and give the suspension by deep IM injection into a large muscle mass (eg, gluteal muscle).175

Pharmacology. Meropenem is a carbapenem with a mechanism of action similar to that of imipenem. Unlike imipenem, meropenem is not appreciably degraded by renal dehydropeptidase-I and thus does not require concomitant administration of a dehydropeptidase inhibitor.175,179,180 (See Notes.)

Administration and Adult Dosage. IV for less severe infections 500 mg–1 g q 8–12 hr; IV for severe or life-threatening infections (eg, meningitis) 2 g q 8 hr.

Special Populations. Pediatric Dosage. IV (<3 months) safety and efficacy not established, but 20 mg/kg q 12 hr has been used; (3 months–12 yr) 10–20 mg/kg q 8 hr; in meningitis 40 mg/kg q 8 hr has been used.179

Geriatric Dosage. Same as adult dosage but adjust for age-related reduction in renal function.

Other Conditions. Reduce dosage in renal impairment. With a Clcr of 26–50 mL/min, give the normal dose q 12 hr; with Clcr of 11–25 mL/min, the dosage is reduced by 50%; with Clcr <10 mL/min, give one-half the dose once daily.179,181

Dosage Forms. Inj 500 mg, 1 g.

Pharmacokinetics. Fate. The pharmacokinetics of meropenem are similar to those of imipenem, although meropenem can be given by IV infusion and bolus.177,179 After IV infusion of 1 g, the peak serum concentration is 39–68 mg/L; the drug distributes well into most tissues and fluids, including the CSF. Plasma protein binding is low and the Vdss is 0.32 ± 0.03 L/kg. Meropenem is primarily eliminated renally by glomerular filtration and tubular secretion. Up to 70% of a dose is recovered unchanged in the urine, with a renal metabolite accounting for the remainder of the dose (up to 30%). Meropenem is appreciably removed by hemodialysis, and a supplemental dose is required after dialysis. Children have pharmacokinetics similar to adults; increased clearance and reduced half-life occur in cystic fibrosis.181

\[ t_{1/2} = 0.9 ± 0.09 \text{ hr}, \] increasing to 6.8–13.7 hr in end-stage renal disease.181

Adverse Reactions. Adverse effects are similar to imipenem; the most common are injection-site reactions, rash, nausea, vomiting, and diarrhea.175,179 Animal studies suggest that meropenem has a lower epileptogenic potential, which has been supported by a low frequency of seizures in clinical trials, including studies in patients with meningitis.179

Precautions. Use with caution in patients with hypersensitivity to penicillins because meropenem can cause immediate hypersensitivity reactions in patients allergic to penicillins.178 Adjust dosage in renal impairment.
Drug Interactions. Probenecid can reduce renal clearance of meropenem and increase its half-life by 38% and AUC by 56%; avoid the combination.

Parameters to Monitor. Obtain renal function tests periodically.

Notes. Meropenem is more active than imipenem against enteric Gram-negative bacilli; the two have equivalent activity against *Pseudomonas aeruginosa* and *Bacteroides fragilis*, and meropenem is slightly less active than imipenem against Gram-positive organisms.¹⁷⁸,¹⁷⁹

**PENICILLIN G AND V SALTS**

Pharmacology. Penicillins G and V have activity against most Gram-positive organisms and some Gram-negative organisms, notably *Neisseria* sp, by interfering with late stages of bacterial cell wall synthesis; resistance is caused primarily by bacterial elaboration of β-lactamases; some organisms have altered penicillin-binding protein targets (eg, enterococci and pneumococci); others have impermeable outer cell wall layers.¹⁴⁵,¹⁴⁶

Administration and Adult Dosage. **PO** (penicillin V) 125–500 mg q 6-8 hr for mild to moderate infections. **IV** (penicillin G) 2–5 million units q 4-6 hr to a maximum of 24 million units/day, depending on infection. **IM** not recommended (very painful); use benzathine or procaine penicillin G as indicated.

Special Populations. Pediatric Dosage. **PO** (penicillin V) (<12 yr) 15–50 mg/kg/day in 3–4 divided doses; (>12 yr) same as adult dosage. **IV (preferably)** or **IM** (penicillin G) (<1 month) 25,000–50,000 units/kg q 6–12 hr; up to 400,000 units/kg/day has been used in meningitis; (>1 month) 100,000–300,000 units/kg/day in 4–6 divided doses.

Geriatric Dosage. Same as adult dosage but adjust for age-related reduction in renal function.

Other Conditions. With the usual oral dosage, no dosage adjustment is required in patients with impaired renal function; however, in treating more severe infections with larger IV dosages, careful adjustment is necessary.¹⁸²

Dosage Forms. (Penicillin G) **Inj** (as potassium salt) 1, 5, 10, 20 million units; **Inj** 1, 2, 3 million units/50 mL (frozen); **Inj** (as sodium salt) 5 million units. (Penicillin V) **Susp** 25, 50 mg/mL; **Tab** 125, 250, 500 mg (250 mg = 400,000 units).

Patient Instructions. Take this (oral) drug with a full glass of water on an empty stomach (1 hour before or 2 hours after meals) for best absorption; refrigerate solution.

Missed Doses. Take this drug at regular intervals. If you miss a dose, take it as soon as you remember. If it is about time for the next dose, take that dose only. Leave at least 4–6 hours between doses. Do not double the dose or take extra.

Pharmacokinetics. **Fate.** (Penicillin G) A peak of 20 mg/L is achieved with a dose of 12 million units IV. Widely distributed in body tissues, fluids, and cavities, with biliary levels up to 10 times serum levels; 45–68% plasma protein bound. Penetration into CSF is poor, even with inflamed meninges; however, large parenteral dosages (>20 million units/day) adequately treat meningitis
caused by susceptible organisms. (Penicillin V) Oral absorption is 60–73%, with a peak concentration of 5–6 mg/L after a 500 mg oral dose. It is about 80% plasma protein bound and has poor CNS penetration. For both drugs, 80–85% of the absorbed dose is excreted unchanged in the urine.\textsuperscript{145,146}

\[ t_{1/2} \text{ (Penicillins G and V)} \text{ 30–40 min; 7–10 hr in patients with renal failure; 20–30 hr in patients with hepatic and renal failure.} \text{182} \]

**Adverse Reactions.** Occasionally, nausea or diarrhea occurs after usual oral doses. As with all penicillins, CNS toxicity can occur with massive IV dosages (penicillin G 60–100 million units/day) or excessive dosage in patients with impaired renal function (usually >10–20 million units/day of penicillin G in anuric patients); characterized by confusion, drowsiness, and myoclonus, which can progress to convulsions and result in death. Large dosages of the sodium salt form can result in hypernatremia and fluid overload with pulmonary edema, especially in patients with impaired renal function or CHF. Large dosages of the potassium salt form can result in hyperkalemia, especially in patients with impaired renal function and with rapid infusions. Occasional positive Coombs’ reactions with rare hemolytic anemia have been reported after large IV doses. Interstitial nephritis has been rarely reported after large IV dosages. Hypersensitivity reactions (primarily rashes) occur in 1–10% of patients. Most serious hypersensitivity reactions follow injection rather than oral administration.\textsuperscript{145,178}

**Contraindications.** History of anaphylactic, accelerated (eg, hives), or serum sickness reaction to previous penicillin administration. (See Notes.)

**Precautions.** Use caution in patients with a history of penicillin or cephalosporin hypersensitivity reactions, atopic predisposition (eg, asthma), impaired renal function (hence neonates and geriatric patients), impaired cardiac function, or pre-existing seizure disorder.

**Drug Interactions.** Physically and/or chemically incompatible with aminoglycosides leading to drug inactivation; never mix them together in the same IV solution or syringe. Probencid competes with penicillin for renal excretion, resulting in higher and prolonged serum concentrations.\textsuperscript{145,146}

**Parameters to Monitor.** Obtain renal function tests initially when using high dosages. During prolonged high-dose therapy, monitor renal function tests and serum electrolytes periodically.

**Notes.** Skin testing with penicilloypolylysine (PPL, Pre-Pen) and minor determinant mixture (MDM) can help determine the likelihood of serious reactions to penicillin in penicillin-allergic individuals.\textsuperscript{145,183} Availability of MDM is limited; it is locally available in small amounts only at larger medical centers. Desensitization is recommended in pregnant women with syphilis and may be attempted (rarely) in patients with life-threatening infections that are likely to be responsive only to penicillin, but this is a dangerous procedure and many alternative antibiotics are available.\textsuperscript{145} (See also β-Lactams Comparison Chart.)
| CARBAPENEMS |  |  |  |
|-------------|-----------------------------|-------------------|-----------------|-----------------|-----------------|-----------------|-----------------|
| **Imipenem** | **Inj (IV) 250 plus 250 mg,** | **IV 1–4 g/day (1–2 g/day preferred) in 3 or 4 divided doses; IM 500–750 mg q 12 hr.** | **(<3 months) See monograph; IV (3 mo–3 yr) 25 mg/kg q 6 hr; (>3 yr) 15 mg/kg q 6 hr.** | **Clr 31–70 mL/min:** 75% of usual dosage; **Clr <20–30 mL/min:** 50% of usual dosage; **Clr <20 mL/min:** 25% of usual dosage. | **21–58 (IV 500 mg imipenem)** | **20** | **Very broad activity against most aerobic and anaerobic bacteria. Frequent nausea and dose-related seizure potential.** |
| and Cila-statin Sodium Primaxin | **500 mg; (IM) 500 plus 500 mg,** | **IM 500–750 mg q 12 hr.** | **(<3 months) See monograph; IV (3 mo–3 yr) 25 mg/kg q 6 hr; (>3 yr) 15 mg/kg q 6 hr.** | **Clr 31–70 mL/min:** 75% of usual dosage; **Clr <20–30 mL/min:** 50% of usual dosage; **Clr <20 mL/min:** 25% of usual dosage. | **21–58 (IV 500 mg imipenem)** | **20** | **Very broad activity against most aerobic and anaerobic bacteria. Frequent nausea and dose-related seizure potential.** |
| **Meropenem** | **Inj 500 mg, 1 g.** | **IV 500 mg–1 g q 8–12 hr; 2 g q 8 hr in life-threatening infections.** | **(<3 months) See monograph; IV (3 mo–12 yr) 10–20 mg/kg q 8 hr; 40 mg/kg q 8 hr in meningitis.** | **Clr 26–50 mL/min:** usual dose q 12 hr; **Clr 11–25 mL/min:** 50% of usual dose q 12 hr; **Clr <10 mL/min:** 50% of usual dose q 24 hr. | **55** | **2** | **Less active than imipenem against Gm+ and more active against most Gm– bacteria; equivalent against P. aeruginosa and B. fragilis. Less seizure potential than imipenem.** |
| **Merrem** | **Inj 500 mg, 1 g.** | **IV 500 mg–1 g q 8–12 hr; 2 g q 8 hr in life-threatening infections.** | **(<3 months) See monograph; IV (3 mo–12 yr) 10–20 mg/kg q 8 hr; 40 mg/kg q 8 hr in meningitis.** | **Clr 26–50 mL/min:** usual dose q 12 hr; **Clr 11–25 mL/min:** 50% of usual dose q 12 hr; **Clr <10 mL/min:** 50% of usual dose q 24 hr. | **55** | **2** | **Less active than imipenem against Gm+ and more active against most Gm– bacteria; equivalent against P. aeruginosa and B. fragilis. Less seizure potential than imipenem.** |

(continued)
<table>
<thead>
<tr>
<th>DRUG CLASS AND DRUG</th>
<th>DOSAGE FORMS</th>
<th>ADULT DOSAGE</th>
<th>PEDIATRIC DOSAGE</th>
<th>ADULT DOSAGE IN RENAL IMPAIRMENT^a</th>
<th>PEAK SERUM LEVELS (MG/L)^b</th>
<th>PERCENTAGE PROTEIN BOUND</th>
<th>COMMENTS</th>
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<tbody>
<tr>
<td><strong>CEPHALOSPORINS, FIRST-GENERATION</strong></td>
<td></td>
<td></td>
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<td></td>
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</tr>
<tr>
<td>Cefadroxil Duricef Various</td>
<td>Cap 500 mg Susp 25, 50, 100 mg/mL Tab 1 g.</td>
<td>PO 1–2 g/day in 1 or 2 divided doses; PO for endocarditis prophylaxis 2 g 1 hr prior to dental procedure.</td>
<td>PO 30 mg/kg/day in 1 or 2 divided doses; PO for endocarditis prophylaxis 50 mg/kg 1 hr prior to dental procedure.</td>
<td>PO 1 g, then 500 mg at intervals below: Cl&lt;sub&gt;r&lt;/sub&gt; 26–50 mL/min: 12 hr; Cl&lt;sub&gt;r&lt;/sub&gt; 10–25 mL/min: 24 hr; Cl&lt;sub&gt;r&lt;/sub&gt; &lt;10 mL/min: 36 hr.</td>
<td>12–16</td>
<td>20</td>
<td>Spectrum similar to cefazolin.</td>
</tr>
<tr>
<td><strong>Cefazolin Sodium</strong></td>
<td>Inj 250, 500 mg, 1, 5, 10, 20 g.</td>
<td>IM or IV 250 mg–2 g q 6–12 hr; (usually q 8 hr), to a maximum of 12 g/day. IM or IV for surgical prophylaxis: 1g 30–60 min prior to surgery; IM or IV for endocarditis prophylaxis 1g within 30 min prior to upper airway procedure.</td>
<td>IM or IV (neonates &lt;1 month) 25 mg/kg q 8–12 hr; (infants &gt;1 month) 50–100 mg/kg/day in 3 divided doses to a maximum of 6 g/day. IM or IV for endocarditis prophylaxis 25 mg/kg within 30 min of procedure.</td>
<td>Cl&lt;sub&gt;r&lt;/sub&gt; 10–30 mL/min: 50% of usual dose q 12 hr; Cl&lt;sub&gt;r&lt;/sub&gt; &lt;10 mL/min: 50% of usual dose q 24 hr.</td>
<td>185</td>
<td>75–85</td>
<td>Good Gm+ coverage (including S. aureus), plus some Gm– activity (E. coli, Klebsiella spp.). Sodium = 2 mEq/g</td>
</tr>
</tbody>
</table>

(continued)
### β-LACTAMS COMPARISON CHART (continued)

<table>
<thead>
<tr>
<th>DRUG CLASS AND DRUG</th>
<th>DOSAGE FORMS</th>
<th>ADULT DOSAGE</th>
<th>PEDIATRIC DOSAGE</th>
<th>ADULT DOSAGE IN RENAL IMPAIRMENT&lt;sup&gt;a&lt;/sup&gt;</th>
<th>PEAK SERUM LEVELS (MG/L)&lt;sup&gt;b&lt;/sup&gt;</th>
<th>PERCENTAGE PROTEIN BOUND</th>
<th>COMMENTS</th>
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</thead>
<tbody>
<tr>
<td><strong>Cephalexin</strong></td>
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<tr>
<td>Kellex</td>
<td>Cap 250, 500 mg</td>
<td>PO 250 mg–1 g q 6 hr; to a maximum of 4 g/day.</td>
<td>PO 25–50 mg/kg/day in divided doses q 6 hr; severe infections may require 50–100 mg/kg/day, to a maximum of 3 g/day.</td>
<td>C&lt;sub&gt;r&lt;/sub&gt; 10–50 mL/min: 18–38</td>
<td>6</td>
<td>Oral absorption is almost complete; spectrum similar to cefazolin.</td>
<td></td>
</tr>
<tr>
<td>Keftab</td>
<td>Drp 100 mg/mL</td>
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</tr>
<tr>
<td>Various</td>
<td>Susp 25, 50 mg/mL</td>
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<tr>
<td></td>
<td>Tab 250, 500 mg, 1 g.</td>
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<tr>
<td><strong>Cephapirin Sodium</strong></td>
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<tr>
<td>Cefadyl</td>
<td>Inj 1 g.</td>
<td>IM or IV 500 mg–1 g q 4–6 hr, to a maximum of 12 g/day.</td>
<td>IM or IV (&lt;3 months) not well studied; (children) 40–80 mg/kg/day in divided doses q 6 hr.</td>
<td>C&lt;sub&gt;r&lt;/sub&gt; 10–50 mL/min; usual dose q 6–8 hr</td>
<td>10–20</td>
<td>45–50</td>
<td>Spectrum similar to cefazolin. Sodium = 1.2 mEq/g.</td>
</tr>
<tr>
<td>Various</td>
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</tbody>
</table>

<sup>a</sup> Cl<sub>r</sub> = creatinine clearance in mL/min.

<sup>b</sup> Serum levels are for adult patients unless otherwise noted.
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<th>PEAK SERUM LEVELS (MG/L)&lt;sup&gt;b&lt;/sup&gt;</th>
<th>PERCENTAGE PROTEIN BOUND</th>
<th>COMMENTS</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Cephradine</strong></td>
<td>Cap 250, 500 mg</td>
<td>PO 250 mg–1 g q 6 hr to a maximum of 4 g/day.</td>
<td>PO same as cephalexin.</td>
<td>PO same as cephalexin.</td>
<td>10–20 (PO)</td>
<td>10–20</td>
<td>Oral form comparable to cephalexin; spectrum similar to cefazolin.</td>
</tr>
<tr>
<td>Velosef</td>
<td>Susp 25, 50 mg/mL</td>
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</tr>
<tr>
<td>Various</td>
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</tbody>
</table>

**CEPHALOSPORINS, SECOND-GENERATION**

| Cefaclor | Cap 250, 500 mg | PO 250–500 mg q 8 hr | PO 20–40 mg/kg/day in divided doses q 8 hr to a maximum of 2 g/day. | Cl<sub>cr</sub> 10–50 mL/min: 50% of usual dosage; Cl<sub>cr</sub> <10 mL/min: 25% of usual dosage. | 10 | 25 | Spectrum similar to cefazolin, but includes some ampicillin-resistant H. influenzae. |
| Ceclor | SR Tab 375, 500 mg | SR Tab 375–500 mg q 12 hr. | | | | | |
| Various | Susp 25, 37.5, 50, 75 mg/mL | | | | | | |

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<table>
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<th>PEDIATRIC DOSAGE</th>
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<th>PEAK SERUM LEVELS (MG/L)&lt;sup&gt;b&lt;/sup&gt;</th>
<th>PERCENTAGE PROTEIN BOUND</th>
<th>COMMENTS</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cefaman-dole</td>
<td>Inj 1, 2, g.</td>
<td>IM or IV 500 mg–1 g q 4–8 hr.; life threatening infections may require 2 g q 4 hr.</td>
<td>IM or IV 50–150 mg/kg/day in divided doses q 4–8 hr.</td>
<td>Cl&lt;sub&gt;r&lt;/sub&gt; 10–50 mL/min: 50% of usual dose q 8 hr; Cl&lt;sub&gt;r&lt;/sub&gt; &lt;10 mL/min: 25% of usual dose q 12 hr.</td>
<td>80–90</td>
<td>56</td>
<td>NMTT side chain. Spectrum similar to cefuroxime. Sodium = 3.3 mEq/g.</td>
</tr>
<tr>
<td>Nafate</td>
<td>Disodium</td>
<td>Sodium</td>
<td>Mandol</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Cefonicid</td>
<td>Inj 1 g.</td>
<td>IM or IV 500 mg–2 g/day as a single dose.</td>
<td>Not established.</td>
<td>IM or IV 7.5 mg/kg, then 25–50% of usual dose given: Cl&lt;sub&gt;r&lt;/sub&gt; 10–50 mL/min: q 24–48 hr; Cl&lt;sub&gt;r&lt;/sub&gt; &lt;10 mL/min: q 3–5 days.</td>
<td>220</td>
<td>83–98&lt;sup&gt;c&lt;/sup&gt;</td>
<td>Poor activity against Staphylococcus spp. Unbound drug levels low and excreted rapidly because of saturable protein binding. Sodium = 3.7 mEq/g.</td>
</tr>
<tr>
<td>Sodium</td>
<td>Monocid</td>
<td></td>
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<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Cefotetan</td>
<td>Inj 1, 2, 10 g.</td>
<td>IM or IV 500 mg–2 g q 12–24 hr.</td>
<td>Not established.</td>
<td>IM or IV give usual dose at intervals below: Cl&lt;sub&gt;r&lt;/sub&gt; 10–30 mL/min: q 24 hr; Cl&lt;sub&gt;r&lt;/sub&gt; &lt;10 mL/min: q 48 hr.</td>
<td>140–180</td>
<td>78–91&lt;sup&gt;c&lt;/sup&gt;</td>
<td>NMTT side chain. Spectrum similar to cefoxitin. Sodium = 3.5 mEq/g. Reconstitute IM with 0.5% lidocaine.</td>
</tr>
<tr>
<td>Disodium</td>
<td>Cefotan</td>
<td></td>
<td></td>
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<td></td>
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</tr>
</tbody>
</table>

<sup>a</sup> Calculated using standard formulas for renal impairment. <br>
<sup>b</sup> Serum concentrations are approximate. <br>
<sup>c</sup> Values may vary depending on the specific drug and clinical scenario.
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<th>Drug Class and Drug</th>
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<th>Pediatric Dosage</th>
<th>Adult Dosage in Renal Impairment&lt;sup&gt;a&lt;/sup&gt;</th>
<th>Peak Serum Levels (mg/L)&lt;sup&gt;b&lt;/sup&gt;</th>
<th>Percentage Protein Bound</th>
<th>Comments</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cefoxitin Sodium</td>
<td>Inj 1, 2, 10 g.</td>
<td>IV 1–2 g q 6–8 hr.</td>
<td>IV 80–160 mg/kg/day in divided doses q 4–6 hr.</td>
<td>$C_L &lt; 10$ mL/min: 25% of usual dose q 12 hr.</td>
<td>110</td>
<td>75</td>
<td>Gm+ activity less than cefazolin, but better Gm− and anaerobic activity. Sodium = 2.3 mEq/g.</td>
</tr>
<tr>
<td>Mefoxin</td>
<td>Tab 250, 500 mg</td>
<td>PO 500 mg daily–bid.</td>
<td>PO (6 mo–12 yr) 15 mg/kg q 12 hr.</td>
<td>$C_L &lt; 30$ mL/min: 50% of usual dose at same interval.</td>
<td>10.5</td>
<td>36</td>
<td>Spectrum similar to cefaclor, but more active against H. influenzae.</td>
</tr>
<tr>
<td>Cefuroxime Sodium</td>
<td>Inj 750 mg, 1.5 g; IM or IV 750 mg–1.5 g q 6–8 hr; to a maximum of 6 g/day.</td>
<td>IM or IV (neonates) 10–25 mg/kg q 12 hr; (children) 50–100 mg/kg/day in divided doses q 6–8 hr.</td>
<td>IM or IV: $C_L &lt; 10$ mL/min: usual dose q 12 hr.</td>
<td>100 (IV 1.5 g)</td>
<td>33–50</td>
<td>Gm+ activity similar to cefazolin, but better Gm− activity, including H. influenzae. Sodium = 2.4 mEq/g.</td>
<td></td>
</tr>
<tr>
<td>Kefurox Susp 25, 50 mg/mL Tab 125, 250, 500 mg.</td>
<td>PO 125–500 mg q 12 hr.</td>
<td>PO 15–40 mg/kg/day in divided doses q 12 hr.</td>
<td>—</td>
<td>3.6 (PO)</td>
<td>33–50</td>
<td>Do not interchange suspension and tablets.</td>
<td></td>
</tr>
</tbody>
</table>

<sup>a</sup> Renal impairment levels: 10–50 mL/min: 50% of usual dose q 6–8 hr; $C_L < 10$ mL/min: 25% of usual dose q 12 hr.

<sup>b</sup> Sodium = 2.3 mEq/g.
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<tr>
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<th>PEAK SERUM LEVELS (MG/L)&lt;sup&gt;b&lt;/sup&gt;</th>
<th>PERCENTAGE PROTEIN BOUND</th>
<th>COMMENTS</th>
</tr>
</thead>
<tbody>
<tr>
<td>Loracarbef</td>
<td>Cap 200, 400 mg/mL.</td>
<td>PO 200–400 mg q 12–24 hr.</td>
<td>PO (6 mo–12 yr) 7.5–15 mg q 12 hr.</td>
<td>Cl&lt;sub&gt;r&lt;/sub&gt; 10–49 mL/min: 50% of usual dosage; Cl&lt;sub&gt;r&lt;/sub&gt; &lt;10 mL/min: usual dose q 3–5 days.</td>
<td>6.8 (PO 200 mg)</td>
<td>25</td>
<td>Carbacephem analogue of cefaclor with similar spectrum; must be taken on an empty stomach.</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>CEPHALOSPORINS, THIRD-GENERATION</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Cefdinir</strong></td>
</tr>
<tr>
<td>Omnicef</td>
</tr>
<tr>
<td>Tab 200 mg.</td>
</tr>
<tr>
<td>PO 200 q 12 hr.</td>
</tr>
<tr>
<td>PO (1 or 2 doses)</td>
</tr>
<tr>
<td>Cl&lt;sub&gt;r&lt;/sub&gt; &lt;30 mL/min: 300 mg/day.</td>
</tr>
</tbody>
</table>

| **Cefditoren**                   |
| Spectracef                       |
| Tab 200 mg.                      |
| PO 200 q 12 hr.                  |
| PO (<12 yr) not established.     |
| Cl<sub>r</sub> <50 mL/min: reduce dosage. | 2.6 | 88 | Spectrum similar to cefdinir and cefpodoxime but more active. |

| **Cefixime**                     |
| Suprax                           |
| Tab 200, 400 mg/mL.              |
| PO 400 mg/day in 1 or 2 doses.   |
| PO for gonorrhea 400 mg once.    |
| Cl<sub>r</sub> 20–60 mL/min: 75% of usual dosage; Cl<sub>r</sub> <20 mL/min: 50% of usual dosage. | 4.9 | 70 | More active than cefuroxime or cefaclor against *H. influenzae*, but less Gm+ activity. |
### β-LACTAMS COMPARISON CHART (continued)

<table>
<thead>
<tr>
<th>DRUG CLASS AND DRUG</th>
<th>DOSAGE FORMS</th>
<th>ADULT DOSAGE</th>
<th>PEDIATRIC DOSAGE</th>
<th>ADULT DOSAGE IN RENAL IMPAIRMENT *</th>
<th>PEAK SERUM LEVELS (MG/L) *</th>
<th>PERCENTAGE PROTEIN BOUND</th>
<th>COMMENTS</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cefoperazone Sodium</td>
<td>Inj 1, 2, 10 g.</td>
<td>IM or IV 2–8 g/day in divided doses q 12 hr.</td>
<td>IM or IV (neonates) 50 mg/kg/dose q 12 hr; (children) 50–75 mg/kg q 8–12 hr.</td>
<td>No change.</td>
<td>125</td>
<td>85–95</td>
<td>Less active than ceftazidime against <em>P. aeruginosa</em>. NMTT side chain. Sodium = 1.5 mEq/g.</td>
</tr>
<tr>
<td>Cefobid Various</td>
<td>Inj 500 mg, 1, 2, 10 g.</td>
<td>IM or IV 1–2 g q 8–12 hr; life-threatening infections may require 2 g q 6 hr.</td>
<td>IM or IV (neonates ≤1 week) 50 mg/kg q 12 hr; (neonates 1–4 weeks) 50 mg/kg q 8 hr; (infants &gt;4 weeks) 50–200 (200 in meningitis) mg/kg/day in divided doses q 4–6 hr.</td>
<td>Clr, 10–50 mL/min; usual dose q 8–12 hr; &lt;10 mL/min usual dose q 24 hr.</td>
<td>40–100</td>
<td>37</td>
<td>Good Gm+ and Gm- activity except for <em>P. aeruginosa</em>; modest anti-anaerobic activity. Sodium = 2.2 mEq/g.</td>
</tr>
<tr>
<td>Desacetylcefotaxime</td>
<td>—</td>
<td>—</td>
<td>—</td>
<td>1–65</td>
<td></td>
<td></td>
<td>Active metabolite of cefotaxime.</td>
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</tbody>
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(continued)
### β-LACTAMS COMPARISON CHART (continued)

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<th>PEDIATRIC DOSAGE</th>
<th>ADULT DOSAGE IN RENAL IMPAIRMENT*</th>
<th>PEAK SERUM LEVELS (MG/L)*</th>
<th>PERCENTAGE PROTEIN BOUND</th>
<th>COMMENTS</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cefpodoxime Proxetil Vanitin</td>
<td>Tab 100, 200 mg Susp 10, 20 mg/mL</td>
<td>PO 100–400 mg q 12 hr; PO for gonorrhea 200 mg once.</td>
<td>PO 5 mg/kg q 12 hr.</td>
<td>Clcr ≤30 mL/min: usual dose given q 24 hr.</td>
<td>18–30</td>
<td>2.9 (PO 200 mg)</td>
<td>Spectrum similar to cefixime, but better Gm+ activity.</td>
</tr>
<tr>
<td>Ceftazidime Ceptaz Fortaz Tazicef Tazidime</td>
<td>Inj 500 mg, 1, 2, 6 g.</td>
<td>IM or IV 500 mg–2 g q 8–12 hr.</td>
<td>IM or IV (≤1 month) 30 mg/kg/dose q 12 hr; (&gt;1 month) 30–50 mg/kg/dose q 8 hr.</td>
<td>Clcr 30–50 mL/min: 50% of usual dose q 12–24 hr. Clcr 15–30 mL/min: 1 g q 24 hr. Clcr ≤15 mL/min: 500 mg q 24–48 hr.</td>
<td>17</td>
<td>70–90</td>
<td>Best activity against <em>P. aeruginosa</em>; poor Gm+ activity. Sodium = 2.3 mEq/g.</td>
</tr>
<tr>
<td>Ceftibuten Cedax</td>
<td>Cap 400 mg Susp 18, 36 mg/mL</td>
<td>PO 400 mg q 24 hr.</td>
<td>PO 9 mg/kg/day in 1 dose.</td>
<td>Clcr 30–49 mL/min: 4.5 mg/kg or 200 mg q 24 hr; Clcr ≤30 mL/min: 2.25 mg/kg or 100 mg q 24 hr.</td>
<td>11</td>
<td>60–77</td>
<td>Spectrum similar to cefixime.</td>
</tr>
<tr>
<td><strong>β-LACTAMS COMPARISON CHART (continued)</strong></td>
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<th>PEAK SERUM LEVELS (MG/L)</th>
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<th>COMMENTS</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ceftriaxone Disodium</td>
<td>Inj 250, 500 mg, 1, 2, 10 g.</td>
<td>IM or IV 1–2 g/day as a single dose; IV for meningitis 2 g q 12 hr; IM for gonorrhea 250 mg once.</td>
<td>IM or IV 50–100 (100 in meningitis) mg/kg/day in 2 divided doses.</td>
<td>No change. (See Comments.)</td>
<td>151</td>
<td>83–96°</td>
<td>Spectrum similar to cefotaxime. Reduce dose with concurrent renal and hepatic dysfunction. Sodium = 3.6 mEq/g.</td>
</tr>
<tr>
<td>Sodium Ceftriaxone</td>
<td>Ceftriaxone</td>
<td>Rocephin</td>
<td></td>
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</table>

- Spectrum similar to cefotaxime except slightly more active against anaerobes. Sodium = 2.6 mEq/g.
- Spectrum similar to cefotaxime. Reduce dose with concurrent renal and hepatic dysfunction. Sodium = 3.6 mEq/g.
### β-LACTAMS COMPARISON CHART (continued)

<table>
<thead>
<tr>
<th>DRUG CLASS AND DRUG</th>
<th>DOSAGE FORMS</th>
<th>ADULT DOSAGE</th>
<th>PEDIATRIC DOSAGE</th>
<th>ADULT DOSAGE IN RENAL IMPAIRMENT&lt;sup&gt;a&lt;/sup&gt;</th>
<th>PEAK SERUM LEVELS (MG/L)&lt;sup&gt;b&lt;/sup&gt;</th>
<th>PERCENTAGE PROTEIN BOUND</th>
<th>COMMENTS</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>CEPHALOSPORINS, FOURTH-GENERATION</strong></td>
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<tr>
<td>Cefepime Maxipime</td>
<td>Inj 500 mg, 1, 2 g.</td>
<td>IM or IV 500 mg–2 g q 12 hr; 2 g q 8 hr may be required for pseudomonal infections and febrile neutropenia.</td>
<td>IM or IV (2 months–16 yr) febrile neutropenia 50 mg/kg q 8 hr; other infections 50 mg/kg q 12 hr.</td>
<td>Cl&lt;sub&gt;r&lt;/sub&gt; 30–60 mL/min: usual dose q 24 hr; Cl&lt;sub&gt;r&lt;/sub&gt; 10–29 mL/min: 50% of usual dose q 24 hr; Cl&lt;sub&gt;r&lt;/sub&gt; &lt;10 mL/min: 25% of usual dose q 24 hr.</td>
<td>79</td>
<td>16–19</td>
<td>Spectrum similar to ceftazidime; more active against Gm+ organisms; also active against resistant Enterobacter spp.</td>
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<tr>
<td><strong>MONOBACTAM</strong></td>
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<tr>
<td>Aztreonam Azactam</td>
<td>Inj 500 mg, 1, 2 g.</td>
<td>IM or IV 0.5–2 g q 6–12 hr.</td>
<td>Safety and efficacy not established. IV 30 mg/kg q 6–8 hr (50 mg/kg q 6–8 hr in cystic fibrosis, to a maximum of 200 mg/kg/day).</td>
<td>Cl&lt;sub&gt;r&lt;/sub&gt; 10–30 mL/min: 50% of usual dosage; Cl&lt;sub&gt;r&lt;/sub&gt; &lt;10 mL/min: 25% of usual dosage.</td>
<td>164</td>
<td>60</td>
<td>Spectrum similar to ceftazidime against aerobic Gm− organisms only. No cross-allergenicity in penicillin-allergic patients.</td>
</tr>
<tr>
<td>DRUG CLASS AND DRUG</td>
<td>DOSAGE FORMS</td>
<td>ADULT DOSAGE</td>
<td>PEDIATRIC DOSAGE</td>
<td>ADULT DOSAGE IN RENAL IMPAIRMENT&lt;sup&gt;a&lt;/sup&gt;</td>
<td>PEAK SERUM LEVELS (MG/L)&lt;sup&gt;b&lt;/sup&gt;</td>
<td>PERCENTAGE PROTEIN BOUND</td>
<td>COMMENTS</td>
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<tr>
<td><strong>PENICILLIN G AND V</strong></td>
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<tr>
<td><strong>Penicillin G</strong></td>
<td>Inj 1, 5, 10, 20 million units</td>
<td>IV 2–5 million units q 4–6 hr.</td>
<td>IV (neonates) 25,000–50,000 units/kg q 6–12 hr; (&gt;1 month) 100,000–300,000 units/kg/d in divided doses q 4–6 hr.</td>
<td>Cl&lt;sub&gt;r&lt;/sub&gt; 10–50 mL/min: 75% of dosage; Cl&lt;sub&gt;r&lt;/sub&gt; &lt;10 mL/min: 25–50% of dosage.</td>
<td>1.5–2.7 (IV 500 mg)</td>
<td>60</td>
<td>Gm+ (except most Staphylococcus strains), some Gm– (Neisseria spp.), and anaerobes (except B. fragilis). Poor oral absorption. Potassium = 1.7 mEq/ million units.</td>
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<tr>
<td><strong>Potassium</strong></td>
<td>Inj 1, 2, 3 million units/50 mL (frozen)</td>
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<td><strong>Various</strong></td>
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<tr>
<td><strong>Penicillin G Benzathine</strong></td>
<td>Inj 300,000, 600,000, 1.2 million units/mL.</td>
<td>IM for Strep. pharyngitis 1.2 million units once; IM for syphilis (early) 2.4 million units once; (late) 2.4 million units/week for 3 weeks.</td>
<td>IM for Strep. pharyngitis (&lt;27 kg) 300,000–600,000 units once; (&gt;27 kg) 900,000 units once.</td>
<td>No change.</td>
<td>0.063 (IM 600,000 units)</td>
<td>60</td>
<td>Use limited to syphilis and Strep. pharyngitis. For IM use only.</td>
</tr>
</tbody>
</table>

(continued)
<table>
<thead>
<tr>
<th>DRUG CLASS AND DRUG</th>
<th>DOSAGE FORMS</th>
<th>ADULT DOSAGE</th>
<th>PEDIATRIC DOSAGE</th>
<th>ADULT DOSAGE IN RENAL IMPAIRMENT</th>
<th>PEAK SERUM LEVELS (MG/L)</th>
<th>PERCENTAGE PROTEIN BOUND</th>
<th>COMMENTS</th>
</tr>
</thead>
<tbody>
<tr>
<td>Penicillin G</td>
<td>Inj 300,000, 600,000 units/mL.</td>
<td>IM 600,000–2.4 million units q 12–24 hr (0.6–4.8 million units/day divided q 12–24 hr).</td>
<td>IM (neonates) 50,000 units/kg/day in 1–2 divided doses; (&gt;27 kg) 900,000 units once daily.</td>
<td>No change.</td>
<td>0.9 (IM 300,000 units)</td>
<td>60</td>
<td>For IM use only.</td>
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<tr>
<td>Procaine</td>
<td>Various</td>
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<tr>
<td>Penicillin V</td>
<td>Tab 125, 250, 500 mg</td>
<td>PO 125–500 mg q 6–8 hr.</td>
<td>PO 15–50 mg/kg/day in 3–4 divided doses.</td>
<td>No change.</td>
<td>3–8</td>
<td>78</td>
<td>Spectrum similar to penicillin G. About 60% absorbed; preferred oral form of penicillin.</td>
</tr>
<tr>
<td>Potassium</td>
<td>Tab 125, 250, 500 mg</td>
<td>PO 125–500 mg q 6–8 hr.</td>
<td>PO 15–50 mg/kg/day in 3–4 divided doses.</td>
<td>No change.</td>
<td>3–8</td>
<td>78</td>
<td>Spectrum similar to penicillin G. About 60% absorbed; preferred oral form of penicillin.</td>
</tr>
<tr>
<td>Various</td>
<td>Tab 125, 250, 500 mg</td>
<td>PO 125–500 mg q 6–8 hr.</td>
<td>PO 15–50 mg/kg/day in 3–4 divided doses.</td>
<td>No change.</td>
<td>3–8</td>
<td>78</td>
<td>Spectrum similar to penicillin G. About 60% absorbed; preferred oral form of penicillin.</td>
</tr>
</tbody>
</table>

**ANTISTAPHYLOCOCCAL PENICILLINS**

<table>
<thead>
<tr>
<th>DRUG CLASS AND DRUG</th>
<th>DOSAGE FORMS</th>
<th>ADULT DOSAGE</th>
<th>PEDIATRIC DOSAGE</th>
<th>ADULT DOSAGE IN RENAL IMPAIRMENT</th>
<th>PEAK SERUM LEVELS (MG/L)</th>
<th>PERCENTAGE PROTEIN BOUND</th>
<th>COMMENTS</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cloxacin Sodium</td>
<td>Cap 250, 500 mg</td>
<td>PO 250–500 mg q 6 hr.</td>
<td>PO (&lt;20 kg) 50–100 mg/kg/day in divided doses q 6 hr; (&gt;20 kg) same as adult dosage.</td>
<td>No change.</td>
<td>7–18</td>
<td>94</td>
<td>Used primarily for S. aureus infections. Suspension may be better tolerated than dicloxacillin.</td>
</tr>
<tr>
<td>Cloxapen</td>
<td>Susp 25 mg/mL</td>
<td>PO 250–500 mg q 6 hr.</td>
<td>PO (&lt;20 kg) 50–100 mg/kg/day in divided doses q 6 hr; (&gt;20 kg) same as adult dosage.</td>
<td>No change.</td>
<td>7–18</td>
<td>94</td>
<td>Used primarily for S. aureus infections. Suspension may be better tolerated than dicloxacillin.</td>
</tr>
<tr>
<td>Tegopen</td>
<td>Susp 25 mg/mL</td>
<td>PO 250–500 mg q 6 hr.</td>
<td>PO (&lt;20 kg) 50–100 mg/kg/day in divided doses q 6 hr; (&gt;20 kg) same as adult dosage.</td>
<td>No change.</td>
<td>7–18</td>
<td>94</td>
<td>Used primarily for S. aureus infections. Suspension may be better tolerated than dicloxacillin.</td>
</tr>
<tr>
<td>Various</td>
<td>Tab 125, 250, 500 mg</td>
<td>PO 125–500 mg q 6–8 hr.</td>
<td>PO 15–50 mg/kg/day in 3–4 divided doses.</td>
<td>No change.</td>
<td>3–8</td>
<td>78</td>
<td>Spectrum similar to penicillin G. About 60% absorbed; preferred oral form of penicillin.</td>
</tr>
</tbody>
</table>

(continued)
### β-LACTAMS COMPARISON CHART (continued)

<table>
<thead>
<tr>
<th>DRUG CLASS AND DRUG</th>
<th>DOSAGE FORMS</th>
<th>ADULT DOSAGE</th>
<th>PEDIATRIC DOSAGE</th>
<th>ADULT DOSAGE IN RENAL IMPAIRMENT*</th>
<th>PEAK SERUM LEVELS (MG/L)³</th>
<th>PERCENTAGE PROTEIN BOUND</th>
<th>COMMENTS</th>
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<tbody>
<tr>
<td><strong>Dicloxacillin</strong></td>
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<td>Sodium</td>
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<td>Dynapen</td>
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<td>Pathocillin</td>
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<tr>
<td>Various</td>
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<tr>
<td></td>
<td>Cap 125, 250,</td>
<td>PO 125–500 mg q</td>
<td>PO 12.5–25 mg/</td>
<td>No change.</td>
<td>7–18</td>
<td>98</td>
<td>Comparable to clox-</td>
</tr>
<tr>
<td></td>
<td>500 mg</td>
<td>q 6 hr.</td>
<td>kg/day in divided</td>
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<td>acillin.</td>
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<td></td>
<td>Susp 12.5 mg/mL</td>
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<td>doses q 6 hr.</td>
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<td><strong>Nafcillin</strong></td>
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<td>Sodium</td>
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<td></td>
<td>Cap 250 mg</td>
<td>IV 500 mg–2 g</td>
<td>IV (neonates &lt;7 days) 25 mg/kg q</td>
<td>No change.</td>
<td>3.4 (PO)</td>
<td>89</td>
<td>Comparable to ox-</td>
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<tr>
<td></td>
<td>Inj 500 mg,</td>
<td>q 4–6 hr.</td>
<td>8–12 hr; (neonates &gt;7 days) 25 mg/</td>
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<td>acillin. Revers-</td>
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<td>1, 2, 4, 10 g.</td>
<td>PO 250 mg–1 g</td>
<td>kg q 6–8 hr.</td>
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<td>ible neutropenia may</td>
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<td>q 4–6 hr.</td>
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<td>be more common with</td>
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<td>Sodium</td>
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<td>nafcillin. Poorly</td>
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<td>Unipen</td>
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<td>absorbed orally;</td>
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<td>cloxacillin or</td>
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<td>dicoxacillin prefe-</td>
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<td>rred. N sodium = 2.9 mEq/g.</td>
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<tr>
<td>DRUG CLASS AND DRUG</td>
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<td>ADULT DOSAGE</td>
<td>PEDIATRIC DOSAGE</td>
<td>ADULT DOSAGE IN RENAL IMPAIRMENT</td>
<td>PEAK SERUM LEVELS (MG/L)</td>
<td>PERCENTAGE PROTEIN BOUND</td>
<td>COMMENTS</td>
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<tr>
<td><strong>Oxacillin Sodium</strong></td>
<td>Cap 250, 500 mg Susp 50 mg/mL</td>
<td>IV 250 mg–2 g q 4–6 hr; PO 500 mg–1 g q 6 hr, but not recommended.</td>
<td>IV (≤14 days) 25 mg/kg q 8–12 hr; (15–30 days) 25 mg/kg q 6 hr; (children) same as adult dosage. PO 50–100 mg/kg/day in 4–6 divided doses.</td>
<td>No change.</td>
<td>2.5 (PO) 40 (IV)</td>
<td>92</td>
<td>Poorly absorbed orally; cloxacillin or dicloxacillin preferred. Rare hepatic toxicity. IV sodium = 2.9 mEq/g.</td>
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<tr>
<td><strong>Bactocill</strong></td>
<td>Inj 250, 500 mg</td>
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<tr>
<td><strong>Prostaphlin</strong></td>
<td>1, 2, 4, 10 g.</td>
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<tr>
<td><strong>AMPICILLIN DERIVATIVES</strong></td>
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<tr>
<td><strong>Amoxicillin Amoxil</strong></td>
<td>Cap 250, 500 mg Chew Tab 125, 200, 250, 400 mg Dp 50 mg/mL Susp 25, 50 mg/mL Tab 500, 875 mg.</td>
<td>PO 250–500 mg tid, or 500–875 mg bid, to a maximum of 4.5 g/day; PO for endocarditis prophylaxis 2 g 1 hr before procedure.</td>
<td>PO 20–40 mg/kg/day in 3 divided doses; PO for endocarditis prophylaxis 50 mg/kg 1 hr before procedure.</td>
<td>Cl <em>p</em>, 10–30 mL/min: 250–500 mg bid; Cl <em>p</em>, &lt;10 mL/min: 250–500 mg q 24 hr.</td>
<td>9</td>
<td>20</td>
<td>Spectrum similar to ampicillin, but better bioavailability (85%) and less diarrhea.</td>
</tr>
</tbody>
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(continued)
### β-LACTAMS COMPARISON CHART (continued)

<table>
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<tr>
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<th>PEDIATRIC DOSAGE</th>
<th>ADULT DOSAGE IN RENAL IMPAIRMENT&lt;sup&gt;a&lt;/sup&gt;</th>
<th>PEAK SERUM LEVELS (MG/L)&lt;sup&gt;b&lt;/sup&gt;</th>
<th>PERCENTAGE PROTEIN BOUND</th>
<th>COMMENTS</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Ampicillin Sodium</strong></td>
<td>Cap 250, 500 mg</td>
<td>PO 250–500 mg qid; IM or IV 500 mg–3 g q 4–6 hr, to a maximum of 12 g/day.</td>
<td>PO (&lt;20 kg) 50–100 mg/kg/day in 2–4 divided doses; PO or IV (&gt;20 kg) 100–400 mg/kg/day in divided doses q 4–6 hr.</td>
<td>Cl&lt;sub&gt;cr&lt;/sub&gt; &lt;20 mL/min: same dose q 12 hr.</td>
<td>4 (PO) 58 (IV)</td>
<td>22</td>
<td>About 50% oral bio-availability; GI side effects and rashes are frequent. IV sodium = 3 mEq/g.</td>
</tr>
<tr>
<td><strong>EXTENDED-SPECTRUM PENICILLINS</strong></td>
<td>Various</td>
<td>IV 3–4 g q 4–6 hr.</td>
<td>IV (&lt;7 days) 50–100 mg/kg q 12 hr; (neonates &gt;7 days) 50–100 mg/kg q 6–8 hr; (children) 300 mg/kg/ day in divided doses q 4–6 hr to a maximum of 24 g/day.</td>
<td>Cl&lt;sub&gt;cr&lt;/sub&gt; 10–30 mL/min: 3 g q 8 hr; Cl&lt;sub&gt;cr&lt;/sub&gt; &lt;10 mL/min: 2 g q 8 hr.</td>
<td>263 (IV 4 g)</td>
<td>35</td>
<td>Spectrum similar to ticarcillin, but better enterococcal coverage. Least active drug in this class against <em>P. aeruginosa</em>. Sodium = 1.85 mEq/g.</td>
</tr>
<tr>
<td>DRUG CLASS AND DRUG</td>
<td>DOSAGE FORMS</td>
<td>ADULT DOSAGE</td>
<td>PEDIATRIC DOSAGE</td>
<td>ADULT DOSAGE IN RENAL IMPAIRMENT&lt;sup&gt;a&lt;/sup&gt;</td>
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<tr>
<td><strong>Piperacillin</strong></td>
<td>Inj 2, 3, 4, 40 g.</td>
<td>IV 3–4 g q 4–6 hr, to a maximum of 24 g/day</td>
<td>Not well established. IV (neonates) 100 mg/kg q 12 hr; (children) 200–300 mg/kg/day in divided doses q 4–6 hr.</td>
<td>Cl&lt;sub&gt;r&lt;/sub&gt; 20–40 mL/min: 3–4 g q 8 hr; Cl&lt;sub&gt;r&lt;/sub&gt; &lt;20 mL/min: 3–4 g q 12 hr.</td>
<td>244 (IV 4 g)</td>
<td>15–20</td>
<td>Best activity against <em>P. aeruginosa</em>. Sodium = 1.85 mEq/g.</td>
</tr>
<tr>
<td>Sodium</td>
<td>Inj 2, 3, 4, 40 g.</td>
<td>IV 3–4 g q 4–6 hr, to a maximum of 24 g/day</td>
<td>Not well established. IV (neonates) 100 mg/kg q 12 hr; (children) 200–300 mg/kg/day in divided doses q 4–6 hr.</td>
<td>Cl&lt;sub&gt;r&lt;/sub&gt; 20–40 mL/min: 3–4 g q 8 hr; Cl&lt;sub&gt;r&lt;/sub&gt; &lt;20 mL/min: 3–4 g q 12 hr.</td>
<td>244 (IV 4 g)</td>
<td>15–20</td>
<td>Best activity against <em>P. aeruginosa</em>. Sodium = 1.85 mEq/g.</td>
</tr>
<tr>
<td>Pipracil</td>
<td>24 g/day nates) 100 mg/kg q 12 hr; (children) 200–300 mg/kg/day in divided doses q 4–6 hr.</td>
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<tr>
<td><strong>Ticarcillin</strong></td>
<td>Inj 1, 3, 6, 20, 30 g.</td>
<td>IV 2–4 g q 4–6 hr, to a maximum of 24 g/day.</td>
<td>IV (neonates ≤7 days and &lt;2 kg) 75 mg/kg q 12 hr; (neonates &gt;7 days and &lt;2 kg or ≤7 days and &gt;2 kg) 75 mg/kg q 8 hr; (neonates &gt;7 days and &gt;2 kg) 75 mg/kg q 6 hr; (children) 200–300 mg/kg/day in divided doses q 6–8 hr.</td>
<td>Cl&lt;sub&gt;r&lt;/sub&gt; 30–60 mL/min: 2 g q 4 hr; Cl&lt;sub&gt;r&lt;/sub&gt; 10–30 mL/min: 2 g q 8 hr; Cl&lt;sub&gt;r&lt;/sub&gt; &lt;10 mL/min: 2 g q 12 hr.</td>
<td>260 (IV 3 g)</td>
<td>50–60</td>
<td>Less active than piperacillin against <em>P. aeruginosa</em>. No activity against Klebsiella spp. More antiplatelet effect than mezlocillin or piperacillin. Sodium = 5.2–6.5 mEq/g.</td>
</tr>
<tr>
<td>Disodium</td>
<td>24 g/day nates) 100 mg/kg q 12 hr; (children) 200–300 mg/kg/day in divided doses q 4–6 hr.</td>
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<tr>
<td>Ticar</td>
<td>24 g/day nates) 100 mg/kg q 12 hr; (children) 200–300 mg/kg/day in divided doses q 4–6 hr.</td>
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</tr>
</tbody>
</table>

<sup>a</sup> Cl<sub>r</sub> = creatinine clearance.

<sup>b</sup> Sodium = equivalent of sodium citrate or sodium bicarbonate.
### β-LACTAMS COMPARISON CHART (continued)

<table>
<thead>
<tr>
<th>Drug Class and Drug</th>
<th>Dosage Forms</th>
<th>Adult Dosage</th>
<th>Pediatric Dosage</th>
<th>Adult Dosage in Renal Impairment*</th>
<th>Peak Serum Levels (mg/L)b</th>
<th>Percentage Protein Bound</th>
<th>Comments</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>PENICILLIN AND β-LACTAMASE INHIBITOR COMBINATIONS</strong></td>
<td></td>
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<td></td>
</tr>
<tr>
<td>Amoxicillin and Clavulanate</td>
<td>Chew Tab 125 mg amoxicillin plus 31.25 mg clavulanate, 200 mg amoxicillin plus 28.5 mg clavulanate, 250 mg amoxicillin plus 62.5 mg clavulanate, 400 mg amoxicillin plus 57 mg clavulanate; Susp 25 mg amoxicillin plus 6.25 mg clavulanate/mL, 40 mg amoxicillin plus 5.7 mg clavulanate, 50 mg amoxicillin plus 12.5 mg clavulanate, 80 mg amoxicillin plus 11.4 mg clavulanate/mL. Tab 250 mg amoxicillin plus 125 mg clavulanate, 500 mg amoxicillin plus 125 mg clavulanate, 875 mg amoxicillin plus 125 mg clavulanate.</td>
<td>PO “250” or “500” tablet q 8 hr or “875” tablet q 12 hr.</td>
<td>PO 20–40 mg/kg/day (of amoxicillin) in 3 divided doses or 45 mg/kg/day in 2 divided doses.</td>
<td>Cl\text{r} 10–30 mL/min: 250–500 mg bid Cl\text{r} &lt;10 mL/min 250–500 mg q 24 hr.</td>
<td>9 (PO 500 mg amoxicillin) 2.6 (PO 125 mg clavulanate)</td>
<td>20 (amoxicillin) 22 (clavulanate)</td>
<td>Active against ampicillin-resistant <em>S. aureus</em>, <em>B. fragilis</em>, and β-lactamase-producing Enterobacteriaceae. More diarrhea than with amoxicillin. Do not substitute 2 “250” tablets for 1 “500” tablet.</td>
</tr>
</tbody>
</table>
**β-LACTAMS COMPARISON CHART (continued)**

<table>
<thead>
<tr>
<th>DRUG CLASS AND DRUG</th>
<th>DOSAGE FORMS</th>
<th>ADULT DOSAGE</th>
<th>PEDIATRIC DOSAGE</th>
<th>ADULT DOSAGE IN RENAL IMPAIRMENT</th>
<th>PEAK SERUM LEVELS (MG/L)b</th>
<th>PERCENTAGE PROTEIN BOUND</th>
<th>COMMENTS</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ampicillin Sodium and Sulbactam</td>
<td>Inj 1 g ampicillin plus 500 mg sulbactam/vial, 2 g ampicillin plus 1 g sulbactam/vial, 10 g ampicillin plus 5 g sulbactam/vial.</td>
<td>IM or IV 1.5–3 g of the combination q 6–8 hr, to a maximum of 12 g/day.</td>
<td>IM or IV (3 months–12 yr) 150–300 mg of the combination q 6 hr to a maximum of 12 g/day.</td>
<td>Clcr 15–30 mL/min: same dose q 12 hr; Clcr 5–14 mL/min: same dose q 24 hr.</td>
<td>Clcr (IV 1 g ampicillin) 58; Clcr (IV 500 mg sulbactam) 30</td>
<td>22 (ampicillin); 38 (sulbactam)</td>
<td>Spectrum similar to Augmentin. Sodium = 5 mEq/1.5 g.</td>
</tr>
<tr>
<td>Piperacillin Sodium and Tazobactam</td>
<td>Inj 2.25, 3.375, 4.5, 40.5 g (0.5 g tazobactam) and 4 g piperacillin.</td>
<td>IV 3.375–4.5 g q 4–6 hr; 3.375 g q 4 hr or 4.5 g q 6 hr for P. aeruginosa.</td>
<td>Safety and efficacy not established.</td>
<td>Clcr 20–40 mL/min: 2.25 g q 6 hr; Clcr &lt;20 mL/min: 2.25 g q 8 hr.</td>
<td>Clcr (IV 4 g piperacillin) 400; Clcr (IV 0.5 g tazobactam) 34</td>
<td>15–20 (piperacillin); 1 (tazobactam)</td>
<td>Similar spectrum to Timentin, but better activity against P. aeruginosa and enterococci. Sodium = 2.35 mEq/g of piperacillin.</td>
</tr>
<tr>
<td>Ticarcillin Disodium and Clavulanate Potassium</td>
<td>Inj 3.1, 31 g (100 mg clavulanate/3 g ticarcillin).</td>
<td>IV 3.1 g q 4–6 hr.</td>
<td>IV (≥3 months) &lt;60 kg: 50 mg/kg (of ticarcillin) q 4–6 hr; ≥60 kg: same as adult dosage.</td>
<td>Clcr 30–60 mL/min: 3.1 g q 6 hr; Clcr &lt;10 mL/min: 3.1 g q 12 hr.</td>
<td>Clcr (IV 3 g ticarcillin) 260; Clcr (IV 100 mg clavulanate) 8</td>
<td>50–60 (ticarcillin); 8 (clavulanate)</td>
<td>Improved activity over ticarcillin against S. aureus, H. influenzae, and anaerobes, but not P. aeruginosa or E. cloacae. Sodium = 4.7 mEq/g of ticarcillin.</td>
</tr>
</tbody>
</table>

*aUsual dose means individual doses given at the specified interval; usual dosage means total daily dosage.

*bAverage peak serum concentrations following administration of a 500 mg oral dose or a 1 g IV infusion over 30 min, except as noted.

*cConcentration dependent.

*dWith dosages recommended in marked renal impairment, clavulanate concentrations may provide ineffective synergism with ticarcillin.173

From references 172–174 and 184–186 and product information.
AZITHROMYCIN  Zithromax

Pharmacology. Azithromycin is a macrolide with a 15-membered ring (making it an azalide) that is slightly less active than erythromycin against Gram-positive bacteria but substantially more active against Moraxella (Branhamella) catarrhalis, Haemophilus sp., Legionella sp., Neisseria sp., Bordetella sp., Mycoplasma spp., and Chlamydia trachomatis. The drug also has activity against aerobic Gram-negative bacilli and Mycobacterium avium and is comparable to erythromycin in its activity against Campylobacter sp. It is the most active macrolide for Toxoplasma gondii, including activity against the cyst form.\(^{191–194}\)

Administration and Adult Dosage. PO for mild to moderate acute bacterial exacerbations of COPD, pneumonia, pharyngitis or tonsillitis, and uncomplicated skin and skin structure infections 500 mg as a single dose on the first day followed by 250 mg/day on days 2–5 for a total dosage of 1.5 g. PO for nongonococcal urethritis and cervicitis caused by C. trachomatis or for chancroid (Haemophilus ducreyi) 1 g as a single dose.\(^{195}\) PO for treatment of M. avium complex in AIDS patients 500 mg/day in combination with ethambutol.\(^ {196}\) PO for prophylaxis of M. avium complex in AIDS patients 1.2 g once weekly alone or in combination with rifabutin 300 mg/day.\(^ {197}\) PO for endocarditis prophylaxis 500 mg 1 hr before procedure. IV for pelvic inflammatory disease 500 mg/day for 1–2 days, followed by PO 250 mg/day to complete 7 days of therapy. IV for community-acquired pneumonia 500 mg/day for at least 2 days followed by PO 500 mg/day to complete 7–10 days of therapy.

Special Populations. Pediatric Dosage. PO for otitis media (≥6 months) 10 mg/kg as a single daily dose on day 1, followed by 5 mg/kg/day as a single dose on days 2–5. PO for streptococcal pharyngitis/tonsillitis (≥2 yr) 12 mg/kg/day as a single dose for 5 days.\(^{198–200}\) PO for endocarditis prophylaxis 15 mg/kg 1 hr before procedure. IV (<16 yr) safety and efficacy not established.

Geriatric Dosage. Same as adult dosage.

Other Conditions. Dosage reduction may be needed in severe hepatic impairment, but guidelines are not available.

Dosage Forms. Tab 250, 600 mg; Susp 20, 40 mg/mL; Pwdr for Oral Susp 1 g; Inj 500 mg.

Patient Instructions. Take the oral suspension with a full glass of water on an empty stomach (1 hour before or 2 hours after meals) for best absorption. Tablets may be taken without regard to meals. Do not take aluminum- or magnesium-containing antacids with azithromycin.

Missed Doses. Take this drug at regular intervals. If you miss a dose, take it as soon as you remember. If it is about time for the next dose, take that dose only. Leave at least 12 hours between doses. Do not double the dose or take extra.

Pharmacokinetics. Fate. Oral bioavailability is 37%. After a 500 mg oral capsule, a peak serum concentration of 0.41 mg/L (0.55 μmol/L) is achieved in 2 hr. Plasma protein binding is 7–50%, primarily to α₁-acid glycoprotein.
Azithromycin penetrates macrophages and polymorphonuclear leukocytes, accounting for intracellular concentrations that are 40-fold extracellular concentrations. Azithromycin is widely distributed throughout the body, and tissue concentrations (including the CNS) range from 10- to 150-fold higher than those in serum. Tissue concentrations peak 48 hr after administration, and high concentrations persist for several days after drug discontinuation. Elimination is polyphasic, reflecting rapid initial distribution into tissues, followed by slow elimination. \( V_d \) is 23–31 L/kg; Cl is 38 L/hr in adults. Azithromycin is metabolized in the liver and eliminated largely through biliary excretion; only 6% is excreted unchanged in urine.\(^{192,194,201,202}\)

\( t_{1/2} \). Terminal phase \( 11–68+ \) hr.\(^{192,202}\)

**Adverse Reactions.** The drug is well tolerated. Frequent adverse effects are mild to moderate diarrhea, nausea, and abdominal pain. Headache and dizziness occur occasionally. Rash, angioedema, hepatomegaly, and cholestatic jaundice are reported rarely.\(^{192}\)

**Contraindications.** Hypersensitivity to any macrolide.

**Precautions.** Use during pregnancy only if clearly needed. Use caution in patients with impaired hepatic function or severely impaired renal function.

**Drug Interactions.** Azithromycin does not interact with hepatic cytochrome P450 enzymes and, unlike erythromycin and clarithromycin, is not associated with these types of interactions.\(^{203}\)

**Parameters to Monitor.** Baseline and periodic liver function tests during prolonged therapy.

### CLARITHROMYCIN

**Pharmacology.** Clarithromycin is a semisynthetic macrolide antibiotic that is slightly more active than erythromycin against Gram-positive bacteria, *Moraxella (Branhamella) catarrhalis*, and *Legionella* sp. It is very active against *Chlamydia* sp. and superior to other macrolides in its activity against *Mycobacterium avium* complex (MAC).\(^{192,194,201,204,205}\)

**Administration and Adult Dosage.** PO for respiratory and skin infections 250–500 mg bid; PO for MAC in AIDS patients 500 mg bid. PO for endocarditis prophylaxis 500 mg 1 hr before procedure. PO for eradication of *Helicobacter pylori* 500 mg tid in combination with proton pump inhibitors and other drugs. (See Gastrointestinal Drugs, Treatment of *Helicobacter pylori* Infection in Peptic Ulcer Disease Chart.)\(^{206}\)

**Special Populations.** *Pediatric Dosage.* PO for community-acquired pneumonia 15 mg/kg q 12 hr for 10 days; PO for other indications 7.5 mg/kg bid, to a maximum of 500 mg bid. PO for endocarditis prophylaxis 15 mg/kg 1 hr before the procedure.

**Geriatric Dosage.** Same as adult dosage.

**Other Conditions.** Reduce dosage by 50% with Cl\(_r\) <30 mL/min.

**Dosage Forms.** Tab 250, 500 mg; Susp 25, 37.5, 50 mg/mL.
Pharmacokinetics. 

Fate. Clarithromycin is acid-stable and absorbed well with or without food. Bioavailability is 55%, with peak serum concentrations of about 2 mg/L attained after a 400 mg oral dose. The hydroxy metabolite is active and may be synergistic in vitro with the parent drug. \( t_{1/2} \). (Clarithromycin) 4.5 hr; (hydroxy-metabolite) 4–9 hr.

Adverse Reactions. Similar to erythromycin, but clarithromycin has better GI tolerance.

Contraindications. Hypersensitivity to any macrolide antibiotic; concurrent use with certain other drugs. (See Drug Interactions.)

Precautions. Use with caution in severe renal or hepatic function impairment; dosage reduction is advised.

Drug Interactions. Clarithromycin has a lower affinity for CYP3A4 than erythromycin and therefore has fewer clinically important drug interactions; however, its use is contraindicated with astemizole or cisapride. Serum concentrations of theophylline and carbamazepine also can be increased by clarithromycin.

ERYTHROMYCIN AND SALTS

Pharmacology. Erythromycin is a bacteriostatic macrolide antibiotic with a spectrum similar to that of penicillin G; it is also active against Mycoplasma pneumoniae and Legionella pneumophila. It acts by binding to the 50S ribosomal subunit, inhibiting protein synthesis. Gram-positive organisms develop resistance via R-factor mediated alteration of the binding site. Gram-negative organisms are resistant because of cell wall impermeability.

Administration and Adult Dosage. (See Macrolide Antibiotics Comparison Chart.) For gastroparesis 200 mg IV of the lactobionate salt, 250 mg PO of the ethylsuccinate salt or 500 mg PO of the base 15–120 min before meals and at bedtime appear to be effective.

Special Populations. Pediatric Dosage. (See Macrolide Antibiotics Comparison Chart.)

Geriatric Dosage. Same as adult dosage.

Other Conditions. Dosage adjustment is probably unnecessary in renal impairment.

Dosage Forms. (See Macrolide Antibiotics Comparison Chart.)

Patient Instructions. Take this drug with a full glass of water on an empty stomach (1 hour before or 2 hours after meals) for best absorption. Refrigerate the suspension.

Missed Doses. Take this drug at regular intervals. If you miss a dose, take it as soon as you remember. If it is about time for the next dose, take that dose only. Leave at least 4–6 hours between doses. Do not double the dose or take extra.

Pharmacokinetics. Fate. Oral absorption varies widely with the salt and dosage forms (see Macrolide Antibiotics Comparison Chart), with peak serum concentrations occurring from 30 min (suspension) to 4 hr (coated tablet) after administration. However, enteric-coated erythromycin base tablets, stearate tablets, and esto-
late capsules produce equivalent erythromycin serum levels when administered to fasting subjects. Food or restricted water intake (ie, <20 mL) with a dose dramatically lowers the absorption of the stearate form. The drug is 83 ± 5% plasma protein bound and widely distributed into most tissues, cavities, and body fluids except the brain and CSF (even with meningeal inflammation). \( V_d \) is 0.6 ± 0.1 L/kg; \( \text{Cl} \) is 0.55 ± 0.25 L/hr/kg. Erythromycin is partially metabolized in the liver by CYP3A3/4 and excreted primarily as unchanged erythromycin with high concentrations in the bile and feces. Only 12–15% of an IV dose is excreted unchanged in urine.\(^{52,207}\)

\[ t_{1/2} = 1.6 ± 0.7 \text{ hr} \]; unchanged or slightly prolonged in anuric patients, based on minimal data; prolonged in cirrhosis.\(^{52}\)

**Adverse Reactions.** Frequent GI distress. IM form is very painful, despite local anesthetic (butamben) in the product, and might produce sterile abscesses. IV administration frequently produces pain, venous irritation, and phlebitis. Mild elevations of serum hepatic enzymes occur frequently. Transient deafness occurs occasionally with high dosages.\(^{207,211}\) Rare, but potentially serious, reversible intrahepatic cholestatic jaundice occurs primarily with the estolate and ethylsuccinate forms, usually in adults after 10–14 days of therapy, although it can occur after the first dose if there is a history of previous use. Prodrome includes malaise, nausea, vomiting, fever, and abdominal pain (which can be severe and misdiagnosed as acute surgical abdomen). Symptoms resolve in 1–2 weeks, and serum enzymes return to normal over several months.

**Contraindications.** Concurrent use with astemizole, cisapride or pimozide; IM form in patients with hypersensitivity to local anesthetics of the para-aminobenzoic acid type (eg, procaine); hepatic dysfunction (estolate and ethylsuccinate forms).

**Precautions.** Pregnancy. Use with caution in patients with liver disease because of possibly impaired excretion.

**Drug Interactions.** Erythromycin inhibits CYP3A4 and can reduce hepatic metabolism of some drugs, including astemizole, carbamazepine, cisapride, cyclosporine, theophylline, triazolam, warfarin, and others.\(^{212}\) (See Contraindications.)

**Parameters to Monitor.** Liver function tests in patients who experience prodromal symptoms (see Adverse Reactions) while receiving the estolate or ethylsuccinate form; check daily for vein irritation and phlebitis in patients receiving IV forms. Closely monitor the effects of other drugs that interact with erythromycin during concurrent use.

**Notes.** Avoid injectable forms if at all possible. Erythromycin is more active in an alkaline environment. Unrelated to its antibacterial effect, erythromycin in low doses binds to motilin receptors in the GI tract to stimulate gastric emptying. It is the most prokinetic macrolide and has been used in gastroparesis and other GI motility disorders.\(^{210,213-216}\)
<table>
<thead>
<tr>
<th>DRUG</th>
<th>DOSAGE FORMS</th>
<th>ADULT DOSAGE</th>
<th>PEDIATRIC DOSAGE</th>
<th>COMMENTS</th>
</tr>
</thead>
<tbody>
<tr>
<td>Azithromycin</td>
<td>Tab 250, 600 mg</td>
<td>PO 500 mg once, then 250 mg/day for 4 days; PO for otitis media or pneumonia.</td>
<td>PO for otitis media or pneumonia. 10 mg/kg once, then 5 mg/kg/day for 4 days; PO for pharyngitis/tonsillitis 12 mg/kg/day for 5 days.</td>
<td>Broader spectrum than erythromycin. Less GI intolerance than erythromycin. Little or no P450 inhibition.</td>
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<td></td>
<td>Susp 20, 40 mg/mL</td>
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<td>Pwdr for Oral Susp 1 g</td>
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<tr>
<td></td>
<td>Inj 500 mg</td>
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<tr>
<td>Zithromax</td>
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<tr>
<td>Clarithromycin</td>
<td>Tab 250, 500 mg</td>
<td>PO 250–500 mg bid; PO for MAC 500 mg bid; PO for H. pylori 500 mg tid with other agents.</td>
<td>PO for pneumonia 15 mg/kg q 12 hr for 10 days; PO for other uses 7.5 mg/kg bid.</td>
<td>Broader spectrum than erythromycin. Food does not decrease absorption. Less GI intolerance than erythromycin. Less inhibition of CYP3A4/3 than erythromycin.</td>
</tr>
<tr>
<td>Biaxin</td>
<td>Tab 250, 500 mg</td>
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<td></td>
<td>Susp 25, 37.5, 50 mg/mL</td>
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<tr>
<td>Dirithromycin</td>
<td>EC Tab 250 mg.</td>
<td>PO 500 mg once daily for 7–14 days.</td>
<td>&lt;12 yr not recommended.</td>
<td>Spectrum similar to erythromycin, but less GI intolerance and little or no P450 inhibition.</td>
</tr>
<tr>
<td>Dynabac</td>
<td></td>
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<tr>
<td>Erythromycin</td>
<td>EC Tab 250, 333, 500 mg</td>
<td>PO 1 g/day in 2–4 doses, to a maximum of 4 g/day.</td>
<td>PO 30–50 mg/kg/day in 4 doses; may double in severe infection.</td>
<td>Food interferes with absorption of uncoated products; EC products appear to be among the best tolerated erythromycin formulations.</td>
</tr>
<tr>
<td>Base</td>
<td>EC Tab 333, 500 mg</td>
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<tr>
<td>E-Mycin</td>
<td>SR Cap 250 mg</td>
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<tr>
<td>Ery-Tab</td>
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<tr>
<td>ERYC</td>
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</tr>
<tr>
<td>Various</td>
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</table>

(continued)
## MACROLIDE ANTIBIOTICS COMPARISON CHART (continued)

<table>
<thead>
<tr>
<th>DRUG</th>
<th>DOSAGE FORMS</th>
<th>ADULT DOSAGE</th>
<th>PEDIATRIC DOSAGE</th>
<th>COMMENTS</th>
</tr>
</thead>
<tbody>
<tr>
<td>Erythromycin</td>
<td>Cap 250 mg</td>
<td>PO 250–500 mg q 6 hr, to a maximum of 4 g/day.</td>
<td>PO 30–50 mg/kg/day in 3–4 doses.^[a]</td>
<td>PO well absorbed; unaffected by food and highly resistant to gastric acid hydrolysis; absorbed as pro-ponate ester which predominates in serum (8:1) and might be less active; rare intrahepatic cholestatic jaundice.^[b]</td>
</tr>
<tr>
<td>Estolate</td>
<td>Susp 25, 50 mg/mL</td>
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<tr>
<td>Ilosone</td>
<td>Tab 500 mg</td>
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<tr>
<td>Various</td>
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<tr>
<td>Erythromycin</td>
<td>Drp 40 mg/mL</td>
<td>PO 400 mg q 6 hr, to a maximum of 4 g/day.</td>
<td>PO 30–50 mg/kg/day in 3–4 doses; may double in severe infection.^[a]</td>
<td>Absorbed better than base; intermedi-ate susceptibility to gastric acid hydrolysis. Absorbed as ester, which predominates in serum (3:1) and might be less active. Rare intrahepatic cholestatic jaundice.^[b]</td>
</tr>
<tr>
<td>Ethylsuccinate</td>
<td>Susp 40, 80 mg/mL</td>
<td></td>
<td></td>
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</tr>
<tr>
<td>E.E.S.</td>
<td>Chew Tab 200 mg</td>
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</tr>
<tr>
<td>EryPed</td>
<td>Tab (coated) 400 mg</td>
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<tr>
<td>Various</td>
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<tr>
<td>Erythromycin</td>
<td>Inj (IV only) 1 g.</td>
<td>IV 15–20 mg/kg/day in 3–4 doses, to a maximum of 4 g/day.</td>
<td>IV same as adult dosage in 2–4 doses; may double in severe infection.^[a]</td>
<td>Painful; phlebitis frequent; avoid use if possible. Infuse over 20–60 min.^[b]</td>
</tr>
<tr>
<td>Gluceptate</td>
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<tr>
<td>Ilotycin</td>
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</tbody>
</table>

^[a] in severe infection.^[b]
<table>
<thead>
<tr>
<th>DRUG</th>
<th>DOSAGE FORMS</th>
<th>ADULT DOSAGE</th>
<th>PEDIATRIC DOSAGE</th>
<th>COMMENTS</th>
</tr>
</thead>
<tbody>
<tr>
<td>Erythromycin Lactobionate</td>
<td>Inj (IV only) 500 mg, 1 g.</td>
<td>Same as erythromycin gluceptate.</td>
<td>Same as erythromycin gluceptate.</td>
<td>Same as erythromycin gluceptate.</td>
</tr>
<tr>
<td>Erythrocin Stearate</td>
<td>Tab (film coated) 250, 500 mg.</td>
<td>PO 1 g/day in 2 or 4 doses, to a maximum of 4 g/day.</td>
<td>PO 30–50 mg/kg/day in 4 doses; may double in severe infections.</td>
<td>Absorption about equal to ethylsuccinate, although food interferes markedly with absorption. Hydrolyzed to free base before absorption.</td>
</tr>
</tbody>
</table>

*In newborns, data are available for erythromycin estolate only, suggesting an oral dosage of 40 mg/kg/day in 2–4 divided doses.

*Despite differences in oral absorption, no clinical studies have shown any salt to be clearly superior in any particular therapeutic use. From references 217–222 and product information.
Quinolones

**CIPROFLOXACIN** Ciloxan, Cipro

**Pharmacology.** Ciprofloxacin is a fluoroquinolone that inhibits bacterial DNA-gyrase, an enzyme responsible for the unwinding of DNA for transcription and subsequent supercoiling of DNA for packaging into chromosomal subunits. It is highly active against aerobic, Gram-negative bacilli, especially Enterobacteriaceae, with MICs often <0.1 mg/L. It is also active against some strains of *Pseudomonas aeruginosa* and *Staphylococcus* spp., with an MIC of 0.5–1 mg/L. However, recent reports indicate increasing resistance to this agent in methicillin-resistant *S. aureus*. It has poor activity against streptococci and anaerobes.223,224

**Administration and Adult Dosage.** PO for uncomplicated UTIs 250 mg q 12 hr. PO for moderate to severe systemic infections 500–750 mg q 12 hr; PO for gonorrhea 250–500 mg once. PO for chancroid 500 mg q 12 hr for 3 days.195 IV 200–400 mg q 12 hr

**Special Populations.** Pediatric Dosage. (<16 yr) safety and efficacy not established. Use has been limited because of the potential for arthropathy. Ciprofloxacin has been used in children 6–16 yr old in limited situations to treat serious infections. IV for *P. aeruginosa* infections in cystic fibrosis 15–30 mg/kg/day in 2–3 divided doses. PO for *P. aeruginosa* infections in cystic fibrosis 20–40 mg/kg/day in 2 divided doses.225

**Geriatric Dosage.** Reduce dosage for age-related reduction in renal function, although dosage reduction is not necessary with only minor age-related renal function changes.226

**Other Conditions.** Reduce dosage by 50% or double the dosage interval when Clr <30 mL/min; special dosage adjustments in patients with cystic fibrosis are not necessary.227

**Dosage Forms.** Inj 200, 400 mg; Susp 50, 100 mg/mL; Tab 100, 250, 500, 750 mg; Ophth Drp (Ciloxan) 3.5 mg/mL (equivalent to 3 mg/mL base); Otic Susp 2 mg plus hydrocortisone 10 mg/mL.

**Patient Instructions.** This drug may be taken with food to minimize stomach upset. Avoid antacid use during treatment; calcium, iron, or zinc supplements can reduce absorption. Avoid excessive exposure to sunlight during ciprofloxacin treatment. Report any tendon pain or inflammation that occurs during therapy.

**Missed Doses.** Take this drug at regular intervals. If you miss a dose, take it as soon as you remember. If it is about time for the next dose, take that dose only. Leave at least 6–8 hours between doses.

**Pharmacokinetics.** Serum Levels. A peak serum level of 4–6 mg/L (12–18 mmol/L) 2 hr after an oral 750–1000 mg dose is proposed as evidence of absorption adequate for tuberculosis therapy.58

**Fate.** About 70–80% absorbed orally; food decreases the rate but not the extent of absorption. Aluminum-, calcium-, or magnesium-containing antacids or sucralfate markedly decrease the extent of absorption. Peak serum concentrations are
3 ± 0.6 mg/L (9 ± 1.8 mmol/L) after a 750 mg oral dose; a 200 mg IV dose infused over 30 min results in a peak concentration of about 3.2 ± 0.6 mg/L. \( V_d \) averages 2 L/kg. Renal clearance averages 0.26 L/hr/kg. Less than 30% is plasma protein bound. Ciprofloxacin attains very high concentrations in many body fluids and tissues, most notably urine, prostate, and pulmonary mucosa. CSF concentrations are <1 mg/L; experience with the drug in the treatment of meningitis is very limited. From 45% and 60% of a parenteral dose is recovered unchanged in urine; the remainder is excreted as four metabolites or eliminated in feces. \( t_{1/2} \) 4.2 ± 0.63 hr, 6.9 ± 2.9 hr in severe renal impairment.

**Adverse Reactions.** GI intolerance (nausea, vomiting, diarrhea, abdominal discomfort) occurs frequently. CNS effects such as headaches and restlessness have occurred in 1–2% of patients. Other CNS effects (eg, dizziness, insomnia, anxiety, irritability, and seizures) have been reported in fewer than 1% of patients. Skin rashes and photosensitivity occur occasionally. Anaphylaxis occurs rarely.

**Contraindications.** Hypersensitivity to any quinolone.

**Precautions.** Pregnancy; lactation.

**Drug Interactions.** Aluminum-, calcium-, or magnesium-containing antacids markedly reduce oral absorption. Although there is some information that spacing administration by \( \geq 2 \) hr might minimize these interactions, it is probably best not to use ciprofloxacin in patients taking long-term antacid therapy. Iron supplements and zinc-containing multivitamins can reduce absorption. Theophylline clearance can be reduced in some patients receiving ciprofloxacin. Patients receiving fluoroquinolones and methylxanthines such as theophylline or caffeine might be at increased risk of CNS toxicity (eg, convulsions). Warfarin metabolism can be impaired by ciprofloxacin, although studies with the fluoroquinolone enoxacin indicate that only the metabolism of the less active (R)-warfarin is affected. Use caution when adding ciprofloxacin in a patient taking warfarin. The solubility of ciprofloxacin is reduced at higher pH values; thus, avoid alkalization of the urine.

**Parameters to Monitor.** Monitor serum theophylline levels closely in patients receiving theophylline. Monitor prothrombin time and signs of bleeding in patients on warfarin.

**OFLOXACIN** Floxin, Ocuflux

**LEVOFLOXACIN** Levaquin, Quixin

**Pharmacology.** Ofloxacin is a systemic fluoroquinolone similar to ciprofloxacin. Levofloxacin is the active L-isomer of ofloxacin that allows higher dosages of the active form to be given with fewer side effects. Ofloxacin has greater activity against *Chlamydia trachomatis*, *Ureaplasma urealyticum*, *Mycoplasma pneumoniae*, and *Mycobacterium tuberculosis* than ciprofloxacin, but less activity against *Pseudomonas aeruginosa*. (See Fluoroquinolones Comparison Chart.)

**Adult Dosage.** (Ofloxacin) IV or PO for systemic infections 400 mg q 12 hr; PO for nongonococcal urethritis 300 mg q 12 hr for 7 days; PO for acute, un-
complicated gonorrhea 400 mg once; IV or PO for urinary tract infections 200 mg q 12 hr. (Levofloxacin) IV or PO 250–500 mg once daily. Reduce the dosage of both drugs in renal impairment. Ophth (Ofloxacin) 1–2 drops q 30 min while awake and q 4–6 hr after retiring for 2 days, then q 1 hr while awake for 4–6 days, then qid until cure. (Levofloxacin) 1–2 drops q 2 hr while awake up to 8 times/day for 2 days, then q 4 hr while awake for 5 days. Otic (Ofloxacin) 10 drops bid for 14 days. (See Fluoroquinolones Comparison Chart.)

**Pediatric Dosage.** PO, IV (<18 yr) safety and efficacy not established. Ophth (<1 yr) safety and efficacy not established; (ofloxacin, levofloxacin) same as adult dosage. Otic (Ofloxacin) 5 drops bid for 10 days.

**Dosage Forms.** (Ofloxacin) Inj 200, 400 mg; Tab 200, 300, 400 mg; Ophth Drp (Ocuflox) 3 mg/mL; Otic Drp 3 mg/mL. (Levofloxacin) Inj 5, 25 mg/mL; Tab 250, 500 mg; Ophth Drp (Quixin) 5 mg/mL.

**Pharmacokinetics.** Ofloxacin is >95% bioavailable orally. A peak serum concentration of 8–12 mg/L (22–33 mmol/L) 2 hr after an oral dose of 600–800 mg is proposed as evidence of absorption adequate for tuberculosis therapy. Ofloxacin is predominantly renally excreted with a half-life of 5–7 hr.

**Adverse Reactions.** (See Ciprofloxacin.)

**Drug Interactions.** Ofloxacin does not alter hepatic metabolism of methylxanthine compounds (eg, caffeine, theophylline). However, like other fluoroquinolones, cations markedly reduce the absorption of this agent.
<table>
<thead>
<tr>
<th>DRUG</th>
<th>DOSAGE FORMS</th>
<th>ADULT DOSAGE</th>
<th>DOSAGE IN RENAL IMPAIRMENT</th>
<th>ORAL BIOAVAILABILITY (PERCENT)</th>
<th>PEAK SERUM LEVELS (MG/L)²</th>
<th>COMMENTS³</th>
</tr>
</thead>
</table>
| **Ciprofloxacin** | Tab 100, 250, 500, 750 mg  
Inj 200, 400 mg  
Susz 50, 100 mg/mL  
Ophth Drp 0.3%  
Cipro HC Otic 2.5, 5 mL  
Otcsus 2 mg plus hydrocortisone 10 mg/mL. | PO 250–750 mg q 12 hr; PO for gonorrhea 500 mg once; IV 200–400 mg q 12 hr; Ophth 2 drops q 15 min–4 hr. | Cl₂ 30–50 mL/min: PO 250–500 mg q 12 hr; IV usual dosage; Cl₂ 5–29 mL/min: PO 250–500 mg q 18 hr; IV 200–400 mg q 18–24 hr; Dialysis: PO 250–500 mg q 24 hr after dialysis. | 60–80 | 3 ± 0.6 (PO 750 mg)  
3.2 ± 0.6 (IV 200 mg) | Most active against P. aeruginosa. Do not use against Gm+ organisms such as S. aureus. |
| **Cipro** |              |              |                            |                                 |                           |           |
| **Ciloxan** |              |              |                            |                                 |                           |           |
| **Cipro HC Otic** |              |              |                            |                                 |                           |           |
| **Enoxacin** | Tab 200, 400 mg. | PO 200–400 mg q 12 hr; PO for gonorrhea 400 mg once. | Cl₂ <30 mL/min: usual dose q 24 hr. | 83–90 | 5.5 (PO 400 mg) | Most potent inhibitor of theophylline metabolism. |

(continued)
## FLUROQUINOLONES COMPARISON CHART (continued)

<table>
<thead>
<tr>
<th>DRUG</th>
<th>DOSAGE FORMS</th>
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<th>DOSAGE IN RENAL IMPAIRMENT</th>
<th>ORAL BIOAVAILABILITY (PERCENT)</th>
<th>PEAK SERUM LEVELS (MG/L)³</th>
<th>COMMENTS³</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Gatifloxacin</strong></td>
<td>Tab 200, 400 mg PO or IV 200–400 mg q 24 hr.</td>
<td>Cl&lt;sub&gt;r&lt;/sub&gt; &lt;40 mL/min: 400 mg once, then 200 mg/day.</td>
<td>93</td>
<td>4.3 (PO 400 mg)</td>
<td>4.6 (IV 400 mg)</td>
<td>No effect on theophylline metabolism. Can be taken with or without food. Enhanced Gm+ activity. Prolongs QTc interval in some patients.</td>
</tr>
<tr>
<td><strong>Levofloxacin</strong></td>
<td>Tab 250, 500 mg PO or IV 250–500 mg q 24 hr.</td>
<td>Cl&lt;sub&gt;r&lt;/sub&gt; 20–49 mL/min: usual dose once, then 250 mg q 24 hr; Cl&lt;sub&gt;r&lt;/sub&gt; 10–19 mL/min: usual dose once, then 250 mg q 48 hr; Hemodialysis or peritoneal dialysis: 500 mg once, then 250 mg q 48 hr.</td>
<td>95–100</td>
<td>5.7 (PO 500 mg)</td>
<td>6.4 (IV 500 mg)</td>
<td>Active S(−) enantiomer of ofloxacin. Levofloxacin is not appreciably removed from the body during hemodialysis or peritoneal dialysis.</td>
</tr>
<tr>
<td><strong>Lomefloxacin</strong></td>
<td>Tab 400 mg PO 400 mg once daily.</td>
<td>Cl&lt;sub&gt;r&lt;/sub&gt; &lt;40 mL/min: PO 400 mg once, then 200 mg/day.</td>
<td>95–98</td>
<td>3.5 (PO 400 mg)</td>
<td>4.5 (400 mg)</td>
<td>No effect on theophylline metabolism. Long half-life of 8 hr. Relatively weak antibacterial. Phototoxic.</td>
</tr>
<tr>
<td><strong>Moxifloxacin</strong></td>
<td>Tab 400 mg PO 400 mg once daily.</td>
<td>No change required.</td>
<td>95</td>
<td>4.5 (400 mg)</td>
<td>4.5 (400 mg)</td>
<td>No effect on theophylline or warfarin metabolism. No food interactions. Enhanced activity against common community-acquired pneumonia pathogens. Prolongs QTc in some patients.</td>
</tr>
</tbody>
</table>

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<table>
<thead>
<tr>
<th>DRUG</th>
<th>DOSAGE FORMS</th>
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<td></td>
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</tr>
<tr>
<td>Norfloxacin</td>
<td>Tab 400 mg</td>
<td>PO 200–400 mg q 12 hr;</td>
<td>Cl&lt;sub&gt;r&lt;/sub&gt; &lt;30 mL/min:</td>
<td>30–40 (estimated)</td>
<td>1.4–1.6 (PO 400 mg)</td>
<td>Used in urinary and GI tract infections, sexually transmitted diseases and prostatitis only because of poor oral bioavailability.</td>
</tr>
<tr>
<td>Noroxin</td>
<td>Ophth Drp 0.3% (Chibroxin)</td>
<td>PO for gonorrhea 800 mg once; Ophth 1 or 2 drops q 2 hr–qid.</td>
<td>PO 400 mg/day.</td>
<td></td>
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<tr>
<td>Chibroxin</td>
<td>5 mL.</td>
<td></td>
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</tr>
<tr>
<td>Ofloxacin</td>
<td>Tab 200, 300, 400 mg</td>
<td>PO or IV 200–400 mg q 12 hr.</td>
<td>Cl&lt;sub&gt;r&lt;/sub&gt; 10–50 mL/min:</td>
<td>95–100</td>
<td>3.5–5.3 (PO 400 mg)</td>
<td>Most active against Chlamydia spp.; little effect on theophylline metabolism.</td>
</tr>
<tr>
<td>Floxin</td>
<td>Inj 200, 400 mg</td>
<td>PO for gonorrhea 400 mg once. Ophth 1 or 2 drops q 2–4 hr for 2 days, then qid up to 5 days.</td>
<td>usual dose q 24 hr; Cl&lt;sub&gt;r&lt;/sub&gt; &lt;10 mL/min:</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Ocuflox</td>
<td>Ophth Drp 0.3% (Ocuflox).</td>
<td></td>
<td>50% of usual dose q 24 hr.</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Sparfloxacin</td>
<td>Tab 200 mg.</td>
<td>PO 400 mg once, then 200 mg q 24 hr.</td>
<td>Cl&lt;sub&gt;r&lt;/sub&gt; &lt;50 mL/min:</td>
<td>90</td>
<td>0.62–0.71 (PO 200 mg)</td>
<td>Can be taken with or without food. No effect on theophylline metabolism. Pneumococcal activity superior to ciprofloxacin or ofloxacin. Phototoxic.</td>
</tr>
<tr>
<td>Zagam</td>
<td></td>
<td></td>
<td>400 mg once, then 200 mg q 48 hr.</td>
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<thead>
<tr>
<th>DRUG</th>
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<th>ADULT DOSAGE</th>
<th>DOSAGE IN RENAL IMPAIRMENT</th>
<th>ORAL BIOAVAILABILITY (PERCENT)</th>
<th>PEAK SERUM LEVELS (MG/L) a</th>
<th>COMMENTSb</th>
</tr>
</thead>
<tbody>
<tr>
<td>Trovafloxacin</td>
<td>Tab 100, 200 mg</td>
<td>PO 200 mg/day c</td>
<td>No adjustment needed in renal impairment or dialysis.</td>
<td>88</td>
<td>1.1 (PO 100 mg)</td>
<td>Broad spectrum of activity.</td>
</tr>
<tr>
<td>Alatrofloxacin</td>
<td>Inj 5 mg/mL.</td>
<td>IV 200–300 mg/day c</td>
<td></td>
<td>3.3</td>
<td>3.3 (PO 300 mg)</td>
<td>Penetrates into CSF better than other quinolones.</td>
</tr>
<tr>
<td>Mesylate</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>May have benefit against resistant organisms.</td>
</tr>
<tr>
<td>Trovan</td>
<td></td>
<td></td>
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</tbody>
</table>

aPeak serum concentrations following administration of the dose shown in parentheses.
bAll fluoroquinolones are associated with tendon rupture. Discontinue therapy at the first sign of tendon pain or inflammation, and patients should refrain from exercise until the diagnosis of tendinitis can be confidently excluded.203

cTrovafloxacin has been associated with serious liver injury, leading to liver transplantation or death. Reserve trovafloxacin for patients with serious, life-, or limb-threatening infections who receive their initial therapy in an inpatient facility (eg, hospital, long-term nursing care facility). Do not use trovafloxacin when a safer, alternative antimicrobial regimen will be effective. From references 202 and 223–239.
Sulfonamides

TRIMETHOPRIM AND SULFAMETHOXAZOLE
Bactrim, Septra, Various

Pharmacology. Sulfamethoxazole (SMZ) is a synthetic analogue of paraaminobenzoic acid (PABA), which competitively inhibits the synthesis of dihydropteric acid (an inactive folic acid precursor) from PABA in microorganisms. Trimethoprim (TMP) acts at a later step to inhibit the enzymatic reduction of dihydrofolic acid to tetrahydrofolic acid. The most important determinant of efficacy is usually the level of susceptibility to TMP; resistance to the combination is uncommon but appears to be increasing worldwide. The combination is active against many bacteria except anaerobes, Pseudomonas aeruginosa, and many Streptococcus faecalis spp. It is also highly active and effective against the protozoan Pneumocystis carinii. TMP/SMZ has in vitro activity against methicillin-resistant Staphylococcus aureus (MRSA), but clinical success has been variable and unpredictable.

Administration and Adult Dosage. PO for UTI 160 mg of TMP and 800 mg of SMZ q 12 hr for 10–14 days. PO for prophylaxis of recurrent UTI 40 mg TMP and 200 of SMZ at bedtime 3 times a week. PO for shigellosis 160 mg of TMP and 800 of SMZ q 12 hr for 5 days. IV for severe Gram-negative infections or shigellosis 8–10 mg/kg/day of TMP and 40–50 mg/kg/day of SMZ, in 2–4 equally divided doses, q 6–12 hr for 5 days for shigellosis and up to 14 days for severe UTI. PO or IV for P. carinii pneumonia (PCP) 12.5–20 mg/kg/day of TMP and 62.5–100 mg/kg/day of SMZ, in 2–4 equally divided doses, for up to 21 days. PO for PCP infection prophylaxis 160 mg of TMP and 800 mg of SMZ once daily; intermittent dosage (eg, 5 times a week) is also used. In patients with HIV infection, the drug is indicated if there was a previous episode of PCP or CD4 counts are <200 cells/μL. (See Notes.)

Special Populations. Pediatric Dosage. PO for UTI or shigellosis (2 months–12 yr) 8 mg/kg/day of TMP and 40 mg/kg/day of SMZ (Susp 1 mL/kg/day) in 2 equally divided doses; (>12 yr) same as adult mg/kg dosage. PO for otitis media same as UTI dosage. IV for severe Gram-negative infection or shigellosis (>2 months) same as adult mg/kg dosage. PO or IV for PCP same as adult mg/kg dosage. PO for P. carinii infection prophylaxis 150 mg/m²/day of TMP and 750 mg/m²/day of SMZ, in divided doses, given 3 days a week.

Geriatric Dosage. Reduce dosage for age-related reduction in renal function, although dosage reduction is not necessary with only minor age-related renal function changes. (See Precautions.)

Other Conditions. For a Clr <30 mL/min, give normal dosage for 1–6 doses; with a Clr of 15–30 mL/min, follow with 50% of the usual dosage; with a Clr <15 mL/min, follow with 25–50% of the usual dosage in 1 or 2 divided doses. Give patients on hemodialysis a normal dose after each dialysis procedure. For systemic infections treated with higher dosages, monitor serum levels.

Dosage Forms. Susp 8 mg/mL of TMP and 40 mg/mL of SMZ; Tab 80 mg of TMP and 400 mg of SMZ (single strength), 160 mg of TMP and 800 mg of SMZ (double strength); Inj 16 mg/mL of TMP and 80 mg/mL of SMZ.
Patient Instructions. Take this medication with a full 8 fluid ounce glass of water on an empty stomach (1 hour before or 2 hours after meals) for best absorption. Drink several additional glasses of water daily, unless directed otherwise.

Missed Doses. Take this drug at regular intervals. If you miss a dose, take it as soon as you remember. If it is about time for the next dose, take that dose only. If you are taking the drug once a day, leave at least 10–12 hours between doses. If you are taking the drug twice a day, leave at least 5–6 hours between doses. Do not double the dose or take extra.

Pharmacokinetics. Serum Levels. Trimethoprim levels >5 mg/L (>17 μmol/L) and SMZ peak levels of about 100 mg/L (396 μmol/L) may be required in PCP. Fate. TMP and SMZ are 90–100% absorbed orally. In normal adults, peak serum concentrations of 0.9–1.9 mg/L (3.1–6.5 μmol/L) of TMP and 20–50 mg/L (79–198 μmol/L) of SMZ occur about 1–4 hr after 160 mg of TMP and 800 mg of SMZ. An additional 10–20 mg/L of SMZ exists in the serum as inactive metabolites. IV infusion of 160 mg of TMP and 800 mg of SMZ over 1 hr produces peak serum levels of 3.4 mg/L (11.7 μmol/L) of TMP and 46.3 mg/L (183 μmol/L) of SMZ. TMP and SMZ are widely distributed in the body, although TMP is much more widely distributed because of its greater lipophilicity. TMP is 45% plasma protein bound and has a V₆ of 1–2 L/kg; SMZ is 60% plasma protein bound and has a V₆ of 0.36 L/kg. TMP concentrations in various tissues and fluids (including the prostate, bile, and sputum) are several times greater than concomitant serum concentrations; CSF concentrations in normal adults are approximately 50% of serum concentrations. Nearly all TMP is excreted in the urine within 24–72 hr, 50–75% as unchanged drug. SMZ undergoes extensive liver metabolism, producing N⁴-acetylated and N⁴-glucuronidated derivatives; 85% is excreted in the urine within 24–72 hr, 10–30% as unchanged drug. The pharmacokinetics of these drugs are essentially unchanged when given in combination. The pH of the urine influences renal excretion of both drugs but does not markedly alter overall elimination.

t₁/₂. 11 ± 2.3 hr for TMP and 8 ± 0.4 hr for SMZ in normal adults. 20–30 hr or more for TMP in severe renal failure; 18–24 hr for SMZ in anuria.

Adverse Reactions. GI irritation including nausea, vomiting, and anorexia occurs frequently, and frequency and severity appear to be dose related. Rashes and other hypersensitivity reactions similar to those caused by other sulfonamides occur occasionally. In patients with AIDS, allergic skin reactions, rash (usually diffuse, erythematous or maculopapular, and pruritic) are frequent and might be associated with fever, leukopenia, neutropenia, thrombocytopenia, and increased transaminase levels. Desensitization has been successful. (See Notes.) In patients without underlying myelosuppression and treated with conventional dosages, the frequency of megaloblastic anemia and other hematologic disorders is rare but might be higher in folate-deficient patients. Hepatotoxicity and nephrotoxicity are rare; renal dysfunction can occur in patients with pre-existing renal disease, but it is reversible. Allergic skin reactions, including toxic epidermal necrolysis, exfoliative dermatitis, Stevens-Johnson syndrome, erythema multiforme, and fixed drug eruptions, occur rarely. Other rare adverse effects are cholestatic jaundice,
pancreatitis, pseudomembranous colitis, hyperkalemia, myalgia, headache, insomnia, fatigue, ataxia, vertigo, depression, and anaphylaxis.243

**Contraindications.** Pregnancy; infants <2 months; history of hypersensitivity reaction to sulfonamide derivatives or trimethoprim; megaloblastic anemia caused by folate deficiency. Lactation is stated by manufacturer to be a contraindication, but risk is probably limited to nursing infants <2 months of age.

**Precautions.** G-6-PD deficiency; impaired renal or hepatic function. Adverse reactions can be more frequent in the elderly, especially with impaired hepatic or renal function or in those taking thiazide diuretics.

**Drug Interactions.** The effects of methotrexate, sulfonylureas, and warfarin are increased when used with trimethoprim-sulfamethoxazole. Enhanced bone marrow suppression can occur with the combination of trimethoprim/sulfamethoxazole and mercaptopurine. A decreased effect of cyclosporine and an increased risk of nephrotoxicity can occur. High-dose trimethoprim-sulfamethoxazole with di-danosine can increase the risk of pancreatitis. Phenytoin clearance can be decreased with concurrent use.

**Parameters to Monitor.** Baseline and periodic CBC counts for patients on long-term or high-dose treatment. Monitor SMZ serum levels in patients treated for PCP if absorption is questionable or response is poor. In patients with AIDS, monitor for hypersensitivity skin reactions (rash and urticaria).

**Notes.** Protect all dosage forms from light. The efficacy and safety of TMP and SMZ have been demonstrated in numerous infectious conditions (eg, chronic UTI, chronic bronchitis, sepsis, enteric fever, prostatitis, endocarditis, meningitis, and gonorrhea), and the combination is considered an effective alternative to conventional therapy in most cases.241,246 Efficacy of TMP and SMZ in the treatment of PCP is equivalent to pentamidine, which makes the combination the therapy of choice because of its greater safety and lower cost.245 Oral desensitization or rechallenge with TMP/SMZ has been successful in permitting continued use in patients with AIDS who experience hypersensitivity reactions.248

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**Tetracyclines**

**DOXYCYCLINE AND SALTS**

**Vibramycin**

**Pharmacology.** Tetracyclines are broad-spectrum bacteriostatic compounds that inhibit protein synthesis at the 30S ribosomal subunit. Activity includes Gram-positive, Gram-negative, aerobic, and anaerobic bacteria, as well as spirochetes, mycoplasmas, rickettsiae, chlamydiae, and some protozoa. Many bacteria have developed plasmid-mediated resistance. Most Enterobacteriaceae and *P. aeruginosa* are resistant. Doxycycline is somewhat more active than other tetracyclines against anaerobes and facultative Gram-negative bacilli.249,250

**Administration and Adult Dosage.** PO 100 mg q 12 hr for 2 doses, then 50–100 mg/day in 1 or 2 doses, depending on the severity of the infection, to a maximum of 200 mg/day. PO for uncomplicated chlamydial genital infections 100 mg bid for at least 7 days. PO for primary and secondary syphilis 100 mg tid for at least 10 days. PO for prophylaxis against travelers’ diarrhea 200 mg
en route, then 100 mg/day for duration of travel (6 weeks maximum). **PO for malaria prophylaxis in short-term (<4 months) travelers** 100 mg/day beginning 1–2 days before travel to malarious areas and for 4 weeks after leaving the area. **IV** 200 mg in 1 or 2 divided doses for 1 day, followed by 100–200 mg/day, infused at a concentration of 0.1–1 g/L over 1–4 hr; double maintenance dosage in severe infections. **Intrapleural for pleural effusions** 500 mg in 25–30 mL of NS has been used; most patients require 2–4 infusions for maximum efficacy. Not for SC or IM use.

**Special Populations. Pediatric Dosage.** Not recommended ≤8 yr. PO (>8 yr, <45 kg) 2.2 mg/kg q 12 hr for 2 doses, then 2.2–4.4 mg/kg/day in 1 or 2 divided doses, depending on the severity of the infection; (>45 kg) same as adult dosage.

**PO for malaria prophylaxis in short-term (<4 months) travelers** (>8 yr) 2.2 mg/kg/day to a maximum of 100 mg/day beginning 1–2 days before travel to malarious areas and for 4 weeks after leaving the area. **IV** (<45 kg) 4.4 mg/kg in 1 or 2 divided doses for 1 day followed by 2.2–4.4 mg/kg/day in 1 or 2 divided doses, infused at a concentration of 0.1–1 g/L over 1–4 hr; (>45 kg) same as adult dosage.

**Geriatric Dosage.** Same as adult dosage.

**Other Conditions.** No dosage adjustment is necessary in renal impairment.

**Dosage Forms.** Cap (as hyclate) 20, 50, 100 mg; Tab (as hyclate) 50, 100 mg; Cap (as monohydrate) 50, 100 mg; Susp (as monohydrate) 5 mg/mL (reconstituted); Syrup (as calcium) 10 mg/mL; Inj (as hyclate) 100, 200 mg.

**Patient Instructions.** Take doxycycline by mouth with a full glass of water on an empty stomach; if stomach upset occurs, the drug may be taken with food or milk but not with antacids or iron products. Avoid prolonged exposure to direct sunlight while taking this drug.

**Missed Doses.** Take this drug at regular intervals. If you miss a dose, take it as soon as you remember. If it is about time for the next dose, take that dose only. Leave at least 6–8 hours between doses. Do not double the dose or take extra.

**Pharmacokinetics. Onset and Duration.** Duration of protection against travelers’ diarrhea is about 1 week after drug discontinuation.

**Fate.** About 93% is orally absorbed, producing a peak of 3 mg/L (6.5 μmol/L) 2–4 hr after administration of a 200 mg dose; antacids and iron can markedly impair oral absorption; milk causes about a 30% decrease in bioavailability and food has little effect. Widely distributed in the body, penetrating most cavities including CSF (12–20% of serum levels). The drug is 88 ± 5% plasma protein bound, \( V_d = 0.75 \pm 0.32 \text{ L/kg} \); \( C_l = 0.032 \pm 0.01 \text{ L/hr/kg} \). About 41 ± 19% is excreted unchanged in the urine in normal adults; the remainder is eliminated in feces via intestinal and biliary secretion. \( t_{1/2} = 16 \pm 6 \text{ hr} \) in normal adults; slightly prolonged in severe renal impairment.

**Adverse Reactions.** IV administration frequently produces phlebitis. Oral doxycycline causes less alteration of intestinal flora than other tetracyclines but can cause nausea and diarrhea with equal frequency. It binds to calcium in teeth and bones, which can cause discoloration of teeth in children, especially during...
growth; however, doxycycline has a lower potential for this effect than most other tetracyclines. In contrast to other tetracyclines, doxycycline is not very antiana-

bolic and will not further increase azotemia in renal failure. Phototoxic skin reac-

tions occur occasionally.249,250,252

Contraindications. Hypersensitivity to any tetracycline.

Precautions. Not recommended in pregnancy or in children ≤8 yr because per-

manent staining of the child’s teeth will occur. Use with caution in severe hepatic dysfunc-

tion. The syrup contains sulfites.

Drug Interactions. Antacids containing di- or trivalent cations, bismuth salts, or

zinc salts interfere with absorption of oral tetracyclines. Oral iron salts lower
doxycline serum levels, even of IV doxycycline, by interfering with absorption
and enterohepatic circulation. Barbiturates, carbamazepine, and phenytoin can en-

hance doxycycline hepatic metabolism, possibly decreasing its effect. Tetracy-

clines can interfere with enterohepatic circulation of contraceptive hormones,
causing menstrual irregularities and possibly unplanned pregnancies. Combined
use of tetracyclines with the bactericidal agents such as penicillins can result in
decreased activity in some infections.

Parameters to Monitor. Check for signs of phlebitis daily during IV use.

Notes. Doxycycline is the tetracycline of choice because it is better tolerated than
other tetracyclines, although tetracyclines are the drugs of choice for very few in-
fecions.249,250 Each vial contains 480 mg of ascorbic acid per 100 mg of doxycy-
cline hyclate for injection. (See Tetracyclines Comparison Chart.)

Pharmacology. Tetracycline has an antimicrobial spectrum of activity similar to
that of doxycycline. Current uses are for treatment of infection caused by Chlamy-
dia sp., Mycoplasma sp., and Brucella spp.249,250 It is also used as a treatment for
acne and in some regimens against Helicobacter pylori. (See Gastrointestinal
Drugs, Treatment of Helicobacter pylori Infection in Peptic Ulcer Disease.)

Adult Dosage. PO 1–2 g/day in 2–4 divided doses. Reduce dosage, or preferably
use another drug, in severe renal or hepatic impairment.

Pediatric Dosage. Not recommended ≤8 yr; PO (>8 yr) 25–50 mg/kg/day in
2–4 divided doses.

Dosage Forms. (See Tetracyclines Comparison Chart.)

Pharmacokinetics. Tetracycline is well absorbed from the GI tract. Multivalent
cations chelate tetracyclines and inhibit absorption; warn patients to avoid concur-
rent antacids, dairy products, iron, or sucralfate. The half-life of tetracycline is
about 10 hr, increasing to as high as 108 hr in anuria.52

Adverse Reactions. GI irritation is frequent and can result in esophageal ulceration
if the drug is taken at bedtime with insufficient fluid. Disruption of bowel
flora occurs frequently and can result in diarrhea, candidiasis, or rarely
pseudomembranous colitis. Antianabolic effects produce elevated BUN, hyper-
phosphatemia, and acidosis in patients with renal failure. Acute fatty infiltration
of the liver with pancreatitis occurs rarely with large (>2 g) IV doses, especially in
pregnancy; avoid tetracyclines in pregnancy. Do not give tetracyclines to children <8 yr because of binding of calcium in teeth and resultant discoloration.

**Contraindications.** (See Doxycycline.)

**Precautions.** (See Doxycycline.)

**Drug Interactions.** Oral absorption is markedly inhibited by di- and trivalent cations (eg, antacids, iron salts). (See also Doxycycline Interactions.)
<table>
<thead>
<tr>
<th>DRUG</th>
<th>DOSAGE FORMS</th>
<th>ADULT DOSAGE</th>
<th>PERCENTAGE ORAL ABSORPTION</th>
<th>HALF-LIFE (HOURS)</th>
<th>PERCENTAGE EXCERTED UNCHANGED IN URINE</th>
<th>COMMENTS</th>
</tr>
</thead>
<tbody>
<tr>
<td>Demeclocycline</td>
<td>Cap 150 mg</td>
<td>PO 600 mg/day in 2–4 divided doses.</td>
<td>66</td>
<td>15</td>
<td>40–60</td>
<td>42 Most phototoxic tetracycline; causes nephrogenic diabetes insipidus rarely.</td>
</tr>
<tr>
<td>Hydrochloride</td>
<td>Tab 150, 300 mg.</td>
<td>PO for SIADH 300 mg tid–qid.</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Doxycycline Calcium</td>
<td>Cap 20, 50, 100 mg</td>
<td>PO 100 mg q 12 hr for 2 doses, then 50–100 mg/day in 1–2 divided doses; IV 200 mg in 1–2 divided doses on day 1, then 100–200 mg/day.</td>
<td>93</td>
<td>16 ± 6</td>
<td>12–22</td>
<td>41 Safest in renal failure because of its lack of accumulation and lack of antianabolic effects. Well tolerated when given IV.</td>
</tr>
<tr>
<td>Doxycycline Hyclate</td>
<td>Tab 50, 100 mg</td>
<td>Syrup 10 mg/mL</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Doxycycline Monohydrate</td>
<td>Inj 100, 200 mg</td>
<td>PO or IV 200 mg initially, then 100 mg q 12 hr.</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Minocycline Hydrochloride</td>
<td>Cap 50, 75, 100 mg</td>
<td>Susp 10 mg/mL</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Minocin Various</td>
<td>Inj (IV only) 100 mg</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

(continued)
# Tetracyclines Comparison Chart (continued)

<table>
<thead>
<tr>
<th>Drug</th>
<th>Dosage Forms</th>
<th>Adult Dosage</th>
<th>Percentage Oral Absorption</th>
<th>Half-Life (Hours)</th>
<th>Percentage Excreted Unchanged in Urine</th>
<th>Comments</th>
</tr>
</thead>
<tbody>
<tr>
<td>Oxytetracycline</td>
<td>Cap 250 mg</td>
<td>PO 1–2 g/day in 2–4 divided doses; IM 250 mg once</td>
<td>58</td>
<td>9</td>
<td>47–66</td>
<td>70 Seldom used. IM produces lower serum levels than oral.</td>
</tr>
<tr>
<td>Oxytetracycline Hydrochloride</td>
<td>Inj (IM only, contains 2% lidocaine) 50, 125 mg/mL</td>
<td>daily to 300 mg/day in 2–3 doses.</td>
<td></td>
<td>10.6 ± 5</td>
<td>57–108</td>
<td>60 (See monograph.)</td>
</tr>
<tr>
<td>Oxytetracycline Calcium</td>
<td>Various</td>
<td>Top (soln) for acne apply in the morning and evening.</td>
<td>77</td>
<td>10.6 ± 5</td>
<td>57–108</td>
<td>60 (See monograph.)</td>
</tr>
<tr>
<td>Tetracycline</td>
<td>Cap 100, 250, 500 mg</td>
<td>PO 1–2 g/day in 2–4 divided doses.</td>
<td>77</td>
<td>10.6 ± 5</td>
<td>57–108</td>
<td>60 (See monograph.)</td>
</tr>
<tr>
<td>Tetracycline Hydrochloride</td>
<td>Various</td>
<td>Top Soln 2.2 mg/mL</td>
<td>Top Oint 3%</td>
<td>77</td>
<td>10.6 ± 5</td>
<td>57–108</td>
</tr>
</tbody>
</table>

From reference 52 and product information.
Atovaquone is a highly lipophilic hydroxynaphthoquinone with activity against *Pneumocystis carinii*, *Toxoplasma gondii*, and *Plasmodium* sp. It is a structural analogue of ubiquinone, a small hydrophobic respiratory chain electron carrier molecule found in mitochondria. The mechanism of antipneumocystis activity by atovaquone is unclear but might be inhibition of the mitochondrial electron transport chain, which inhibits pyrimidine synthesis and leads to inhibition of nucleic acid and ATP synthesis.

**Administration and Adult Dosage.** PO for PCP treatment 750 mg bid for 21 days; PO for PCP prophylaxis 1.5 g once daily. (See Notes.)

**Special Populations.** *Pediatric Dosage.* Safety and efficacy not established.

**Geriatric Dosage.** (>65 yr) not evaluated, but dosage adjustment appears not to be necessary.

**Other Conditions.** Dosage alteration is not required with renal or hepatic impairment.

**Dosage Forms.** Susp 150 mg/mL.

**Patient Instructions.** It is extremely important to take this medication with food to increase absorption; failure to do so might limit response to therapy. Shake the suspension gently before use.

**Missed Doses.** Take this drug at regular intervals. If you miss a dose, take it as soon as you remember. If it is about time for the next dose, take that dose only. Leave at least 6–8 hours between doses. Do not double the dose or take extra.

**Pharmacokinetics.** *Serum Levels.* Steady-state serum levels >14 mg/L (>38 μmol/L) are correlated with survival in patients with PCP; serum levels <6 mg/L (<16 μmol/L) might be ineffective.

**Fate.** Atovaquone exhibits slow, irregular absorption, depending on the formulation. A high-fat meal increases absorption of the suspension 2.3-fold compared with the fasting state. A peak concentration of 11.5 mg/L (31 μmol/L) is achieved with a single 750 mg dose of the suspension. Oral administration of 750 mg bid as the suspension produces a steady-state level of 24 mg/L (65 μmol/L). More than 99.9% is protein bound, and the drug does not appear to cross the blood–brain barrier well. It appears to undergo enterohepatic cycling with >94% excreted over 21 days in the feces, with no metabolite identified and <0.6% renally excreted.

\[ t_{1/2} \approx 67 \pm 10 \text{ hr}. \]

**Adverse Effects.** Maculopapular rash occurs frequently, but many patients can continue atovaquone therapy; in most instances, the rash resolves without sequelae. GI disturbances such as abdominal pain, nausea, vomiting, and diarrhea occur in more than 10% of patients. Fever, headaches, and insomnia also have been reported frequently. Elevations of hepatic transaminases and hyponatremia occur frequently (1–10% of patients) but do not require cessation of therapy.
Contraindications. Severe diarrhea or malabsorption syndrome because pre-existing diarrhea is associated with poor outcome, presumably as a result of decreased absorption and serum levels.

Precautions. Lactation. Consider alternative therapy in patients who cannot take the drug with food or with GI disorders that might decrease oral absorption.

Drug Interactions. Rifampin can decrease atovaquone serum levels.


Notes. Use atovaquone only in the treatment of mild to moderate episodes of PCP. Atovaquone has been used for prevention of PCP in patients who cannot tolerate or who have failed other traditional prevention medication; although clinical trials are ongoing the safety, efficacy, and optimal dosage for this indication are not well established.

Malarone tablets contain atovaquone 250 mg and proguanil 100 mg/tablet; Malarone Pediatric Tablets contain atovaquone 62.5 mg and proguanil 25 mg/tablet. Malarone is used for prevention or treatment of chloroquine-resistant malaria.

CHLORAMPHENICOL AND SALTS

Pharmacology. Chloramphenicol is a broad-spectrum bacteriostatic antibiotic isolated from Streptomyces venezuelae and is particularly useful against ampicillin-resistant Haemophilus influenzae, Salmonella sp., rickettsial infections such as Rocky Mountain Spotted Fever, typhoid fever, most anaerobic organisms, and many vancomycin-resistant enterococci. It inhibits protein synthesis by binding the 50S ribosomal subunit and might be bactericidal against some bacteria including pneumococci, meningococci, and H. influenzae. Resistance occurs because of impermeability of the cell wall or bacterial production of chloramphenicol acetyltransferase, a plasmid-mediated enzyme that acetylates chloramphenicol into a microbiologically inert form.\textsuperscript{209,250,256,257}

Administration and Adult Dosage. PO or IV 50–100 mg/kg/day in 4 divided doses, depending on severity, location, and organism, to a maximum of 4 g/day. IM not recommended.

Special Populations. Pediatric Dosage. PO or IV (<7 days or <2 kg) 25 mg/kg once daily; (neonates >7 days and >2 kg) 25 mg/kg q 12 hr; (older infants and children) 50–100 mg/kg/day given q 6 hr. These regimens produce unpredictable levels, and serum level monitoring is recommended.\textsuperscript{253} IM not recommended.

Geriatric Dosage. Same as adult dosage.

Other Conditions. Reduce dosage with impaired liver function as guided by serum levels; no alteration necessary in impaired renal function.\textsuperscript{253}

Dosage Forms. Cap 250 mg; Inj (as sodium succinate) 1 g (100 mg/mL when reconstituted); Ophth Oint 10 mg/g; Ophth Pwdr for Soln 25 mg/vial; Ophth Soln 5 mg/mL.

Pharmacokinetics. Serum Levels. Therapeutic peak 10-20 mg/L; therapeutic trough 5–10 mg/L. (See Adverse Reactions.)
Fate. Well absorbed orally with 75–90% bioavailability and a peak serum level of 12 mg/L after administration of 1 g to adults. IV 1 g produces levels of 5–12 mg/L (15–37 μmol/L) 1 hr after administration to normal adults. In infants and young children, hydrolysis of succinate to the active form can be slow and incomplete. IM administration produces serum levels of active drugs that are 50% lower than the equivalent oral dose. The drug attains therapeutic levels in most body cavities, the eye, and CSF; it is 53% plasma protein bound. $V_d$ is $0.94 \pm 0.06$ L/kg; $Cl$ is $0.14 \pm 0.01$ L/hr/kg. Most of the drug is eliminated by glucuronidation in the liver followed by excretion in the urine; the remainder is excreted in the urine unchanged. The rate of glucuronidation and renal elimination is greatly reduced in neonates; 6.5–80% of succinate can be excreted unhydrolyzed. Urine concentrations can be inadequate to treat UTIs, especially in patients with moderately to severely impaired renal function. A small amount (2–4%) of a dose appears in the bile and feces, mostly as the glucuronide. $t_{1/2}$ is 4 ± 2 hr in healthy adults; extremely prolonged and variable in neonates, infants, and young children. Unpredictable in patients with impaired liver function. Some normal patients and patients with impaired renal function exhibit impaired free drug elimination.

Adverse Reactions. Serum levels >25 mg/L (>77 μmol/L) frequently produce reversible bone marrow depression with reticulocytopenia, decreased hemoglobin, increased serum iron and iron-binding globulin saturation, thrombocytopenia, and mild leukopenia. The drug inhibits iron uptake by bone marrow, and anemic patients do not respond to iron or vitamin B12 therapy while receiving chloramphenicol. This anemia most often follows parenteral therapy, large dosages, long duration of therapy, or impaired drug elimination. Complete recovery usually occurs within 1–2 weeks after drug discontinuation. Aplastic anemia occurs rarely (1/12,000 to 1/50,000) and can be fatal. It is not dose related and can occur long after a short course of oral or parenteral therapy; its occurrence after ophthalmic or parenteral use is controversial. Fatal cardiovascular-respiratory collapse (gray syndrome) can develop in neonates given excessive dosages. This syndrome is associated with serum levels of about 50–100 mg/L (155–310 μmol/L). A similar syndrome has been reported in children and adults given large overdoses.

Contraindications. Trivial infections; prophylactic use; uses other than those for which it is indicated.

Precautions. Pregnancy; lactation. Use with caution in patients with liver disease (especially cirrhosis, ascites, and jaundice) or pre-existing hematologic disorders or patients receiving other bone marrow depressants. It can cause hemolytic episodes in patients with G-6-PD deficiency; observe dosage recommendations closely in neonates and infants.

Drug Interactions. Chloramphenicol inhibits CYP2C9 and increases serum concentrations of phenytoin, warfarin, and sulfonylurea oral hypoglycemic agents. Phenytoin, phenobarbital, and rifampin can decrease serum levels of chloramphenicol.

Parameters to Monitor. CBC with platelet and reticulocyte counts before and frequently during therapy; serum iron and iron-binding globulin saturation also
might be useful. Liver and renal function tests before and occasionally during therapy. Monitor serum levels weekly because of variability in Pharmacokinetics. More frequent monitoring might be necessary in patient with hepatic dysfunction and during long-term (>2 weeks) therapy.

**Pharmacology.** Clindamycin is a semisynthetic 7-chloro, 7-deoxylincomycin derivative that is active against most Gram-positive organisms except enterococci and *Clostridium difficile*. Gram-negative aerobes are resistant, but most anaerobes are sensitive. It inhibits bacterial protein synthesis by binding to the 50S ribosomal subunit, it is bactericidal or bacteriostatic depending on the concentration, organism, and inoculum.260,261

**Administration and Adult Dosage.** PO 150–450 mg q 6-8 hr; PO for prevention of endocarditis in patients at risk undergoing dental, oral, or upper respiratory tract procedures and who are allergic to penicillin 600 mg 1 hr before procedure.219 IM or IV 600 mg–2.7 g/day in 2–4 divided doses, to a maximum of 4.8 g/day. IV for endocarditis prophylaxis 600 mg within 30 min before a dental procedure. Single IM doses >600 mg are not recommended; infuse IV no faster than 30 mg/min. Top for acne apply bid. Vag for bacterial vaginosis 1 applicatorful hs for 7 days.

**Special Populations.** Pediatric Dosage. PO (<10 kg) give no less than 37.5 mg q 8 hr; (>10 kg) 8–25 mg/kg/day in 3 or 4 divided doses; PO for endocarditis prophylaxis 20 mg/kg 1 hr before a dental procedure. IM or IV (<1 month) 15–20 mg/kg/day in 3 or 4 divided doses; the lower dosage may be adequate for premature infants; (>1 month) 15–40 mg/kg/day in 3 or 4 divided doses (not less than 300 mg/day in severe infection, regardless of weight). IV for endocarditis prophylaxis 20 mg/kg within 30 min before a dental procedure.

**Geriatric Dosage.** Same as adult dosage.

**Other Conditions.** Dosage adjustment is unnecessary in renal impairment or cirrhosis, although the effect of acute liver disease is unknown.253,260,262

**Dosage Forms.** Cap (as hydrochloride) 75, 150, 300 mg; Soln (as palmitate) 15 mg/mL (reconstituted); Inj (as phosphate) 150 mg/mL; Top Soln (as phosphate) 1%; Top Gel (as phosphate) 1%; Vag Crm 2%.

**Patient Instructions.** Report any severe diarrhea or blood in the stools immediately and do not take antidiarrheal medication. Do not refrigerate the reconstituted oral solution because it will thicken.

**Missed Doses.** Take this drug at regular intervals. If you miss a dose, take it as soon as you remember. If it is about time for the next dose, take that dose only. Leave at least 4–6 hours between doses. Do not double the dose or take extra.

**Pharmacokinetics.** Fate. Absorption is nearly 87% and is the same from the capsule or the solution; food can delay, but not decrease, absorption. The palmitate and phosphate esters are absorbed intact and rapidly hydrolyzed to the active base. Unhydrolyzed phosphate ester usually constitutes <20% of the total peak serum level after parenteral clindamycin but can increase to 40% in patients with im-
paired renal function. A 500 mg oral dose produces a peak serum level of 5–6 mg/L (12–14 μmol/L) in 1 hr. A 300 mg IM dose produces a peak level of 5–6 mg/L 1–2 hr postinjection. A 600 mg IV dose infused over 30 min produces a peak serum level of 10 mg/L (23 μmol/L). The drug is widely distributed throughout the body except the CSF. It is 94% plasma protein bound; Vd is 1.1 ± 0.3 L/kg; Cl is 0.28 ± 0.08 L/hr/kg. There is hepatic metabolism and excretion of active forms in the bile. From 5% to 10% of the absorbed dose is recovered as unchanged drug and active metabolites in the urine within 24 hr.52,253,260,263

$\frac{1}{2}$, 2.9 ± 0.7 hr; increased in premature infants;52 unchanged or slightly increased in severe renal disease; might be increased or unchanged in liver disease.261

**Adverse Reactions.** After oral administration, anorexia, nausea, vomiting, cramps, and diarrhea occur frequently.253,260,262 Oral and rarely parenteral clindamycin can cause severe, sometimes fatal, pseudomembranous colitis (PMC), which might be clinically indistinguishable at onset from non-PMC diarrhea.260 Antibiotic-associated PMC is secondary to overgrowth of toxin-producing *Clostridium difficile*. Symptoms usually appear 2–9 days after initiation of therapy. PMC has been reported after topical administration.262 PMC is terminated in many patients by discontinuing the antibiotic immediately; however, if diarrhea is severe or does not improve promptly after discontinuation, treat with oral metronidazole or vancomycin.260,261 The value of corticosteroids, cholestyramine, and antispasmodics in the management of antibiotic-associated diarrhea and PMC has not been established.260 Antidiarrheals such as diphenoxylate or loperamide may worsen PMC and should not be used.

**Precautions.** Pregnancy; lactation. Use with caution in neonates <4 weeks of age and in patients with liver disease. Discontinue immediately if severe diarrhea occurs. Drug accumulation might occur in patients with severe concomitant hepatic and renal dysfunction, but data are lacking.

**Drug Interactions.** Clindamycin might enhance the action of nondepolarizing neuromuscular blocking agents. Kaolin-pectin mixture delays but does not decrease oral absorption of clindamycin.

**Parameters to Monitor.** Observe for changes in bowel frequency.

**Notes.** Oral solution is stable for 2 weeks at room temperature after reconstitution; do not refrigerate.

**LINEZOLID**

**Pharmacology.** Linezolid belongs to a new class of anti-infective agents known as oxazolidinones. It inhibits protein synthesis by binding to the bacterial 23S ribosomal RNA of the 50S subunit and prevents the formation of a functional 70S initiation complex inhibiting bacterial translation. It has bacteriostatic activity against staphylococci and enterococci including vancomycin-resistant *Enterococcus faecium* and *faecalis* and bactericidal activity against most streptococcal strains. In vitro the spectrum also includes certain Gram-negative and anaerobic organisms. Linezolid is a reversible, nonselective inhibitor of monoamine oxidase.264

**Administration and Adult Dosage.** PO or IV 400–600 mg q 12 hr.
Special Populations. Pediatric Dosage. Safety not established in infants and children. PO or IV a dosage of 10 mg/kg q 12 hr has been used.

Geriatric Dosage. Same as adult dosage.

Other Conditions. No dosage adjustment necessary in renal or hepatic insufficiency.

Dosage Forms. Tab 400, 600 mg; Inj 2 mg/mL; Susp 20 mg/mL.

Patient Instructions. This drug may be taken without regard to meals. Avoid concurrent use of diet pills and cough-and-cold remedies and restrict consumption of aged foods high in tyramine. (See Foods That Interact with MAO Inhibitors Chart in the Antidepressants chapter.)

Missed Doses. Take this drug at regular intervals. If you miss a dose, take it as soon as you remember. If it is about time for the next dose, take that dose only.

Pharmacokinetics. Fate. Rapidly absorbed orally; bioavailability is approximately 100% and not affected by food. A single dose of 600 mg achieves a peak of 12.7 mg/L when administered orally and 12.9 mg/L IV. Plasma protein binding is 31% and linezolid is readily distributed into well-perfused tissues. Linezolid is primarily metabolized by oxidation. Nonrenal clearance accounts for approximately 65% of the total clearance. Children appear to have a higher average clearance.

$t_{1/2}$. 4.7–5.4 hr in adults.

Adverse Reactions. Adverse reactions are usually mild to moderate and the most commonly reported are diarrhea, headache, and nausea. Occasional reactions are oral and vaginal candidiasis, hypertension, dyspepsia, abdominal pain, pruritus, and tongue discoloration. Treatment periods beyond 28 days have not been evaluated and are not recommended.

Precautions. Pregnancy; lactation. Linezolid can lead to pseudomembranous colitis, so it is an important consideration if patients present with diarrhea. Avoid large quantities of food containing tyramine (>100 mg/meal) with linezolid. Use caution with pre-existing myelosuppression, other drugs that cause myelosuppression, or chronic infection with previous or concomitant antibiotic therapy. Myelosuppression (including anemia, leukopenia, pancytopenia and thrombocytopenia) has been reported; consider discontinuing therapy if this occurs or worsens. Myelosuppression is usually reversible after drug discontinuation.

Drug Interactions. Linezolid is not metabolized by cytochrome P450 and does not inhibit or induce the activities of clinically important CYP isoforms. By inhibiting MAO, it can interact with adrenergic and serotonergic agents such as phenylpropanolamine and pseudoephedrine; reduce initial doses of epinephrine and dopamine and titrate to response.

Parameters to Monitor. CBC with platelet counts before and during weekly therapy.

Notes. Although the drug is effective for many types of infections, it should generally be reserved for treating resistant organisms.
Pharmacology. Metronidazole is a synthetic nitroimidazole active against *Trichomonas vaginalis* (trichomoniasis), *Entamoeba histolytica* (amebiasis), and *Giardia lamblia* (giardiasis); it is bactericidal against nearly all obligate anaerobic bacteria including *Bacteroides fragilis*. It is inactive against aerobic bacteria and requires microbial reduction by a nitroreductase enzyme to form highly reactive intermediates that disrupt bacterial DNA and inhibit nucleic acid synthesis, leading to cell death.  

Administration and Adult Dosage. PO or IV for anaerobic infections 15 mg/kg (usually 1 g) initially, followed by 7.5 mg/kg (usually 500 mg) q 6–8 hr, to a maximum of 2 g/day. Infuse each IV dose over 1 hr. PO for antibiotic-associated colitis 250 mg qid for 7–10 days. PO for trichomoniasis 2 g as a single dose or in 2 doses on the same day, or 500 mg bid for 7 days. PO for giardiasis 250 mg tid for 5 days. PO for symptomatic intestinal amebiasis (amebic dysentery) 750 mg tid for 10 days. PO for extraintestinal amebiasis 750 mg tid for 10 days; some practitioners include a drug effective against the intestinal cyst form because occasional failures with metronidazole therapy have been reported. PO for bacterial vaginosis 500 mg bid for 7 days, or 2 g as a single dose. PO for giardiasis 250 mg tid for 5 days. PO for symptomatic intestinal amebiasis (amebic dysentery) 15 mg/kg/day in 3 divided doses for 5 days, to a maximum of 750 mg/day. PO for amebic dysentery or extraintestinal amebiasis 35–50 mg/kg/day in 3 divided doses for 10 days, to a maximum of 2.5 g/day.

Special Populations. Pediatric Dosage. IV for anaerobic infections (preterm infants) 15 mg/kg once, then 7.5 mg/kg q 24–48 hr; (term infants) 15 mg/kg once, then 7.5 mg/kg q 12–24 hr; (infants >1 week old and children) same as adult mg/kg dosage. PO for giardiasis 15 mg/kg/day in 3 divided doses for 5 days, to a maximum of 750 mg/day. PO for extraintestinal amebiasis 35–50 mg/kg/day in 3 divided doses for 10 days, to a maximum of 2.5 g/day.

Geriatric Dosage. (>65 yr) decreased clearance can result in accumulation of the drug. Dosage reduction or changing dosage interval to once or twice daily are reasonable modifications to avoid potential adverse reactions. Other Conditions. No dosage alteration required with renal impairment. Patients with substantial liver dysfunction metabolize metronidazole slowly, with resultant accumulation of metronidazole and its metabolites in the serum. For such patients, it has been suggested that dosage intervals be increased to 12–24 hr, although specific guidelines are not available.  

Dosage Forms. Cap 375 mg; SR Tab 750 mg; Tab 250, 500 mg; Inj 500 mg; Crm 1%; Top Gel 0.75%; Vag Gel 0.75%.

Patient Instructions. This drug may be taken with food to minimize stomach upset. It can cause a harmless dark discoloration of the urine and metallic taste in the mouth. Nausea, vomiting, flushing, and faintness can occur if alcohol is taken during therapy with this drug.

Missed Doses. Take this drug at regular intervals. If you miss a dose, take it as soon as you remember. If it is about time for the next dose, take that dose only. Leave at least 4–6 hours between doses. Do not double the dose or take extra.
Pharmacokinetics. Serum Levels. Not used clinically.

Fate. IV 500 mg q 12 hr over 1 hr produces steady-state peak and trough levels of 23.6 mg/L (138 μmol/L) and 6.7 mg/L (39 μmol/L), respectively. IV 500 mg q 8 hr over 1 hr produces steady-state peak and trough levels of 27.4 mg/L (160 μmol/L) and 15.5 mg/L (91 μmol/L), respectively. Well absorbed orally with levels similar to those after IV infusion; 250 and 500 mg doses produce peak concentrations of 4–6 mg/L (23–35 μmol/L) and 10–13 mg/L (58–76 μmol/L), respectively, at 1–2 hr in adults. Bioavailability of vaginal gel is 53–58%. Less than 20% plasma protein bound; wide distribution with therapeutic levels in many tissues, including abscesses, bile, bone, breast milk, CSF, and saliva. Vd is 0.85 ± 0.25 L/kg; Cl is 0.07 ± 0.02 L/hr/kg. Extensively metabolized in the liver by hydroxylation, oxidation, and glucuronide formation; 44–80% excreted in the urine in 24 hr, about 6–18% as unchanged drug.265,266 

t 1/2. 6–10 hr in adults; not increased with impaired renal function; prolonged variably with severe hepatic impairment.52,265,266

Adverse Effects. Metallic taste in mouth and GI complaints occur frequently with high dosages. Occasional dizziness, vertigo, and paresthesias have been reported with very high dosages. Reversible mild neutropenia reported occasionally.209,265 Reversible, rare, but severe peripheral neuropathy can occur with high dosages given over prolonged periods. Antibiotic-associated colitis has been reported rarely with oral metronidazole. The IV preparation is occasionally associated with phlebitis at the infusion site. Experimental production of tumors in some rodent species and mutations in bacteria have raised concern regarding potential carcinogenicity; to date, human epidemiologic research has not detected an appreciable risk, although more data are needed.265

Contraindications. First trimester of pregnancy, although there is no direct evidence of teratogenicity in humans or animals.209

Precautions. Pregnancy; lactation; active CNS disease or neutropenia; hepatic impairment.

Drug Interactions. Disulfiram-like reactions are reported with concurrent alcohol use but are uncommon. Confusion and psychotic episodes have been reported with concurrent disulfiram; avoid this combination, if possible. Metronidazole inhibits CYP2C9, CYP3A3/4, and CYP3A5-7 and can affect the metabolism of many drugs; the best documented is an enhanced hypoprothrombinemic response to warfarin. Phenytoin metabolism also might be inhibited. It is also a substrate of CYP2C9.

Parameters to Monitor. Before and after the completion of any lengthy or repeated courses of therapy, monitor WBC count. Monitor signs of toxicity in patients with severe liver disease.266

Notes. The treatment of asymptomatic trichomoniasis is controversial. Signs of endocervical inflammation or erosion on physical examination are considered an indication for treatment. Also, most practitioners treat asymptomatic male consorts because lack of such treatment might be a cause of treatment failure or recurrent infection of the female partner.195 Metronidazole has been used in combination regimens to treat Helicobacter pylori–infected patients with duodenal or
gastric ulcers. (See Gastrointestinal Drugs, Treatment of *Helicobacter pylori* in Peptic Ulcer Disease Comparison Chart.) Although it is slightly less effective than **vancomycin**, metronidazole is considered by some to be the drug of choice for antibiotic-associated pseudomembranous colitis because of its lower cost and the emergence of vancomycin-resistant enterococci.

**NITROFURANTOIN**

**Pharmacology.** Nitrofurantoin is a synthetic nitrofuran that is active against most bacteria that cause UTIs except *P. aeruginosa*, *Proteus* sp., many *Enterobacter* sp., and *Klebsiella* spp. The drug is used primarily to prevent recurrent UTIs but is also effective in the treatment of uncomplicated UTIs.

**Adult Dosage.** PO for UTI (macrocrystals) 50–100 mg qid with meals and hs for treatment; (Macrobid) 100 mg bid for 7 days. PO for chronic suppression 50–100 mg hs. The drug should be taken with food.

**Pediatric Dosage.** PO for treatment of UTI 5–7 mg/kg/day in 4 divided doses, to a maximum of 400 mg/day; PO for chronic UTI suppression 1 mg/kg/day in 1–2 doses, to a maximum of 100 mg/day.

**Dosage Forms.** Cap (macrocrystals) 25, 50, 100 mg; Susp 5 mg/mL; Cap 100 mg, containing 25 mg as macrocrystals and 75 mg in an SR form (Macrobid).

**Pharmacokinetics.** Well absorbed orally; however, serum and extraurinary tissue concentrations are subtherapeutic. About 60% of drug is metabolized to inactive metabolites; 25–35% is excreted in urine with a urine concentration of about 200 mg/L from an average dose.

**Adverse Reactions.** Adverse effects are primarily nausea, vomiting, and diarrhea and are dose related; use of the macrocrystalline form and administration with food can minimize GI distress. Hypersensitivity reactions such as rash occur only rarely. Acute allergic pneumonitis is reversible with discontinuation of therapy. Chronic interstitial pulmonary fibrosis also occurs occasionally with long-term therapy and might be irreversible. Ascending polyneuropathy associated with prolonged high-dose therapy or use of the drug in renal failure is only slowly reversible. Intravascular hemolysis can occur in patients with severe G-6-PD deficiency. Although the drug is mutagenic in mammalian cells, there is no clinical evidence of carcinogenicity or teratogenicity.

**PENTAMIDINE ISETHIONATE**

**Pharmacology.** Pentamidine is an aromatic diamidine used in the treatment of trypanosomiasis and PCP. Pentamidine inhibits dihydrofolate reductase, interferes with anaerobic glycolysis, inhibits oxidative phosphorylation, and limits nucleic acid and protein synthesis, but the mechanism by which pentamidine kills *P. carinii* is unclear.

**Administration and Adult Dosage.** IV (preferred) or IM 3–4 mg/kg/day as a single dose for 2–3 weeks; infuse IV over 60 min. **Inhal for PCP prophylaxis in high-risk HIV-infected patients** 300 mg q 4 weeks via Respirgard II nebulizer. (See Notes.)

**Special Populations.** **Pediatric Dosage.** Same as adult dosage.
Geriatric Dosage. Same as adult dosage.

Other Conditions. Dosage adjustment does not appear necessary in renal impairment.270

Dosage Forms. Inj 300 mg; Inhal 300 mg.

Pharmacokinetics. Serum Levels. Not used clinically.

Fate. Negligible oral absorption. Peak serum levels of 0.5–3 mg/L (1.5–8.8 µmol/L) occur after 4 mg/kg IV infusion. Serum levels are very low after inhalation (<0.1 mg/L). About 70% plasma protein bound; distributed widely in tissues, with highest concentrations found in spleen, liver, kidneys, and adrenal glands. Vc is 3 L/kg; terminal Vd is 190 ± 70 L/kg; Cl is 1.08 ± 0.42 L/hr/kg. There are no data on the effects of liver impairment. Less than 20% of a dose is excreted unchanged in urine.52,271

t¹⁄₂. α phase 1.2 ± 0.6 hr; terminal elimination half-life is up to 29 ± 25 days, suggesting rapid tissue uptake with slow release and subsequent urinary excretion.270

Adverse Reactions. With IV administration, nephrotoxicity occurs in up to 25% of patients, hypoglycemia in up to 27%, and hypotension in up to 10% of patients. Fever, rash, leukopenia, and liver damage occur occasionally. Hyperglycemia, type 1 diabetes mellitus, and pancreatitis have been reported. Pentamidine-induced torsades de pointes occurs rarely. IM injection frequently produces pain and abscess formation at the injection site. With aerosolized pentamidine, reversible bronchoconstriction and unpleasant taste occur frequently. Severe adverse reactions are less frequent, but reports of pancreatitis, hyperglycemia, and cutaneous eruptions have occurred rarely, suggesting some systemic absorption.269

Precautions. Use with caution in diabetes mellitus.

Drug Interactions. IV pentamidine can increase the risk of hypocalcemia with foscarnet; avoid this combination, if possible, although inhaled pentamidine does not seem to be a risk factor.

Parameters to Monitor. Obtain serum glucose, Crs, BUN, liver function tests, electrolytes, and CBC and platelet counts daily. Monitor blood pressure after administration.

Notes. Concomitant therapy with pentamidine and trimethoprim-sulfamethoxazole appears to offer no benefit and might be additively toxic. There is concern about occupational exposure with inhalation therapy. No studies have determined the health effects of exposure to pentamidine itself; however, transmission of tuberculosis to health care workers has been attributed in part to the use of aerosolized pentamidine among clinic patients coinfected with HIV and tuberculosis. Health care workers administering aerosolized pentamidine should wear masks and protective eye wear.269

QUINUPRISTIN AND DALFOPRISTIN

Pharmacology. Quinupristin and dalfopristin are streptogramin antibiotics that are naturally occurring compounds isolated from Streptomyces pristinaspiralis. Quinupristin, a derivative of pristinamycin IA, and dalfopristin, a derivative of pristinamycin IIA, are combined in a fixed ratio of 30:70 (w/w). This combination
inhibits protein synthesis by sequential binding to the 50S subunit of bacterial ribosomes; its synergistic activity can be caused by binding of dalfopristin, altering conformation of the ribosome such that its affinity for quinupristin is increased. Individually, pristinamycin I and II are bacteriostatic, but in combination they are bactericidal against Gram-positive bacteria, including MRSA. Synergy has been reported with vancomycin against MRSA and multiply resistant enterococci. It also has activity against anaerobic organisms, but most Gram-negative organisms such as the Enterobacteriaceae, *Acinetobacter* spp., and *P. aeruginosa* are resistant. Although the drug is effective for many types of infections, it should generally be reserved for treating resistant organisms such as vancomycin-resistant *Enterococcus faecium*. It has no activity against *Enterococcus faecalis*.

**Adult Dosage.** IV 7.5 mg/kg (of the combination) q 8–12 hr infused in D5W over 60 min. Consider reducing dosage in patients with hepatic impairment who do not tolerate the usual dosage. However, specific guidelines have not been established.

**Dosage Forms.** Inj 500 mg (quinupristin 150 mg and dalfopristin 350 mg)/10 mL vial.

**Pharmacokinetics.** The pharmacokinetics are complex and not fully elucidated. Peak concentrations are 2.4–2.8 mg/L (2.3–2.7 μmol/L) for quinupristin and 6.2–7.2 mg/L (9–10.4 μmol/L) for dalfopristin after a 7.5 mg/kg dose in healthy volunteers. Quinupristin is about 90% protein bound and dalfopristin is 10–36% bound in vitro. Clearance of both drugs decreases with repeated doses and in obese patients. Dalfopristin might have an active metabolite. Both drugs are eliminated primarily in feces, with only about 15–20% excreted unchanged in urine. Half-lives are about 1 hr for quinupristin and 0.5–1 hr for dalfopristin, both possibly increased in cirrhosis.

**Adverse Reactions.** Mild to moderate local reactions of itching, pain, and burning at the injection site are frequent and often lead to drug discontinuation. To avoid such side effects, administer the drug through a central venous catheter. Nausea, vomiting, diarrhea, and headache also have been reported frequently. Occasionally, reversible myalgia and arthralgia occur and liver function tests are increased.

**Drug Interactions.** Quinupristin/dalfopristin inhibits CYP3A4 and markedly impairs cyclosporine clearance, requiring cyclosporine dosage reduction.

**TELITHROMYCIN (Investigational—Aventis) Ketek**

**Pharmacology.** Telithromycin is a ketolide antibiotic, a class similar to macrolides with a similar mechanism of action. It has good activity against Gram-positive organisms, especially respiratory pathogens such as *S. aureus, S. pneumoniae, H. influenzae, M. catarrhalis* and some atypical organisms and anaerobes. It is active against some macrolide-resistant Gram-positive cocci.

**Adult Dosage.** PO for community-acquired pneumonia 800 mg once daily. No change required in renal or hepatic dysfunction.

**Dosage Forms.** Tab (investigational).
Pharmacokinetics. Following an 800 mg oral dose, a peak level of 2.3 mg/L occurs in 1 hr. It is primarily metabolized in the liver and about 18% is excreted unchanged in urine. Terminal half-life is about 10 hours with single doses, and about 13 hr with multiple doses.

Adverse Reactions. The most frequent adverse reactions are nausea, diarrhea and GI pain similar to the macrolides. Elevated LFTs and hepatotoxicity reported.

**TRIMETHOPRIM**

Pharmacology. Trimethoprim is a synthetic folate-antagonist antibacterial. (See Trimethoprim and Sulfamethoxazole.) Trimethoprim is effective in acute UTI. It has a potential advantage over the sulfa-containing combination in patients with allergy or toxicity attributed to sulfonamides; however, the relative potential for trimethoprim alone to permit the development of resistance is undetermined. Used alone, trimethoprim is ineffective against *Pneumocystis carinii*, but in combination with dapsone (a sulfone), it is effective in treating mild to moderate PCP.246,247

Adult Dosage. PO for uncomplicated acute UTI 200 mg/day in 1 or 2 doses for 10 days. PO for the treatment of mild to moderate (PaO₂ >60 mm Hg) PCP 20 mg/kg/day in 3 or 4 divided doses with dapsone 100 mg once daily.

Dosage Forms. Tab 100, 200 mg; Soln 10 mg/mL.

Pharmacokinetics. Trimethoprim is rapidly absorbed orally. A 100 mg dose yields a serum concentration of 1 mg/L (3.4 μmol/L) 1–4 hr after the dose. It is 40% plasma protein bound and 50–60% is excreted unchanged in urine. Half-life is 8–10 hr with normal renal function.

Adverse Reactions. Occasional adverse effects are mild thrombocytopenia, nausea, fever, and rash; the frequency appears to be dose related. Methemoglobinemia and dose-related hemolysis have occurred in patients with G-6-PD deficiency receiving dapsone with trimethoprim; it is important to check G-6-PD status before initiating combination therapy.

**TRIMETREXATE**

Pharmacology. Trimetrexate is a lipophilic analogue of methotrexate that inhibits dihydrofolate reductase, leading to the disruption of purine biosynthesis. It has activity against *Pneumocystis carinii* and *Toxoplasma gondii* and has demonstrated modest efficacy against a number of malignancies. It is approved for the treatment of moderate to severe PCP in immunocompromised patients and has been used investigationally in advanced solid tumors alone or in combination with fluorouracil and leucovorin.275–277

Adult Dosage. IV for moderate to severe PCP 45 mg/m²/day infused over 60–90 min for 21 days. Give IV or PO calcium leucovorin 20 mg/m² q 6 hr concomitantly and continue it for 3 days after the end of trimetrexate administration. IV for colorectal cancer 110 mg/m² on day 1, followed by leucovorin 200 mg/m² and a fluorouracil-leucovorin regimen on day 2. IV for advanced urogenital cancer 8 mg/m²/day for 5 days q 3 weeks.278 Reduce dosage by 50% in patients with Cr₆ >1.6 mg/dL.
Dosage Forms. **Inj** 25 mg.

**Pharmacokinetics.** Trimetrexate has at least 2 metabolites, both of which inhibit dihydrofolate reductase. It is eliminated primarily by hepatic metabolism; less than one-third is excreted unchanged in urine. The elimination half-life is 4–12 hr in patients with AIDS and PCP and 8–26 hr in patients with cancer.

**Adverse Reactions.** The primary toxicity is myelosuppression (neutropenia, thrombocytopenia, and anemia); myelosuppression is minimized with concurrent administration of calcium leucovorin. Hypoalbuminemia (≤3.5 g/dL) or hypoproteinemia (≤6 g/dL) increase the risk of severe or life-threatening myelosuppression, presumably because of increased unbound drug levels. Elevated liver function tests, fever, rash, peripheral neuropathy, mucositis, and nausea or vomiting occur frequently. Hypersensitivity reactions and seizures are reported rarely.

**Pharmacology.** Vancomycin is a glycopeptide that binds irreversibly to the cell wall in a manner slightly different from β-lactams. Many Gram-positive cocci and bacilli, including MRSA and *Clostridium difficile*, are inhibited. Most Gram-negative bacteria are resistant, and vancomycin-resistant enterococci have been reported in association with overuse of vancomycin. Glycopeptide intermediate-resistant *S. aureus* have been reported.

**Administration and Adult Dosage.** IV 20–30 mg/kg/day (usually 2 g/day) in 2–4 divided doses as a dilute infusion over 1–2 hr. PO for staphylococcal enterocolitis 2 g/day in 2–4 divided doses. PO for antibiotic-associated colitis 125–500 mg q 6 hr for 7–10 days; retreat with a longer course if relapse occurs. (See Notes.) Not for IM use.

**Special Populations. Pediatric Dosage.** IV (neonates) 20 mg/kg/day; (older infants and children) 40 mg/kg/day in 2–4 divided doses. PO 10–50 mg/kg/day in 4 divided doses. Not for IM use.

**Geriatric Dosage.** Same as adult dosage but adjust for age-related reduction in renal function.

**Other Conditions.** Adjust dosage carefully in renal impairment; Cl is directly related to Clcr. Anuric patients on hemodialysis have been given the usual dose q 3–7 days. Dosage adjustment is unnecessary in liver disease.

**Dosage Forms.** Cap 125, 250 mg; Susp 1, 10 g; Inj 500 mg, 1, 5, 10 g.

**Patient Instructions.** Report pain at infusion site, dizziness, or fullness or ringing in ears with intravenous use; nausea or vomiting with oral use.

**Missed Doses.** (Oral) if you miss a dose, take it as soon as you remember. If it is about time for the next dose, take that dose only.

**Pharmacokinetics. Serum Levels.** Therapeutic range is not well defined. Otitotoxicity has been associated with high serum concentrations but has been noted at lower levels. Peaks thought to be associated with efficacy are 20–40 mg/L (14–28 μmol/L); troughs >15 mg/L (>10 μmol/L) might be excessive.

**Fate.** Oral absorption is negligible, although appreciable serum levels can be observed in patients with renal dysfunction receiving oral vancomycin for
C. difficile-induced antibiotic-associated colitis. Fecal concentrations with PO 500 mg q 6 hr reach 3 mg/g. IV 500 mg produces serum levels of 6–10 mg/L (4–7 μmol/L) in 1 hr. Plasma protein binding is 30 ± 10%. The drug is widely distributed, except into the CSF, although some success has been reported in the treatment of meningitis, particularly in children. \( V_c \) is 0.1–0.15 L/kg; \( V_d \) is 0.39 ± 0.06 L/kg; CI is 0.084 L/hr/kg with normal renal function. In renal impairment, CI (in mL/min) can be estimated as \( 0.79 \times \text{Clcr (in mL/min)} + 3.5 \). Metabolism and biliary excretion are negligible; 80–90% is excreted unchanged in the urine within 48 hr.\(^\text{52,282} \)

\( t_{1/2} \) β phase 5.6 ± 1.8 hr, 6–10 days with renal impairment. No change with hepatic disease.\(^\text{52,279} \)

**Adverse Reactions.** Chills, fever, nausea, and phlebitis can occur frequently, especially with direct injection of undiluted drug (not recommended). Rapid infusion can cause transient systolic hypotension.\(^\text{283} \) The “red man” or “red neck” syndrome of erythema, pruritus, and localized edema is associated with histamine release caused by rapid infusions of doses ≥500 mg; it often does not occur or is less severe with subsequent doses.\(^\text{284} \) Extravasation causes local tissue necrosis. Ototoxicity (auditory and vestibular) and possibly nephrotoxicity occur but have not been definitely linked to high serum levels.\(^\text{281,285} \) Eosinophilia, neutropenia, and urticarial rashes have been reported frequently. Side effects of vancomycin might not be as prevalent today as in the past, perhaps because of changes in the manufacturing process that eliminated some impurities.\(^\text{286} \)

**Precautions.** Pregnancy. Use with caution in patients with impaired renal function or pre-existing hearing loss or in those receiving other ototoxic or nephrotoxic agents.

**Drug Interactions.** Administration with an aminoglycoside can increase the risk of nephrotoxicity.\(^\text{286} \)

**Parameters to Monitor.** With IV use, obtain initial renal function tests and repeat twice weekly during therapy. Routine monitoring of serum levels in patients with normal renal function is not recommended because it has questionable value, but is often performed.\(^\text{285} \) Check for signs of phlebitis daily.

**Notes.** An alternative agent for treatment or prophylaxis of staphylococcal or streptococcal infections when a less toxic agent is inappropriate (eg, penicillin or cephalosporin allergy, or resistant organisms) or has not produced an adequate therapeutic response. Use of vancomycin in antibiotic-associated colitis is becoming less desirable because of the emergence of vancomycin-resistant enterococci. Reserve vancomycin for cases refractory to metronidazole.\(^\text{267} \)

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270. Lake KD, Peterson CD. A simplified method for initiating vancomycin therapy. 
265. Falagas ME, Gorbach SL. Clindamycin and metronidazole. 
263. Van Arsdel PP et al. The value of skin testing for penicillin allergy diagnosis. 
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Antineoplastics. The agents included in this section are those having widespread use in cancer chemotherapy. Agents with therapeutic importance in small patient populations are not included.

Information on the dosage of these drugs has largely been determined empirically, and clinical investigations are continually being performed to find safer and more effective dosage regimens. Thus, dosages in this section should only be considered as guidelines based on the most widely accepted usage at the time of this writing. Because space does not permit detailed discussions of the toxicity, dosage regimens, and other aspects of these drugs, the reader should become familiar with specific agents before initiating treatment. References are provided in this section for more detailed information concerning the proper and safe use of these agents. Specific investigational protocols, if available, also can provide information that is unavailable from other sources, especially with regard to dosage and regimens.

Cancer chemotherapeutic agents as a class are the most toxic drugs in use. Adverse reactions listed represent those most likely to occur with the usual doses and methods of use. Infrequent, but serious, reactions are also listed; however, the lists of adverse reactions are not comprehensive. Nausea and vomiting are important side effects of these agents that can be adequately treated by current antiemetics alone or in combination. To tailor antiemetic therapy better to the emetic potential of the chemotherapy, a standard rating scale is used in these monographs. Several points to remember are that emetogenicity is dose dependent, combinations of chemotherapeutic agents result in greater emetogenic potential than the drug(s) used alone, and emetogenic potentials are best defined in adults and do not necessarily apply to children. The categories of emetogenicity used are as follows:

<table>
<thead>
<tr>
<th>Emetogenicity Category</th>
<th>Percentage of Patients Affected</th>
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<tbody>
<tr>
<td>High</td>
<td>&gt;90</td>
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<tr>
<td>Moderately high</td>
<td>60–90</td>
</tr>
<tr>
<td>Moderate</td>
<td>30–60</td>
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<tr>
<td>Moderately low</td>
<td>10–30</td>
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<tr>
<td>Low</td>
<td>&lt;10</td>
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Class Instructions. Antineoplastics. This drug is very powerful, and some side effects can be expected to occur with its use. Be sure that you understand the possible benefits and dangers of the drug before you begin to take it.

Cytotoxic Agents. Because this drug can decrease your body’s ability to fight infections, report any signs of infection such as fever, shaking chills, or sore throat immediately. Also report any unusual bruising or bleeding, shortness of breath, or painful or burning urination. Avoid the use of aspirin-containing products, and avoid alcohol or use it in moderation. Nausea, vomiting, or hair loss can sometimes occur with this drug. The severity of these effects depends on the individual, the dosage, and other drugs that might be given at the same time. This drug can cause temporary or sometimes permanent sterility in men and women. It also can cause birth defects if the father is taking the drug at the time of conception or if the mother is taking it any time during pregnancy. If you are breast feeding, this drug might appear in the milk and cause problems in your baby; therefore, use an alternate method of feeding your baby.

Missed Doses. This drug should be taken at regular intervals exactly as prescribed. If a dose is missed, it should be taken as soon as it is remembered. If it is almost time for the next dose, only that dose should be taken and the regular dosage schedule should be resumed. The dose should never be doubled or extra doses taken.

Alkylating Agents

ALTRETAMINE Hexalen
Pharmacology. Altretamine (formerly hexamethylmelamine) acts primarily as an alkylating agent. It is used in combination chemotherapy of ovarian cancer and is active in cervical and lung cancers.3–5

Adult Dosage. PO as a single agent 260 mg/m^2/day in 4 divided doses for as long as 2–5 weeks. Lower dosages are required if altretamine is combined with other myelosuppressive agents.

Dosage Forms. Cap 50 mg.

Pharmacokinetics. Oral bioavailability is incomplete and erratic and may be dose dependent. Altretamine is N-demethylated to pentamethylmelamine by hepatic microsomal enzymes. The serum half-life is 4.7–10.2 hr, with >50% of a dose renally excreted in 24 hr and <1% excreted unchanged.

Adverse Reactions. Nausea, vomiting, and abdominal cramps can be dose limiting in some patients. Neurotoxicity is frequent, including agitation, hallucinations, and confusion; these are reversible and amenable to dosage reduction. Anemia, leukopenia, and thrombocytopenia are typically mild.

BUSULFAN Busulfex, Myleran

CHLORAMBUCIL Leukeran

MELPHALAN Alkeran

Pharmacology. These drugs are water-soluble compounds that alkylate DNA, forming a variety of covalent cross-links. The drugs are polyfunctional and can
form more than one covalent bond to susceptible cell constituents (typically the
N7 position of guanine). They are cell-cycle phase nonspecific and chemically sta-
ble enough for oral absorption before appreciable alkylator activation occurs.

**Administration and Dosage.**

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<thead>
<tr>
<th></th>
<th>BUSULFAN</th>
<th>CHLORAMBUCIL</th>
<th>MELPHALAN</th>
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</thead>
<tbody>
<tr>
<td><strong>Administration</strong></td>
<td>PO.</td>
<td>PO.</td>
<td>PO; IV</td>
</tr>
<tr>
<td><strong>Adult Dosage</strong></td>
<td>Up to 8 mg/day (usually 1–3 mg/day).</td>
<td>0.1–0.2 mg/kg/day for 1 day; or 6–12 mg/day maintenance; or 0.4 mg/kg q 2–4 weeks.⁶</td>
<td>PO 7 mg/m² for 4 days; or 2–4 mg/day maintenance for multiple myeloma.⁷</td>
</tr>
<tr>
<td><strong>Pediatric Dosage</strong></td>
<td>CML 0.06–0.12 mg/kg.</td>
<td>Non-Hodgkin’s lymphoma, CLL, nephrotic syndrome, rheumatoid arthritis (initial) 0.1–0.2 mg/kg/day.</td>
<td>—</td>
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<tr>
<td><strong>Geriatric Dosage</strong></td>
<td>Same as adult dosage, but adjust for age-related reduction in renal function.</td>
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**Special Populations. Other Conditions.** Elimination is significantly correlated with the GFR. Studies in nephrectomized animals demonstrate markedly increased myelotoxicity with unadjusted melphalan doses. Thus, one group currently recommends a 50% decrease in the melphalan dosage for BUN >30 mg/dL or Cr >1.5 mg/dL.⁸ Reduce IV melphalan dosage to 75% of normal for WBC counts of 3000–4000/µL or platelet counts of 75,000–100,000/µL or to 50% for WBC counts of 2000–3000/µL or platelet counts of 50,000–75,000/µL, respectively; do not give it with WBC counts <2000/µL or platelet counts <50,000/µL.⁹

**Dosage Forms.** (Busulfan) Tab 2 mg; Inj 60 mg. (Chlorambucil) Tab 2 mg. (Melphalan) Tab 2 mg; Inj 50 mg.

**Patient Instructions.** (See Antineoplastics Class Instructions.)

**Pharmacokinetics.**

**Fate.**

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<th>BUSULFAN</th>
<th>CHLORAMBUCIL</th>
<th>MELPHALAN</th>
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<tr>
<td><strong>Absorption</strong></td>
<td>Reported by manufacturer to be well absorbed orally.</td>
<td>Oral bioavailability is about 87 ± 20% by radiolabeled drug studies, reduced by 10–20% if ingested with food.</td>
<td>Oral bioavailability erratic and incomplete, (mean of 56%, range 25–89%); some patients have no levels after standard doses.</td>
</tr>
<tr>
<td><strong>Distribution</strong></td>
<td>Homogeneous; good ascites penetration; ( V_d ) is 0.99 ±</td>
<td>( V_d ) is 0.29 ± 0.21 L/kg; 99% plasma protein bound.</td>
<td>( V_d ) is 0.45 ± 0.15 L/kg; 90 ± 5% plasma protein bound.</td>
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</tbody>
</table>
**Adverse Reactions.** Emetic potential is low. Nausea and vomiting are rare with long-term administration, although large single doses can be strongly emetogenic. Dose-limiting toxicity for this group is typically myelosuppression, with nadirs of 14–21 days for leukopenia and thrombocytopenia after pulse dosage regimens; daily administration results in chronic low indices with cumulative effects. Blood counts commonly continue to drop after drug discontinuation; fatal pancytopenia has been reported. Therefore, hematologic assessments are important with long-term daily regimens. There might be some selectivity for different normal cell lines by these drugs; busulfan, and perhaps chlorambucil, selectively depresses granulocytes, relatively sparing platelets and lymphoid elements. The nadir for melphalan can be prolonged (4–6 weeks); continuous administration frequently leads to severe myelosuppression (especially platelets) that continues after the drug is discontinued. Pulmonary fibrosis can occasionally occur with all these drugs, especially busulfan; symptoms include cough, dyspnea, and fever; histopathologic changes include bilateral fibrosis. High-dose glucocorticoid therapy might help early evolving pulmonary disease caused by melphalan and chlorambucil, but “busulfan lung” is usually fatal within 6 months of diagnosis. Busulfan frequently causes hyperpigmentation (especially of intertriginous areas) and broad suppression of testicular, ovarian, and adrenal functions (occasionally leading to Addisonian crisis). Long-term daily administration of these drugs predisposes patients to drug-induced carcinogenesis, often heralded by preleukemic pancytopenia and culminating in acute myelocytic leukemia. Allergic hypersensitivity reported, especially with melphalan. With prolonged use, sterility occurs with all alkylators; women seem more sensitive than men.

<table>
<thead>
<tr>
<th>BUSULFAN</th>
<th>CHLORAMBUCIL</th>
<th>MELPHALAN</th>
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<tr>
<td>0.23 L/kg; ex-</td>
<td>Rapid metabolism to</td>
<td>Not actively metabo-</td>
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<td>tensively bound to</td>
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<td>proteins.</td>
<td>active metabolites.</td>
<td>chemical degradation</td>
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<tr>
<td>Metabolism</td>
<td>Cl is 0.16 ± 0.04</td>
<td>to mono- and dity-</td>
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<td>Extensively meta-</td>
<td>L/hr/kg.</td>
<td>droxy products.</td>
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<td>bolized, major</td>
<td>Cl is 0.31 ± 0.17</td>
<td>Cl is 0.04 to mono-</td>
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<td>fraction as meth-</td>
<td>L/hr/kg.</td>
<td>and dihydroxy products.</td>
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<td>anesulfonic acid.</td>
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<td>Cl is 0.27 ± 0.05</td>
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<tr>
<td>L/hr/kg.</td>
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<tr>
<td>Excretion</td>
<td>Less than 1% ex-</td>
<td>Unchanged drug 24-hr</td>
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<tr>
<td>No unchanged drug</td>
<td>creted unchanged</td>
<td>urinary excretion is</td>
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<td>found in urine;</td>
<td>in urine over 24 hr.</td>
<td>10–15% of a dose.</td>
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<td>however, meta-</td>
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<td>bolites are renally</td>
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<td>excreted.</td>
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</tr>
<tr>
<td>$t_{1/2}$</td>
<td>Rapid initial serum</td>
<td>1.3 ± 0.9 hr (un-</td>
</tr>
<tr>
<td>clearance: 90% of</td>
<td>clearance: 90% of</td>
<td>changed drug); 2.5 hr</td>
</tr>
<tr>
<td>dose after 3 min.</td>
<td>changed drug); 2.5 hr</td>
<td>(major metabolite, an</td>
</tr>
<tr>
<td>$t_{1/2}$ is 2.6 ±</td>
<td>(major metabolite, an</td>
<td>aminophenylacetic</td>
</tr>
<tr>
<td>0.5 hr.</td>
<td>aminophenylacetic</td>
<td>acid derivative).</td>
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<td>acid derivative).</td>
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<tr>
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<td>IV: $t_{1/2a}$ 8 min; $t_{1/2b}$ 1.4 ±</td>
</tr>
<tr>
<td></td>
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<td>0.2 hr.$^{10,13,14}$</td>
</tr>
</tbody>
</table>
Contraindications. Documented hypersensitivity; inadequate marrow reserve.

Precautions. See Special Populations for melphalan use in renal impairment.

Drug Interactions. None known.

Parameters to Monitor. WBC and platelet counts at least monthly; reduce dosage at first sign of appreciable myelosuppression (ie, WBC <3000/μL or platelets <75,000/μL). Conversely, assess patients receiving oral melphalan for evidence of mild to moderate myelotoxicity to ensure that some absorption is occurring.

Pharmacology. Carboplatin is a more stable cyclobutane carboxylato derivative of cisplatin that is slowly activated to expose two DNA binding sites on the platinum II coordinate complex. The drug binds to DNA by both inter- and intrastrand cross-links in a fashion similar to, but more delayed than, that with cisplatin. It is more water soluble and commensurately less nephrotoxic than cisplatin. Action is cell-cycle phase nonspecific.

Administration and Adult Dosage. IV for refractory ovarian cancer 360 mg/m² q 4 weeks. Administration by continuous infusion has been reported but is not commonly used.

Special Populations. Pediatric Dosage. Although not specifically labeled for pediatric use, carboplatin has been safely administered to children. IV for recurrent brain tumors 175 mg/m²/week for 4 weeks.

Geriatric Dosage. Same as adult dosage but adjust for age-related reduction in renal function.

Other Conditions. Reduce dosage in patients with reduced renal function, history of prior myelosuppressive therapy, and/or poor bone marrow reserve. Reduce dosage by about 25% if the prior nadir WBC count was <500/μL or the platelet count was <50,000/μL. When Clcr is 41–59 mL/min, a dose of 250 mg/m² is recommended; for Clcr of 16–40 mL/min, 200 mg/m² is recommended. Two prospectively validated formulas for dosage individualization are available. One formula seeks to achieve different target serum AUC values in untreated or pretreated patients: dose (mg) = AUC × (Clcr + 25), where “desired” AUC ranges are 6–8 mg/mL·min for untreated patients and 4–6 mg/mL·min for previously treated patients. The second does not require Clcr estimates and uses a complex mathematical formula.

Dosage Forms. Inj 50, 150, 450 mg.

Patient Instructions. (See Antineoplastics Class Instructions, particularly regarding infection risk.)

Pharmacokinetics. Fate. About 30% of carboplatin is irreversibly bound to plasma proteins; the half-life of this protein-bound fraction is >5 days. Vd is 16–20 L for carboplatin. Carboplatin is slowly hydrolyzed in vivo to a form with two DNA binding sites; the rate of hydrolysis is much slower than the rate of chloride loss with cisplatin. The free (unbound) fraction of carboplatin and its hydrolyzed species are excreted in urine through glomerular filtration and tubular se-
cretion. Urinary elimination accounts for over 65% of drug elimination in patients with normal renal function.

\( t_{1/2} \) (Unbound) \( \alpha \) phase 90 ± 50 min; \( \beta \) phase 180 ± 50 min.\(^{25}\)

**Adverse Reactions.** The emetic potential is moderately high to high but is much less severe than with cisplatin and is easily controlled with antiemetics. Myelosuppression is the primary dose-limiting effect of carboplatin, and thrombocytopenia tends to be more severe than leukopenia; about 25% of previously untreated and 35% of previously treated ovarian cancer patients experience thrombocytopenia. The thrombocytopenic nadir for carboplatin as a single agent is approximately 21 days, and patients with pre-existing renal dysfunction or poor bone marrow reserve have an increased risk for severe thrombocytopenia. Anemia of a mild degree also can occur in up to 90% of patients; in some studies >40% of patients required transfusions and 5% of patients experienced hemorrhage. Diarrhea, abdominal pain, or constipation occur in 6–17% of patients. Nephrotoxicity occurs in 1–22% of patients. Unlike cisplatin, carboplatin does not cause cumulative damage to renal tubules. Transient decreases of 20–30% in some serum electrolytes occur, specifically magnesium, potassium, sodium, and calcium. Hepatic enzyme elevations occur in one-third of patients, but these elevations are not associated with serious or prolonged liver injury. Peripheral neuropathies occur in <10% of patients; however, the risk increases in patients >65 yr or if large dosages of cisplatin have been administered. CNS symptoms occur in ≤5% patients, and ototoxicity occurs in 1% of patients. Occasional reactions are allergic hypersensitivity, alopecia, and various cardiovascular events (eg, embolism, cerebrovascular accident, cardiac failure).

**Contraindications.** The manufacturer lists pre-existing renal impairment and myelosuppression as contraindications, but the drug has been given with appropriate dosage modification. (See Special Populations, Other Conditions.)

**Precautions.** Use with caution in patients with hearing impairment or reduced renal function, or if extensive prior chemotherapy has been administered. Patients with prior cisplatin therapy are at a higher risk for nephrotoxic and neurotoxic sequelae. Vigorous hydration and diuretics usually are not required with carboplatin.

**Drug Interactions.** Myelotoxicity of carboplatin is additive with other myelotoxic drugs. Concurrent use of other nephrotoxic drugs (eg, aminoglycosides) can delay carboplatin elimination and enhance toxicity. Although not well documented, cisplatin interactions also can occur with carboplatin, but at a lesser intensity.

**Parameters to Monitor.** Measure \( \text{Cl}_{\text{cr}} \) before dosage calculation. Monitor platelet and granulocyte counts and Crs during therapy.

**CISPLATIN**

**Pharmacology.** Cisplatin is a planar coordinate dichlorodiamino compound of platinum in the +II valence state. It is aquated in vivo to a positively charged species that can alkylate nucleophilic sites in DNA such as purine and pyrimidine bases. Its action is cell-cycle phase nonspecific.

**Administration and Adult Dosage.** IV bolus or continuous infusion (usually with aggressive hydration) single doses of up to 120 mg/m\(^2\) have been used.\(^{26}\) IV
in the Einhorn testicular cancer regimen 20 mg/m²/day for 5 days.27 (See Notes.)

**Special Populations.**  
**Pediatric Dosage.** IV 10–20 mg/m²/day for 4–5 days, repeat q 3–4 weeks. IV maximum single dose is 100 mg/m² given q 2–3 weeks.26

**Geriatric Dosage.** Same as adult dosage but adjust for age-related reduction in renal function.

**Other Conditions.** Reduce dosage in renal impairment; specific dosage reduction guidelines have not been established.

**Dosage Forms.** Inj 50, 100 mg.

**Patient Instructions.** (See Antineoplastics Class Instructions.) Be prepared for severe nausea and vomiting after drug administration.

**Pharmacokinetics.**  
**Serum Levels.** In vitro cell culture data suggest cytotoxicity at levels of 50 mg/L for 1 hr or 5 mg/L for 8 hr.

**Fate.** Peak serum levels of free platinum after a 100 mg/m² bolus are about 3.4 mg/L when given with mannitol (12.5 g) and 2.7 mg/L without mannitol.28 Over 90% of platinum is protein bound to RBCs, albumin, and prealbumin. It is freely distributed to most organs including kidneys, liver, skin, and lungs and has minimal accumulation in CSF only after repeated doses. Cumulative 24-hr urinary excretion of platinum is 20% with mannitol, 40% without.

\[ t_{1/2}. (\text{Free platinum}) 59 \text{ min (with mannitol); 48 min (without mannitol).} \]  
Terminal half-life is 58–73 hr, probably reflecting slow release of protein-bound drug.28,29

**Adverse Reactions.** Emetic potential is high. Nausea and vomiting are severe and often prolonged (days) and can be managed with aggressive prophylaxis using a serotonin 5HT₃-antagonist, butyrophenone (eg, droperidol), metoclopramide, a high-dose glucocorticoid, or a combination. Primary toxicity is dose-related nephrotoxicity, especially proximal tubular impairment. Ototoxicity and elevated hepatic enzymes occur frequently; total dose-related hypomagnesemia and severe cumulative peripheral neuropathy occur. Slight leukopenia, thrombocytopenia, and frequent anemia also occur. Epoetin alfa is useful in preventing severe anemia caused by cisplatin. Rare toxicities include transient cortical dysfunction (blindness) and hypersensitivity (including anaphylaxis).

**Contraindications.** Renal insufficiency (Crₜ >1.5–2 mg/dL or Clₜ <60 mL/min); myelosuppression; hearing impairment; previous anaphylaxis. However, some patients with prior anaphylaxis have been successfully retreated with cisplatin and concomitant antihistamine, epinephrine, and glucocorticoid.

**Precautions.** Use with caution in renal impairment and with other nephrotoxic drugs, especially aminoglycosides.30 Assure adequate hydration before administration. Both furosemide and mannitol are used to decrease platinum nephrotoxicity, although each apparently retards free platinum elimination.

**Drug Interactions.** Cisplatin can enhance nephrotoxicity and ototoxicity of the aminoglycosides. Use with ifosfamide can increase nephrotoxicity and potassium and magnesium loss, especially in children. Furosemide ototoxicity might be increased by cisplatin. Cisplatin can increase methotrexate serum levels and its toxic-
ity. Cisplatin can decrease absorption and serum levels of valproic acid. Phenytoin serum levels can be decreased after cisplatin-containing combination regimens.

**Parameters to Monitor.** Assess renal function before each dose (eg, serial BUN or Cr) and serum magnesium levels periodically.

**Notes.** Reconstitute with sterile water; it may then be mixed in saline-containing solutions. It is stable for 24 hr in mannitol. Do not expose solution to metals (eg, metal drippers or cannulae) because platinum can rapidly plate onto these surfaces. Hydrate the patient with at least 1 L of a saline-containing solution with 20 mEq of KCl and 3 g of MgSO₄/L.³¹

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**Pharmacology.** Cyclophosphamide is inactive in vitro and must be enzymatically activated in the liver to yield active alkylating compounds and toxic metabolites.³²

**Cell-cycle phase nonspecific.**

**Administration and Adult Dosage.** IV or PO alone or in combination regimens 250–500 mg/m² q 3–4 weeks. IV (usually) or PO in high-dose intermittent regimens (including bone marrow transplant) maximum of 40–50 mg/kg given once or over 2–5 days, repeat q 2–4 weeks—these doses are not well tolerated orally. IV doses may be given in any convenient volume of all common IV solutions or by IV push. **Continuous daily administration PO** 1–5 mg/kg/day; during continuous therapy, dosage must be individualized based on patient bone marrow response.

**Special Populations.** **Pediatric Dosage.** IV, PO for malignancies same as adult dosage. PO for nephrotic syndrome 2.5–3 mg/kg/day for up to 8 weeks.

**Geriatric Dosage.** Same as adult dosage.

**Other Conditions.** No dosage alteration appears necessary in renal impairment because differences in toxicity between normals and patients with renal failure have not been reported.³³

**Dosage Forms.** Tab 25, 50 mg; Inj 100, 200, 500 mg, 1, 2 g.

**Patient Instructions.** (See Antineoplastics Class Instructions.) Drink 2–3 quarts of fluids daily (1–2 quarts in smaller children) and urinate frequently; do not take oral doses at bedtime. Report any blood in the urine.

**Pharmacokinetics.** Fate. Oral absorption is 74 ± 22%.¹⁰ Metabolized to active compounds (including the highly toxic nonalkylating aldehyde, acrolein, and the principal alkylator, phosphoramid mustard) primarily by hepatic microsomal mixed-function oxidases. Cyclophosphamide is 13% plasma protein bound; its alkylating metabolites are 50% bound. Vd is 0.78 ± 0.57 L/kg for parent drug; Cl is 0.078 ± 0.03 L/hr/kg.¹⁰ Renal elimination accounts for 6.5 ± 4.3% of unchanged drug and 60% of metabolites,³⁴ with a mean renal clearance of 0.66 L/hr of unchanged drug.³³ Clearance may be reduced in obese patients. Elimination is linear over a wide range of doses.³²

\[ t_{1/2} \] (Serum alkylating activity) 7.5 ± 4 hr, slightly longer in patients on allopurinol or those previously exposed to cyclophosphamide;¹⁰,³³,³⁴ unchanged in renal dysfunction.³³
Adverse Reactions. Emetic potential is moderate to high (>1 g). Nausea, vomiting, and alopecia are frequent and dose dependent. Dose-limiting toxicity is myelosuppression, with a WBC nadir of about 10 days; platelets also are suppressed, perhaps to a lesser extent. Transient, reversible blurred vision occurs frequently. The drug is locally nonirritating. Renally eliminated active metabolites occasionally cause sterile hemorrhagic cystitis, which can resolve slowly, often leading to a fibrotic, contracted bladder. Bladder epithelial changes range from minimal to frank neoplasia. An early sign of cystitis is microscopic hematuria, which can lead to hemorrhage. Prophylactic hydration is recommended. To prevent urotoxicity with high-dose regimens, administer mesna. (See Mesna.) Acetylcysteine (Mucomyst) bladder irrigations can have antidotal activity. Rarely, bladder dysplasia can lead to bladder cancer after very high doses or with concurrent or prior bladder radiation. Cross-allergenicity with other alkylators (eg, mechlorethamine) can occur. Ovarian and testicular function can be permanently lost after high-dose, long-term therapy. Rare reactions are a high-dose fatal cardiomyopathy, “allergic” interstitial pneumonitis, and a transient condition similar to SIADH that is preventable with vigorous isotonic hydration.

Contraindications. Previous life-threatening hypersensitivity to cyclophosphamide; marked leukopenia and thrombocytopenia; hemorrhagic cystitis; severe pulmonary toxicity caused by prior alkylator therapy.

Precautions. Pregnancy. Consider dosage reduction or discontinuation of drug in patients who develop infections.

Drug Interactions. Cyclophosphamide can prolong the action of neuromuscular blocking agents. Allopurinol and cimetidine can enhance cyclophosphamide myelotoxicity.

Parameters to Monitor. Before induction therapy, assess the patient for adequate numbers of WBCs (>3500/µL) and platelets (>120,000/µL). With long-term use, assess these counts at least monthly. Monitor closely for hematuria, especially if the patient has received a large cumulative dosage.

Notes. Do not dilute with benzyl alcohol-preserved solutions. Diluted solution is stable for 24 hr at room temperature and 6 days under refrigeration. Widely used in hematologic and solid malignancies and as an immunosuppressant in a variety of autoimmune disorders.

**Dacarbazine**

**Pharmacology.** Dacarbazine is an imidazole analogue of a purine precursor that alkylates DNA via methylidiazonium in a cell-cycle phase nonspecific fashion. It is used in malignant melanoma with about a 10–20% objective response rate.\(^{36,37}\)

**Adult Dosage.** IV as a single dose up to 850 mg/m\(^2\), repeated in 3–4 weeks. Alternatively, it may be given in a dosage of up to 250 mg/m\(^2\)/day for 5 days, repeated in 3–4 weeks. Reduce the dosage in renal and/or hepatic impairment.

**Dosage Forms.** Inj 100, 200 mg.

**Pharmacokinetics.** Dacarbazine is extensively metabolized, some microsomally mediated (50% by N-demethylation); it is 5% plasma protein bound, with 30–45%
of a dose excreted unchanged in the urine. The drug has an $\alpha$ half-life of 35 min and a $\beta$ half-life of about 5 hr; in one patient with renal and hepatic dysfunction, the terminal half-life increased to 7.2 hr.

**Adverse Reactions.** Nausea and vomiting, which occasionally are severe, occur almost invariably; these can decrease in severity with successive courses of therapy. Dose- and duration-dependent sterility, mutagenicity, and teratogenicity have been reported. Pain on injection also occurs. The dose-limiting toxicity is myelosuppression, with a leukopenic nadir at 21–25 days. Occasionally, a flu-like syndrome of myalgia, fever, and malaise occurs within 1 week of drug administration. Use dacarbazine with caution in patients with pre-existing bone marrow aplasia and avoid exposure to sunlight because of possible photosensitivity reactions. The drug is light sensitive so minimize exposure to light after reconstitution. The reconstituted solution is clear to pale yellow and is stable for 8 hr after reconstitution at room temperature; pink discoloration denotes drug decomposition.

**IFOSFAMIDE**

**Pharmacology.** Ifosfamide is a structural analogue of the alkylating agent cyclophosphamide (CTX). The rate of hepatic conversion of ifosfamide to the active metabolite 4-hydroxyifosfamide is slightly slower than with CTX, although formation of the bladder toxin acrolein is not reduced. The ultimate metabolite ifosformamide mustard cross-links DNA to impair cell division. The drug is always given with mesna to prevent urotoxicity. Although labeled for use in refractory testicular cancer, ifosfamide also has useful activity against soft tissue sarcoma, malignant lymphoma, and small cell lung cancer. Ifosfamide is cell-cycle phase nonspecific.

**Administration and Adult Dosage.** IV for refractory testicular cancer 1.2 g/m²/day over 30 min to 4 hr for 5 days, or 2 g/m²/day for 3 consecutive days. The recommended concurrent IV mesna dose is 20% of the ifosfamide dose, given 15 min before ifosfamide and again at 4 and 8 hr. It can be directly admixed with ifosfamide. The latter two mesna doses can be given orally at twice the dose (ie, each at 40% of the ifosfamide dose) if patient compliance and a lack of emesis can be assured. Alternatively, IV by continuous infusion 5–8 g/m² over 24 hr with mesna added at the same concentration as ifosfamide. However, more severe nephrotoxicity can occur with this regimen.

**Special Populations.** Pediatric Dosage. IV for sarcomas (Ewing’s and osteosarcoma) 1.2 g/m²/day over 30 min for 5 days, each with 3 IV mesna doses, as above.

Geriatric Dosage. Same as adult dosage but adjust for age-related reduction in renal function.

**Other Conditions.** Dosage reduction is indicated in patients with reduced renal function, although specific guidelines are not available.

**Dosage Forms.** Inj 1, 3 g.

**Patient Instructions.** (See Antineoplastics Class Instructions.)

**Pharmacokinetics.** Fate. Ifosfamide, but not its metabolites, penetrates into the CNS; CSF levels are about 38–49% of simultaneous serum levels. Ifosfamide is
metabolized to the active alkylating agent ifosfamide mustard by CYP2B6, which converts ifosfamide to 4-hydroxyifosfamide (which can act as the transport form of the molecule). The 4-hydroxy metabolite is then chemically or enzymatically broken down to active and inactive metabolites. Inactive metabolites include 4-carboxyifosfamide and several dechloroethylated species such as thiodiacetic acid. About 60–80% of a dose is excreted in the urine over 72 hr, including up to 50% of unchanged drug. In addition to 4-hydroxyifosfamide, the bladder irritant acrolein is excreted renally and can accumulate to high concentrations in the urinary bladder.44,45

\[ t_{1/2} = 6.9 \text{ hr}. \]

**Adverse Reactions.** Emetic potential is moderate; nausea and vomiting can be readily managed with antiemetics. Alopecia occurs in most patients treated with ifosfamide. The major dose-limiting effect of ifosfamide is urotoxicity manifested as hematuria. The frequency of microscopic hematuria with ifosfamide and the chemoprotectant mesna are 5–18% of courses;38 gross hematuria is less common (<5%). (See Mesna.) Renal tubular toxicity, manifested by elevations in BUN and Cr₅, occurs in <10% of patients. It is more frequent in patients who are poorly hydrated or have pre-existing abnormal renal function,47 those with renal cell cancer,48 those given high-dose 24-hr continuous ifosfamide infusions,40 and those receiving concomitant treatment with other nephrotoxins.47 Myelosuppression primarily involves leukopenia with a 7- to 14-day nadir. This effect is less severe than with cyclophosphamide and rarely affects platelets. However, in combination with other myelosuppressive drugs, additive leukopenia that is not reduced by mesna can occur. Leukopenia has been particularly severe in nephrectomized patients with renal cell cancer.40 CNS toxicities occur in up to 50% of patients, but risk factors, including dosage, are unclear. The most common effect is slight sedation or somnolence, which rarely proceeds to coma and death. These signs appear within 2 hr and typically remit 1–3 days after drug administration. Other rare CNS neurotoxocities are cerebellar toxicity (ataxia), urinary incontinence, and seizures.49 Some of these CNS effects might be caused by the minor metabolite chloracetaldehyde, which is excreted in the urine and accumulates in renal failure.50 Transient elevation in LFTs is frequently reported but is rarely clinically important. Other occasional toxic effects are allergic reactions, diarrhea, peripheral neuropathy, and stomatitis.

**Contraindications.** Severe pre-existing myelosuppression.

**Precautions.** Because of more severe nephrotoxicity and CNS toxicities, patients with reduced renal function, and particularly nephrectomized renal cell cancer patients, are poor candidates for this agent. Withhold repeat therapy until there is resolution of microscopic hematuria (<10 RBCs per high-power field). An adequate state of hydration is critical to reducing urotoxicity.

**Drug Interactions.** Use with cisplatin can increase nephrotoxicity and potassium and magnesium loss, especially in children. Nephrotoxicity is also enhanced when ifosfamide is combined with other nephrotoxic drugs.

**Parameters to Monitor.** Ensure that renal function and peripheral WBC counts are normal before administration. During therapy, monitor hematuria daily be-
cause dosage reduction or higher mesna dosage can prevent more serious urotoxicity.

**Notes.** Ifosfamide is compatible with D5W, NS, Ringer’s lactate injection, and sterile water. It also can be directly mixed with mesna. Exercise caution to reduce exposure during handling and disposal.

**MECHLORETHAMINE HYDROCHLORIDE**

**Pharmacology.** Mechlorethamine (nitrogen mustard; HN$_2$) is a prototype bis-chloroethylamine, polyfunctional alkylating agent. In solution, the compound readily ionizes to an active form, which can alkylate at a number of nucleophilic protein sites, principally the N$^7$ position of guanine in DNA and RNA. This action is cell-cycle phase nonspecific.

**Administration and Adult Dosage.** IV for Hodgkin’s disease (in the classical MOPP regimen) 6 mg/m$^2$ by careful push on days 1 and 8 of a monthly treatment cycle.$^{51}$ Irritation, spasm, and sclerosis occur in exposed veins; therefore, it is common to begin venipunctures low on the limb and move up serially and administer mechlorethamine last in a combination drug sequence. IV as a single agent up to 0.4 mg/kg as a single monthly dose. Top for mycosis fungoides and psoriasis 10 mg/60 mL of water, applied to the affected body areas 1 or 2 times a day.$^{52}$

**Special Populations.** Pediatric Dosage. IV same as adult dosage.

**Geriatric Dosage.** Same as adult dosage.

**Dosage Forms.** Inj 10 mg.

**Patient Instructions.** (See Antineoplastics Class Instructions.)

**Pharmacokinetics.** Fate. Chemical cyclization occurs in vivo to form positively charged carbonium ions, which rapidly react with various cellular components; unchanged drug cannot be detected in the blood within minutes of administration. Less than 0.01% of unchanged drug is recovered in the urine; however, up to 50% of radioactively labeled products can be found in urine within 24 hr.$^1$

**Adverse Reactions.** Emetic potential is high; nausea and vomiting within the first 3 hr are severe and can last more than 1 day. The major dose-limiting toxicity is myelosuppression: leukopenic nadir occurs at 6–8 days, thrombocytopenic nadir at 10–16 days. Extravasation causes delayed and protracted (months) ulceration and necrosis; a 1/6 molar sodium thiosulfate solution (4 mL of 10% sodium thiosulfate plus 6 mL sterile water) and copious flushing with water may be used as topical antidotes to lessen serious tissue damage. Primary reproductive failure and alopecia are frequent in males and females. IV or topical use can cause maculopapular rashes and sometimes severe sensitivity reactions (anaphylaxis and occasional cross-reactivity with other alkylating agents).

**Contraindications.** Prior severe hypersensitivity reactions; pre-existing profound myelosuppression; infection.

**Precautions.** Give patients with lymphomas (especially “bulky” lymphomas) prophylactic allopurinol 2–3 days before and throughout therapy to prevent hyperuricemia and urate nephropathy after massive tumor lysis. Make every effort to avoid topical contact with this highly vesicant drug by health personnel.
Drug Interactions. None known.

Parameters to Monitor. Pretreatment and at least monthly assessment of bone marrow function, particularly WBC and platelet counts.

Notes. Mechlorethamine is a powerful vesicant and should be prepared with great caution. Use mask and rubber gloves during preparation and avoid inhalation of dust and vapors or contact with skin and mucous membranes, especially the eyes. Use the injection within 1 hr of preparation; topical solution and ointment are stable for 1 month under refrigeration. Because of its extreme acute toxicity, use is limited primarily to malignant lymphomas and topically in mycosis fungoides, a cutaneous non-Hodgkin’s T-cell lymphoma.

Pharmacology. Mitomycin (mitomycin C) is an antibiotic that contains quinone, urethane, and aziridine groups. It is activated chemically and metabolically to alkylating species; it is cell-cycle phase nonspecific, but maximum efficacy is in the G1 and S phases. Mitomycin is used primarily in GI tract tumors intravenously and bladder cancer intravesically.

Adult Dosage. IV as a single agent 10–15 mg/m² in a single dose and repeated q 6 weeks if hematologic toxicity has resolved. IV in combination regimens 5–10 mg/m² repeated in 4–6 weeks. Intravesically in bladder cancer up to 60 mg/week.

Dosage Forms. Inj 5, 20, 40 mg.

Pharmacokinetics. After IV doses of 15 mg/m², the peak serum level is about 1 mg/L (3 μmol/L). The drug is eliminated primarily by hepatic clearance, with about 20% hepatic extraction and 10–30% recovery of unchanged drug in the urine. Cl is 0.3–0.4 L/hr/kg. The drug has an α half-life of 5–10 min after IV injection and a β half-life of 46 min.

Adverse Reactions. Nausea, vomiting, diarrhea, alopecia, and nephrotoxicity occur frequently. The drug also produces sterility, mutagenicity, and teratogenicity. The dose-limiting toxicities are myelosuppression (with a long leukopenic nadir of 3–4 weeks), thrombocytopenia, and anemia, all of which can be cumulative. Monitor the patient carefully for delayed and prolonged myelosuppression. Severe ulceration can occur if the drug is extravasated (topical DMSO may be useful). Interstitial pneumonia, for which a glucocorticoid is helpful, occurs occasionally. Long-term therapy occasionally causes hemolytic-uremic syndrome. Mitomycin is contraindicated in patients with pre-existing severe myelosuppression or anemia.

Nitrosoureas:

Carmustine (BCNU) and lomustine (CCNU) are highly lipid-soluble drugs that are metabolized to active alkylating and carbamoylating moieties. Several key cellular enzymatic steps are inhibited, including those involving...
DNA polymerase and RNA and protein synthesis. There is typically only partial cross-resistance to classical alkylators. The nitrosoureas are cell-cycle phase non-specific and even have activity on G0 (resting phase) cells.

**Administration and Dosage.** *(See also Notes.)*

<table>
<thead>
<tr>
<th>Administration</th>
<th><strong>CARMUSTINE</strong></th>
<th><strong>LOMUSTINE</strong></th>
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<tbody>
<tr>
<td>IV in 100–200 mL</td>
<td>D5W in glass containers only over 15–45 min.</td>
<td>PO only.</td>
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<tr>
<td>Adult Dosage</td>
<td>75–100 mg/m²/day for 1–2 days or 200 mg/m² as a single dose, or 80 mg/m²/day for 3 days. Repeat at 6- to 8-week intervals.</td>
<td>100–130 mg/m² as a single dose, repeat at 6- to 8-week intervals.</td>
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<tr>
<td>Pediatric Dosage</td>
<td>Same as adult dosage.</td>
<td>Same as adult dosage.</td>
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<tr>
<td>Geriatric Dosage</td>
<td>Reduce dosage by 25–50% and/or increase treatment interval to at least 8 weeks.</td>
<td>Same as adult dosage.</td>
</tr>
<tr>
<td>Other Conditions</td>
<td>Treat patients with heavily pretreated bone marrow with 50–75% of the recommended dosage and/or at lengthened treatment intervals (8 weeks minimum).</td>
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**Dosage Forms.** *(Carmustine) Inj 100 mg with alcohol diluent (BiCNU); Wafer 7.7 mg (Gliadel). (Lomustine) Cap 10, 40, 100 mg—commercial packet contains two of each strength for a total of 300 mg. *(See Notes.)*

**Patient Instructions.** *(See Antineoplastics Class Instructions.)* Take lomustine on an empty stomach.

**Pharmacokinetics.** *Fate.*

| **Absorption** | — | Complete after 30 min. |
| **Distribution** | Both drugs are diffusely distributed with decreasing relative concentrations in spleen, liver, and ovaries; both achieve substantial penetration into CNS with simultaneous CSF levels of >50% of serum for intact carmustine and its metabolites and >30% for intact lomustine and its metabolites; enterohepatic cycling of active metabolites is possible and may explain subsequent peaks in nitrosourea serum levels at 1 and 4 hr. | |
| **Metabolism** | Both drugs are rapidly and extensively metabolized (partly by liver microsomal enzymes) to a number of active products that have long serum half-lives compared with the parent compounds. | |
| **Excretion** | 30% urinary drug recovery as metabolites after 24 hr, 65% after 96 hr. | 50% urinary drug recovery as metabolites after 12 hr, 60% after 48 hr; <5% fecal excretion. |
| **t½** | Intact drug 5 min; biologic effect 15–30 min; metabolites, slow decay over 3–4 days. | Intact drug 15 min; cyclohexyl and carbonyl metabolites: α phase 4–5 hr, β phase 30–50 hr; chloroethyl metabolite 72 hr. |
**Adverse Reactions.** Emetic potential is moderately high to high; prophylactic antiemetics are recommended. Major dose-limiting toxicity is delayed and potentially cumulative myelosuppression; nadirs are unusually prolonged, with leukopenia at approximately 35 days and thrombocytopenia at about 30 days. Thus, doses are not repeated more often than q 6 weeks. Carmustine frequently causes severe pain at injection site and venospasm, which can be reduced by slow, dilute infusions. Both drugs can transiently elevate liver enzymes. Pulmonary fibrosis can occur after cumulative dosages >1 g/m²; nephrotoxicity consistently occurs after cumulative dosages of ≥1.5 g/m². Variant carmustine-induced pulmonary fibrosis, highly responsive to early drug discontinuation and a glucocorticoid, has been reported. Other occasional toxicities are CNS effects (e.g., confusion, lethargy, ataxia), stomatitis, and alopecia. In animal models, the nitrosoureas are highly carcinogenic and several clinical cases of leukemia after nitrosourea therapy have been reported.

**Contraindications.** Demonstrated hypersensitivity; marked pre-existing myelosuppression.

**Precautions.** Pregnancy.

**Drug Interactions.** Experimentally in rats, carmustine, lomustine, and the investigational drug semustine are cleared much more rapidly (with reduced antitumor activity) by pretreatment with phenobarbital, which stimulates microsomal enzymes. Conversely, cimetidine can impair metabolism and increase nitrosourea myelotoxicity. Clinical resistance to carmustine and perhaps other nitrosoureas is reduced by concomitant amphotericin B. Digoxin and phenytoin serum levels might be decreased after carmustine-containing combination regimens.

**Notes.** Store carmustine under refrigeration; appearance of an oily film in the vial is evidence of decomposition, and such vials should be discarded. Carmustine is incompatible with sodium bicarbonate. Lomustine absorption is rapid; thus, vomiting 45 min or more after ingestion does not require readministration.

Carmustine implant wafers (Gliadel) are indicated for implantation in the resection cavity of patients undergoing surgery for recurrent glioblastoma multiforme. In a multicenter, placebo-controlled trial, 6-month survival in patients with glioblastoma was 50% greater with carmustine implants than with placebo. The typical adult dosage is 8 wafers (61.6 mg of carmustine) implanted at the time of surgery. In experimental systems, the polymer releases carmustine over 2 to 3 weeks in vivo. Intracranial infections occur at a higher rate (4% vs 1%), and seizures are more frequent in patients with carmustine wafers than in untreated patients. Mild to moderate healing abnormalities occur in 4% of treated patients compared with 1% of untreated patients. Other systemic and CNS side effects are equivalent in treated and untreated patients.

**Pharmacology.** Procarbazine is an N-methylhydrazine derivative that undergoes auto-oxidation and microsomal activation to form several alkylating species, including the diazonium ion and several oxygen free radicals such as H₂O₂, ·OH, and ·O₂ (superoxide). It is cell-cycle phase nonspecific and used in brain tumors and Hodgkin’s and non-Hodgkin’s lymphomas.
Adult Dosage. PO 50–200 mg/m²/day for 10–25 days, repeated in 3–4 weeks. Calculate the dosage based on IBW and reduce dosage for a BUN >40 mg/dL, Cr >2 mg/dL, or serum bilirubin >3 mg/dL.

Dosage Forms. Cap 50 mg.

Pharmacokinetics. The drug is rapidly and well absorbed after oral administration; CNS levels are equal to those in serum after 0.5–1.5 hr. Procarbazine is 70% recovered in the urine, primarily as an acid metabolite, with <5% excreted unchanged.

Adverse Reactions. Frequent CNS side effects include dizziness, headache, ataxia, nightmares, depression, and hallucinations (in up to 30% of patients). Paresthesias also can occur occasionally. Mild to moderate nausea and vomiting occur in 60–90% of patients, but tolerance usually develops rapidly. Dose- and duration-dependent sterility, mutagenicity, and teratogenicity are reported. The drug predisposes patients to secondary acute nonlymphocytic leukemias. The dose-limiting toxicity is myelosuppression with a pancytopenic nadir at 2–3 weeks. Occasional side effects include a flu-like syndrome, allergic pneumonitis, and rash. Procarbazine is contraindicated in patients with severe hypersensitivity to the drug or pre-existing bone marrow aplasia. Periodic evaluations of neurologic status and monthly CBCs may be useful.

Drug Interactions. Avoid concurrent use with MAO inhibitors, alcohol, heterocyclic antidepressants, sympathomimetics, or tyramine-containing foods. Microsomal enzyme-inducing drugs might augment procarbazine cytotoxicity. Procarbazine potentiates barbiturates, narcotics, and other hepatically metabolized drugs.

STREPTOZOCIN

Pharmacology. Streptozocin is a glucose-containing nitrosourea. It has some selective cytotoxic activity in insulinomas and malignant carcinoid and is active to a lesser extent in other adenocarcinomas of the GI tract. The drug inhibits DNA synthesis via inhibition of pyrimidine biosynthesis and blockade of key enzymatic reactions in gluconeogenesis pathways. It is cell-cycle phase nonspecific.

Adult Dosage. IV as a single agent 1–1.5 g/m²/week for 6 weeks, followed by a 4-week observation period; IV in combination 0.5–1 g/m²/day for 5 days q 4–6 weeks.

Dosage Forms. Inj 1 g.

Pharmacokinetics. Streptozocin is highly lipophilic, achieving good CNS penetration. Streptozocin and metabolites have a short distribution phase (t¹⁄₂α 6 min) followed by possibly two elimination phases representing active metabolites (t¹⁄₂β 3.5 hr; t²⁄₃, 40 hr). The drug is rapidly and extensively metabolized (unchanged drug half-life is 35 min), and only 10–20% is excreted unchanged in urine.

Adverse Reactions. Frequent acute toxicities include nausea, vomiting, and phlebitis; carefully avoid extravasation. The drug is moderately myelotoxic but extremely nephrotoxic. Signs of streptozocin nephrotoxicity include various renal tubular defects and proteinuria; adequate hydration can offer some protection. It also selectively destroys pancreatic β cells.
Pharmacology. Temozolomide is a synthetic oral alkylating agent structurally related to dacarbazine. Both are converted in vivo to 3-methyl-(triazen-1-yl)imidazole-4-carboxamide (MTIC). Dacarbazine requires metabolic activation through cytochrome P450 enzymes to form this intermediate, whereas temozolomide is spontaneously converted to MTIC under physiologic conditions. Metabolites of MTIC methylate the O^6 position of guanine in DNA, with additional methylation at the N^7 position, resulting in cytotoxicity.

Adult Dosage. PO for refractory anaplastic astrocytoma 150 mg/m^2 once daily for 5 days initially. Adjust subsequent dosages according to nadir neutrophil and platelet counts (see package insert for specific guidelines). The minimum recommended dose is 100 mg/m^2/day for 5 days q 4 weeks. The recommended maintenance dosage if tolerated is 200 mg/m^2/day for 5 days q 4 weeks. Treatment can be continued until disease progression. Temozolomide has not been studied in severe renal impairment (CrCl <36 mL/min/m^2) or in severe hepatic impairment.

Dosage Forms. Cap 5, 20, 100, 250 mg.

Pharmacokinetics. Temozolomide’s oral bioavailability is 100%; food reduces the rate and extent of absorption. Peak plasma concentrations occur in 0.3–2 hr. Temozolomide is 14% bound to plasma proteins and penetrates the CNS in concentrations of about 30% of plasma levels. V_d is 17–28 L/m^2. At neutral or basic pH, temozolomide rapidly and spontaneously hydrolyzes to MTIC and temozolomide acid metabolite (AM). MTIC is further hydrolyzed to 5-aminoimidazole-4-carboxamide (AIC) and methylhydrazine, the active alkylating agent. Less than 1% of temozolomide is excreted in the feces. Five to 7% of temozolomide, 12% of AIC, 2.3% of AM, and 17% of unidentified polar compounds are excreted renally. No accumulation of temozolomide or metabolites occurs. CI is 5.6–8.5 L/hr/m^2, with a half-life of 1.7–2.3 hr.

Adverse Reactions. The dose-limiting toxicity of temozolomide, myelosuppression, is not cumulative. Thrombocytopenia and leukopenia are dose related and predictable with nadir platelet and leukocyte counts occurring around day 22 of treatment. Anemia and lymphopenia also have been reported. The most frequent adverse effects are nausea, vomiting, constipation, and fatigue. Nausea and vomiting are usually moderate and can be controlled by taking the dose on an empty stomach and using prophylactic antiemetics. Occasional toxicities include headache, diarrhea, pain, fever, anorexia, and increased transaminase levels. Rare side effects include stomatitis, alopecia, flushing, dizziness, rash, and infection. Also reported are vomiting and elevation in liver enzymes.

Contraindications. Hypersensitive to any components of temozolomide or dacarbazine.

Notes. If capsules are accidentally opened, inhalation or contact with skin or mucous membranes should be rigorously avoided. Temozolomide is equivalent to dacarbazine in melanoma and might have less CNS relapse than dacarbazine (which does not penetrate the CNS).
Thiotepa (TESPA, TSPA) is a thiophosphoramide compound that is slowly hydrolyzed to release ethylenimine moieties that alkylate DNA. It is used systemically in the treatment of breast cancer, intracavitarily for bladder or pleural disease, and intrathecally for CNS disease. It is also given in high doses with autologous bone marrow transplantation.

**Adult Dosage.** IV, IM, or SC 0.5 mg/kg monthly or 6 mg/m²/day for 4 days. Reduce the dosage by all routes in patients with pre-existing bone marrow suppression. Intracavitary 60 mg; IT 1–10 mg/m².

**Dosage Forms.** Inj 15 mg.

**Pharmacokinetics.** Thiotepa is slowly metabolized, primarily to TEPA. Total body Cl is 8.5 L/hr/m², with 15% recovered in the urine as TEPA in 24 hr. Thiotepa has an α half-life of 7.5 min and a β half-life of 109 min.

**Adverse Reactions.** Mild nausea and vomiting occur frequently. The dose-limiting toxicity is myelosuppression (of granulocytes and platelets). Myelosuppression can occur after intravesicular or intrapleural administration. Anaphylaxis occurs rarely, and mutagenicity, teratogenicity, and sterility have been reported.

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Cladribine (2CdA) is the 20-chloro analogue of deoxyadenosine. It is a purine nucleoside that is avidly phosphorylated to toxic metabolites that accumulate intracellularly. Lymphocytes, which lack inactivating deaminase activity, are selectively destroyed by inhibition of DNA synthesis and repair. Cladribine is highly active in hairy cell leukemia; other responsive tumors are malignant lymphoma and acute and chronic myelogenous leukemias. It is also promising in the treatment of chronic progressive multiple sclerosis.

**Adult Dosage.** IV for hairy cell leukemia 0.09 mg/kg/day for 7 days by continuous infusion. New dosage regimens are exploring single daily SC injections because of the prolonged intracellular retention of active metabolites.

**Dosage Forms.** Inj 1 mg/mL.

**Pharmacokinetics.** Studies with oral administration indicate a bioavailability of 48%, implying that doubling the IV dose allows oral administration in hairy cell leukemia. The drug has a Vd of 9.2 ± 5.4 L/kg and biphasic elimination with half-lives of 35 min and 6.7 hr. About 40% of a dose is excreted renally as parent drug and metabolites.

**Adverse Reactions.** Frequent adverse reactions are severe neutropenia with fever and infection (70%), anemia (37%), and thrombocytopenia (12%). A flu-like syndrome is also common. Suppression of immune system function because of helper T-lymphocyte depletion can be quite long-lived and presents a risk of systemic opportunistic infections by fungi, bacteria, and/or parasites such as Pneumocystis carinii.
Pharmacology. Cytarabine (cytosine arabinoside, ara-C) is an arabinose sugar analogue of the natural pyrimidine nucleoside deoxycytidine. Cytarabine is converted to the triphosphate derivative, ara-CTP, which interferes with one or more DNA polymerases and is incorporated into DNA strands, leading to DNA fragmentation and chain termination. Once a threshold level of ara-C–mediated DNA damage is exceeded, apoptosis occurs. Cytarabine is cell-cycle S-phase specific, with activity markedly enhanced by continuous administration over several days.

Administration and Adult Dosage. (Conventional) IV for remission induction 100–150 mg/m²/day as a continuous infusion for 5–10 days. Experimental therapy has successfully used induction doses of 2–3 g/m² q 12 hr as a 2-hr infusion for 4–12 doses in refractory AML. IV or SC for remission induction 100 mg/m² q 12 hr for 5–10 days. SC for remission maintenance 70–100 mg/m²/day for 5 days in 4 divided doses. (See Notes.) (Liposomal) Intrathecal for lymphomatous meningitis (induction and consolidation) 50 mg on weeks 1, 3, 5, 7, 9, and 13. (Maintenance) 50 mg on weeks 17, 21, 25, and 29. If neurotoxicity develops, reduce subsequent doses to 25 mg; if it persists, discontinue therapy. Administer each dose over 1–5 min directly into the CSF via an intraventricular reservoir or into the lumbar sac. Give dexamethasone 4 mg PO or IV bid for 5 days beginning on the day of each injection.

Special Populations. Pediatric Dosage. (Conventional) IV or SC same as adult dosage. (Liposomal) Safety and efficacy not established.

Geriatric Dosage. Same as adult dosage.

Dosage Forms. Inj (conventional) 100, 500 mg, 1, 2 g; (liposomal) 50 mg.

Patient Instructions. (Liposomal) Lie flat for 1 hour following administration via lumbar puncture. (See also Antineoplastics Class Instructions.)

Pharmacokinetics. Serum Levels. (Conventional) 50–100 mg/L (0.2–0.4 mmol/L) are required for cytotoxic effects.

Fate. (Conventional) Not systemically available after oral absorption. After injection, there is a large interpatient variation in serum levels attained as measured by various assay techniques. Serum levels of 100–400 mg/L (0.4–1.6 mmol/L) are produced by a 60-min continuous infusion of 300 mg/m². Serum levels up to 240 mg/L (1 mmol/L) are achieved with high-dose regimens. It is widely distributed and deactivated by cytidine deaminase, primarily in the liver. The CSF-to-serum ratio is 0.1–0.14:1 with bolus doses and up to 0.4–0.5:1 with continuous infusion. There is slow elimination from the CSF caused by low CNS deaminating activity; however, to attain therapeutic CSF concentrations after standard IV doses, intrathecal administration is required. Tear fluid concentrations are detectable after high-dose therapy. The drug is about 13% plasma protein bound. Vₐ is 3 ± 1.9 L/kg; Cl is 0.78 ± 0.24 L/hr/kg. The deamination product, uracil arabinoside (ara-U) is inactive and rapidly excreted in the urine; 24 hr after injection, 72% of the dose is recovered in the urine as ara-U, only 11 ± 8% as unchanged drug.

t₁/₂. (Conventional) α phase 1.6–12 min; β phase 2.6 ± 0.6 hr.
**Adverse Reactions.** (Conventional) Emetic potential is moderate (<250 mg) to moderately high (250 mg–1 g); prophylactic antiemetics are very effective. The principal side effect is dose-related myelosuppression with a leukopenic nadir of 3–11 days and a thrombocytopenic nadir of 12–14 days; megaloblastosis is typically noted in the recovering bone marrow and in the rare cases in which anemia develops. Ocular toxicity is frequent with high-dose therapy; typically, conjunctival injection and central punctate corneal opacities occur. Concurrent use of glucocorticoid eye drops is recommended with high-dose therapy.89 Occasionally, mild oral ulceration and a flu-like syndrome, manifested by arthralgias, fever, and sometimes rash, occur. Irreversible cerebellar toxicity (ataxia, cognitive dysfunction) is a risk after cumulative doses of 30 g/m².90 Hepatic enzyme elevation is rare, even with 3 g/m² doses; one instance of SIADH was reported with this large dose.83 Cutaneous small vessel necrotizing vasculitis has occurred rarely after high-dose cytarabine, 3–5 days after initiation of therapy.91 (Liposomal) Arachnoiditis is frequent but sometimes can be related to disease progression or infection. Abnormal gait, confusion, headache, somnolence, asthenia, constipation, nausea, vomiting, peripheral edema, neutropenia, and thrombocytopenia are frequent. Side effects are most likely during the 5 days after a dose.

**Precautions.** (Conventional) Myelosuppression is *not* a contraindication because marrow hypoplasia with complete suppression of the leukemic clone is the desired clinical endpoint; however, extensive supportive facilities must be available during therapy, including WBC and platelet transfusion capability. When IV (conventional) and intrathecal (liposomal) cytarabine are given within a few days of each other, spinal cord toxicity is more likely. Concurrent radiation might increase the rate of adverse reactions due to liposomal cytarabine.

**Drug Interactions.** Digoxin bioavailability from tablets may be decreased after cytarabine-containing combination regimens.

**Parameters to Monitor.** Routine WBC and platelet counts; RBC indices. (Liposomal) Monitor continuously for signs of neurotoxicity.

**Notes.** (Conventional) Patients can be taught sterile technique for self-administration of SC drug for leukemia remission maintenance. The use of small reconstitution volumes (1 mL/100 mg) and rotation of injection sites should be observed. Clinical activity is limited primarily to selected hematologic malignancies (eg, AML, ALL, DHL). The combination of cytarabine and interferon increases the rate of response and prolongs survival in patients with the chronic phase of chronic myelogenous leukemia compared with interferon alone.92

Conventional cytarabine is given by IT injection or intraventricular injection via an implanted Ommaya reservoir to prevent or treat malignant metastases from acute myeloid leukemia and other cancers. The usual adult dosage is 70 mg/m² (or a fixed 100 mg) per dose once or twice weekly. IT doses should not be repeated more often than q 3–5 days in adults. In children, the dose is reduced as follows: (<1 yr) reduce by one-half; (1–2 yr) reduce by one-third; (2–3 yr) reduce by one-sixth. The drug should be diluted only with nonpreserved, isotonic solutions such as NS or, preferably, Ringer’s lactate (because of its buffering capacity). In these dilutions, conventional cytarabine is physically compatible with hydrocortisone sodium succinate and methotrexate if a neutral pH is maintained. The half-life in CSF is 2–11 hr (mean 3.5 hr). Typical toxicities include headache and vomiting,
which are dose and frequency related. Patients with blocked or impaired CSF out-
flow might experience greater toxicity. With frequent, repeated administration,
seizures and paraplegia can occur.88,89

(Liposomal) Use within 4 hr of withdrawal from vial and discard any unused
drug. Do not dilute or mix with any other medications and do not use an in-line
filter.

**Pharmacology.** Fluorouridine is the deoxyribose metabolite of fluorouracil. The
drug inhibits DNA synthesis by binding to thymidylate synthetase in S phase of
cell division.

**Administration and Adult Dosage.** *Intra-arterially for colon cancer metastases
to the liver* 0.1–0.6 mg/kg/day for 1–6 weeks by continuous hepatic artery perfu-
sion.93 Hospitalize patients for at least the first course of therapy.

**Special Populations.** *Pediatric Dosage.* Safety and efficacy not established.

**Geriatric Dosage.** Same as adult dosage.

**Other Conditions.** Reduce dosage when combined with other myelosuppressive
drugs or in patients experiencing severe toxicity (usually mucositis or diarrhea)
from previous doses.

**Dosage Forms.** Inj 500 mg.

**Patient Instructions.** *(See Antineoplastics Class Instructions.)*

**Pharmacokinetics.** *Fate.* Fluorouridine has a high degree (69–92%) of hepatic ex-
traction. A large fraction is converted to the active phosphorylated metabolite
5-fluorodeoxyuridylate monophosphate (FdUMP). Ultimately, the drug is almost
completely metabolized to inactive compounds, which are eliminated by exhala-
tion (60% of a dose) or by urinary excretion (about 10–30% of a dose).94

\[ t_{1/2} < 15 \text{ min.} \]

**Adverse Reactions.** Emetic potential with intra-arterial administration is low. Di-
arrhea and stomatitis occur frequently. Stomatitis can be life-threatening, as can an
unusual dermatitis affecting the hands and feet; both toxicities are much more fre-
quent with prolonged infusions. The primary dose-limiting toxicity of fluorouridine
is myelosuppression, principally leukopenia with some thrombocytopenia. Liver
enzyme elevations occur frequently, but they rarely herald serious hepatic compli-
cations. Local complications involving the hepatic catheter are thrombosis, leak-
age, embolism, and infection. Some catheter placements also can result in gastric
ulcers or biliary sclerosis if their respective arterioles are inadvertently perfused.93

**Contraindications.** Pregnancy; poor nutrition; pre-existing myelosuppression; se-
rious infection.

**Precautions.** Biliary sclerosis can occur, requiring repositioning or removal of
the catheter.

**Drug Interactions.** None known.

**Parameters to Monitor.** Monitor WBC count before and after each treatment.
Observe for diarrhea (fluid and electrolyte status). Monitor for severe hepatic en-
zyme elevations, which might indicate biliary sclerosis.
Notes. Flouxuridine can be administered in NS or D5W and it is compatible with heparin.

FLUDARABINE PHOSPHATE Fludara
Pharmacology. Fludarabine is a fluorinated nucleotide analogue of vidarabine. It is rapidly converted to 2-fluoro-ara-A, which is then phosphorylated to 2-fluoro-ara-ATP, which inhibits DNA synthesis. Fludarabine has little cross-resistance with other agents used for chronic lymphocytic leukemia.

Adult Dosage. IV for B-cell CLL that has not responded to at least one standard alkylating agent regimen 25 mg/m²/day for 5 days given over 30 min in 100–125 mL of D5W or NS. Refrigerate the drug before reconstitution and use within 8 hr after reconstitution.

Dosage Forms. Inj 50 mg.

Pharmacokinetics. The metabolite 2-fluoro-ara-A has a Vd of 98 L/m², a Cl of 8.9 L/hr/m², and a half-life of about 10 hr. About 23% of a dose is excreted in the urine as unchanged 2-fluoro-ara-A, and clearance is proportional to Clcr.

Adverse Reactions. The most frequent adverse effects are myelosuppression (neutropenia, thrombocytopenia, and anemia), fever and chills, infection, rash, myalgia, nausea, vomiting, and diarrhea. Frequent pulmonary symptoms include pneumonia, cough, and dyspnea. Fludarabine produced severe CNS toxicity (ie, blindness, coma, and death) in 36% of patients treated with a dosage of 4 times the currently recommended dosage. Similar CNS toxicity occurs occasionally (≤0.2% of patients) with recommended dosages. Other CNS effects are weakness, visual disturbances, paresthesias, agitation, confusion, and peripheral neuropathy.

FLUOROURACIL Various
Pharmacology. Fluorouracil (5-fluorouracil, 5-FU) is a fluorinated antimetabolite of the DNA pyrimidine precursor uracil. It inhibits thymidine formation, thereby blocking DNA synthesis. Some fluorouracil might be incorporated into RNA, inhibiting subsequent protein synthesis. It is cell-cycle S-phase specific.

Administration and Adult Dosage. Rapid IV 15 mg/kg/week for 4 weeks followed by 20 mg/kg/week until severe toxicity develops. The drug is stopped until resolution is complete, then resumed at 5 mg/kg/week.95 IV “loading course” 12 mg/kg (800 mg maximum) as a single daily dose for 4 days, then 12–15 mg/kg/week is recommended by manufacturer; however, this regimen has been associated with severe, life-threatening bone marrow toxicity.96 IV continuous infusion 1–2 g/day for up to 5 days has been used by special treatment centers; continuous infusion does not consistently increase antitumor efficacy but does appear to lessen hematologic toxicity.97 IV for Dukes’ stage C colon cancer after resection in combination with levamisole 450 mg/m²/day for 5 days initially, then 450 mg/m² once a week beginning in 28 days and continued for 1 yr. (See Notes.) PO doses are associated with low bioavailability and short clinical response. Intra-arterial, intraperitoneal, and intracavitary administration also have been used, although floxuridine is preferred. Top for neoplastic keratoses apply daily for 1–2 weeks as a thin layer with gloved hand or nonmetal applicator. Skin response progresses sequentially through erythema, vesiculation, erosion,
ulceration, necrosis, and regranulation. Treatment is usually stopped once erosion is evident to allow healing to occur over the next 1–2 months. **Vag for condylomata acuminata** 1/3 applicatorful (1.5 g) of 5% cream once a week hs for 10 weeks.98

**Special Populations. Pediatric Dosage.** Generally indicated for adult malignancies, although theoretically; equivalent mg/kg doses could be used in children.

**Geriatric Dosage.** Same as adult dosage.

**Other Conditions.** Base dosage on ideal body weight in obesity or if the patient has excessive fluid retention.

**Dosage Forms.** **Inj** 50 mg/mL; **Top Crm** 1, 5%; **Top Soln** 1, 2, 5%.

**Patient Instructions.** (See Antineoplastics Class Instructions.) Avoid prolonged exposure to strong sunlight; report any severe sores in the mouth immediately.

**Pharmacokinetics. Fate.** Oral doses are erratically and incompletely absorbed, with bioavailability of 28%, and worsened by mixing with acidic fruit juices.98 The drug is 8–12% plasma protein bound. The drug diffuses into effusions and CSF (peak CSF levels of 60–80 nmol/L after a 15 mg/kg IV bolus). Vd is about 25 ± 12 L/kg; Cl is 0.96 ± 0.42 L/hr/kg.10 Extensively and rapidly metabolized, primarily in the liver, to a variety of inactive metabolites that are renally excreted. Up to 15% is renally excreted unchanged, 90% within 6 hr of administration. Fluoroacetate and citrate metabolites found in the CSF are believed to mediate rare CNS (cerebellar) toxicities.

\[ t_{1/2} \alpha \text{ phase about 8 min; } \beta \text{ phase } 11 \pm 4 \text{ min.}^{99} \]

**Adverse Reactions.** Emetic potential is moderately low (<1 g) to moderate (>1 g). Dose-limiting toxicity is myelosuppression (when given by bolus injection) with leukopenic and thrombocytopenic nadirs at 7–14 days. Severe stomatitis 5–8 days after therapy can herald severe impending myelosuppression; this occurs unpredictably with large bolus doses (>12 mg/kg). With continuous infusions, myelosuppression is reduced considerably, but mucositis and diarrhea can be dose limiting. Oral administration increases the severity of the frequent mild diarrhea. GI ulceration is occasionally severe. Cutaneous toxicities include mild to moderate alopecia, hyperpigmentation of skin and veins, and rashes that are often worsened by sunlight. Excessive lacrimation is frequent; occasionally tear duct fibrosis develops. Rare toxicities involve CNS dysfunction manifested by ataxia, confusion, visual disturbances, and headache. Cardiotoxicity occurs rarely.

**Contraindications.** Pregnancy. Pre-existing severe myelosuppression (WBCs <2000/µL, platelet count <100,000/µL); poor nutritional state; serious infections.

**Precautions.** Use with caution in patients with pre-existing coronary artery disease.

**Drug Interactions.** Concurrent allopurinol appears to block one activation pathway, thereby reducing fluorouracil hematologic toxicity. Fluorouracil can inhibit the antipurine effects of methotrexate. The clinical importance of these two interactions is unclear.
Parameters to Monitor. Pretreatment and monthly assessment of bone marrow function, particularly WBC and platelet counts. In the weeks after administration, observe for severe stomatitis, which can herald life-threatening myelosuppression.

Notes. If a precipitate is noted in the ampule, gently warm in a water bath and/or vigorously shake to redissolve. Fluorouracil is physically incompatible with diazepam, doxorubicin, cytarabine, and methotrexate injections. Mild to moderate activity in GI tract tumors and breast cancer; topical application of cream is often curative in superficial skin cancers. Leucovorin has been used with fluorouracil to increase fluorouracil binding to the target enzyme, thymidylate synthetase. Levamisole (Ergamisol) is an immunomodulator used to enhance fluorouracil efficacy in Dukes’ C colon cancer. It is given orally in a dosage of 50 mg q 8 hr for 3 days, q 14 days for 1 yr.

GEMCITABINE

Pharmacology. Gemcitabine is a difluorinated nucleoside analogue of cytarabine that is phosphorylated by intracellular deoxycytidine kinase to the active di- and triphosphate forms. These antimetabolites inhibit ribonucleotide reductase and reduce the normal pool of deoxycytidine triphosphate, respectively. This leads to an inhibition of DNA synthesis (of replication and repair). Compared with cytarabine, gemcitabine is preferentially phosphorylated and retained intracellularly. It is approved for palliative therapy in pancreatic cancer and is also active in breast cancer and non–small cell lung cancer.100–102

Administration and Adult Dosage. IV 1 g/m²/week infused over 30 min for 7 consecutive weeks, followed by 1 week rest, then once weekly for 3 weeks with 1 week rest thereafter.

Dosage Forms. Inj 200 mg, 1 g.

Patient Instructions. (See Antineoplastics Class Instructions.) Take acetaminophen before each dose to reduce flu-like symptoms.

Pharmacokinetics. Fate. A peak serum level of 14.7 mg/L (56 μmol/L) occurs after a 1 g/m² IV dose. The drug is metabolized to active di- and triphosphate forms and also deaminated to inactive difluorodeoxyuridine (dFdU) in liver and blood. Cl is 408 ± 121 L/hr/m² in men and 31% lower in women. Renal elimination of dFdU is 77% of a dose; 5% of a dose is recovered unchanged in urine.100 τ₁/₂ (Gemcitabine) 8–14 min (dose and infusion duration dependent); (dFdU) 10–14 hr.100

Adverse Reactions. Emetic potential is moderately low and well controlled by antiemetics. Thrombocytopenia is the dose-limiting toxicity; cumulative-dosage anemia is next most common. Neutropenia occurs but is rarely dose limiting. A transient, acute flu-like syndrome consisting of fever, fatigue, chills, headache, and arthralgias occurs in most patients. Fever responds to acetaminophen and usually does not recur. Erythematous pruritic maculopapular rashes on the neck and extremities are frequent but usually respond to a tropical glucocorticoid. Hepatic transaminases increase in two-thirds of patients but this is rarely serious. Diarrhea occurs rarely.
Contraindications. Severe pre-existing thrombocytopenia.

Precautions. Thrombocytopenia can lead to serious bleeding and anemia and may require transfusion therapy. Based on a similarity to cytarabine, CNS (cerebellar) toxicities might occur after high cumulative dosages, especially with impaired renal function.

Drug Interactions. None known.

Parameters to Monitor. Monitor platelet count, RBC count, and hemoglobin levels, and serum hepatic transaminase levels monthly.

Notes. Gemcitabine is clinically active in pancreatic cancer, breast cancer, and non-small cell lung cancer, although objective increases in tumor shrinkage and survival are minimal.99–102 Gemcitabine produces primarily palliative responses such as reduced pain and enhanced quality of life with minimal serious toxicity compared with other cytotoxic agent therapies.

METHOTREXATE

Pharmacology. Methotrexate is a folic acid analogue that binds to dihydrofolate reductase, blocking formation of the DNA nucleotide thymidine; purine synthesis is also inhibited. It is most active in S phase.

Administration and Adult Dosage. Single Agent Therapy. IM, IV, or PO for choriocarcinoma 15–30 mg/day for 5 days, repeated q 1–2 weeks for 3–5 courses; IM for mycosis fungoides 50 mg once weekly or 25 mg twice weekly; IM, IV, or PO for head and neck cancer 25–50 mg/m² once weekly (watch for cumulative myelosuppression with continued administration of this regimen). IT for meningeal leukemia 12 mg/m² in a preservative-free, isotonic diluent (eg, Elliott’s B solution, patient’s own CSF, or D5LR); IV high-dose therapy (1–3 g/m²) with leucovorin rescue should be used only by experts in major research centers; IM or PO for psoriasis or arthritis maintenance 5–10 mg initially, then IM, IV, or PO 10–25 mg/week, to a maximum of 50 mg/week, depending on clinical response; long-term daily administration results in increased hepatotoxicity compared with weekly oral or parenteral doses. IM in glucocorticoid-dependent asthma 7.5 mg, then 15 mg 1 week later, with subsequent weekly doses adjusted to 15–50 mg depending on 24-hr serum levels.103 PO for glucocorticoid-dependent asthma 15 mg/week has been used.104 IM for ectopic pregnancy 50 mg/m²; some investigators repeat dose in 1 week if β-hCG levels do not drop.105,106 IM for induction of abortion 50 mg/m², followed in 3–7 days by misoprostol 500–800 μg vaginally; exact timing of misoprostol dosage and oral administration of methotrexate are under investigation.107–109

Combined Modality Therapy. For acute lymphocytic leukemia various schedules are reported for remission-maintenance therapy: IM or IV 30 mg/m² twice weekly, or 7.5 mg/kg/day for 5 days, or PO 2.5 mg/kg/day for 2 weeks; repeat at monthly intervals. IM, IV, or PO for Burkitt’s lymphoma 0.625–2.5 mg/kg/day for 1–2 weeks, then off drug for 7–10 days; IM or IV for breast cancer (combined with cyclophosphamide and fluorouracil) 40 mg/m² on days 1 and 8, then repeat monthly.110

Special Populations. Pediatric Dosage. IM or IV for remission maintenance same as adult dosage for acute lymphoblastic leukemia. IT for meningeal cancer use age-adjusted dosage rather than mg/m² dose.111
Geriatric Dosage. Same as adult dosage but adjust for age-related reduction in renal function.

Other Conditions. Patients with any “third space” fluid (eg, ascites, pleural effusions) should have fluid removed before drug administration because of drug retention and slow release of drug from these compartments. Reduce dosage in renal impairment as follows:

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<tr>
<th>AGE (YR)</th>
<th>IT DOSE (MG)</th>
</tr>
</thead>
<tbody>
<tr>
<td>&gt;3</td>
<td>12</td>
</tr>
<tr>
<td>2–3</td>
<td>10</td>
</tr>
<tr>
<td>1–2</td>
<td>8</td>
</tr>
<tr>
<td>&lt;1</td>
<td>6</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>CREATININE CLEARANCE (ML/Min)</th>
<th>PERCENTAGE OF DOSAGE RECOMMENDED</th>
</tr>
</thead>
<tbody>
<tr>
<td>&gt;50</td>
<td>60–100 (0–40% reduction)</td>
</tr>
<tr>
<td>10–50</td>
<td>30–50 (50–70% reduction)</td>
</tr>
<tr>
<td>&lt;10</td>
<td>15 (85% reduction)</td>
</tr>
</tbody>
</table>

Dosage Forms. Tab 2.5 mg; Inj (as sodium) 2.5, 25 mg/mL (preserved solution); 25 mg/mL (nonpreserved solution); 20, 50 mg, 1 g (nonpreserved powder).

Patient Instructions. (See Antineoplastics Class Instructions.) Inform your physician immediately if any of the following symptoms appear: dry cough, severe diarrhea, or mouth ulcers.

Pharmacokinetics. Serum Levels. After high-dose therapy, a threshold for bone marrow and mucosal toxicity is approximately 1 μmol/L 48 hr after administration. To prevent fatal bone marrow toxicity, keep serum levels below 10 μmol/L at 24 hr, 500 nmol/L at 48 hr, and 50 nmol/L at 72 hr.

Fate. PO and IM absorption are rapid, peaking at 1–2 and 0.1–1 hr, respectively. Oral bioavailability is dose related but averages 30%. After IT administration, the drug slowly diffuses into the bloodstream. About 34% is plasma protein bound; $V_d$ is 0.55 ± 0.19 L/kg; $Cl$ is 0.126 ± 0.048 L/hr/kg. Over 90% of a dose is excreted in the urine, 90% unchanged after IV administration of high doses. Methotrexate solubility is markedly enhanced in slightly alkaline urine and reduced in acidic urine.

ADVERSE REACTIONS. Unless otherwise indicated, these reactions apply to high-dose chemotherapy of malignancies. Emetic potential is moderate. Nearly all reactions are dose and duration related. The primary toxicity is hematologic suppression, principally leukopenia, with the nadir at 7–14 days depending upon the administration schedule (more prolonged with daily administration). Thrombocytopenia and macrocytic anemia, dose-related nephrotoxicity, and ocular irritation.
occur frequently. Hepatotoxicity occurs frequently. Diarrhea and mucosal ulcerations of the mouth and tongue occasionally become severe within 1–3 weeks after administration, sometimes heralding severe myelotoxicity. Erythematous rashes have been reported. Leukoencephalopathy occurs rarely with IV or IT use. Other toxicities after IT use include nausea and vomiting, meningismus, paresthesias, and rarely convulsions. Long-term daily administration in psoriasis has led to hepatocellular damage including fibrotic liver changes and atrophy of the liver; the frequency may be lower with larger intermittent doses. Painful plaque erosion has occurred during psoriasis therapy. Pulmonary toxicity occurs rarely at any dosage and is not always reversible. A single low dose for use in medical abortion is generally well tolerated, with none of the severe reactions reported above.

**Contraindications.** Pregnancy; lactation; severe renal or hepatic dysfunction; psoriasis or rheumatoid arthritis patients with pre-existing immunodeficiency syndromes, blood dyscrasias or anemia.

**Precautions.** Renal function must be determined before administration. Alkalinate the urine before high doses to enhance methotrexate solubility. Concomitant use with radiotherapy can increase the risk of soft tissue and osteonecrosis.

**Drug Interactions.** Concomitant vinca alkaloids (vincristine or vinblastine) can impair methotrexate elimination from the CSF and enhance methotrexate toxicity. Cisplatin, NSAIDs, omeprazole, high-dose penicillins, probenecid, and sulfonamides can increase methotrexate serum levels and toxicity. Salicylate can decrease renal elimination of methotrexate and displace it from plasma protein binding sites. Alcohol can enhance hepatotoxicity of methotrexate. Asparaginase given 1 week before or 24 hr after methotrexate appears to reduce methotrexate hematologic toxicities. Cholesterol-binding resins can decrease oral methotrexate absorption. Broad-spectrum antibiotics can decrease methotrexate serum levels and efficacy after oral administration.

**Parameters to Monitor.** Monitor pretreatment and periodic hepatic, renal, and bone marrow functions (including WBCs, platelets, and RBCs). Follow high doses with 24-hr and/or 48-hr serum methotrexate levels and institution of appropriate leucovorin rescue. Observe for pulmonary symptoms, especially a dry, nonproductive cough and for diarrhea and ulcerative stomatitis.

**Notes.** Reconstitute lyophilized forms with NS, D5W, or Elliott’s B solution (for intrathecal use). Reconstituted solutions are chemically stable for 7 days at room temperature. Methotrexate is physically incompatible with fluorouracil, prednisolone sodium phosphate, and cytarabine. It is clinically useful in a variety of hematologic and solid tumors as well as nonmalignant hyperplastic conditions such as psoriasis. If overdosage occurs, the antidote is calcium leucovorin (citrovorum factor), which can be given IV or IM in methotrexate-equivalent doses up to 75 mg q 6 hr for 4 doses. A delay of >36 hr lessens the chance of rescue.

**PENTOSTATIN**

**Pharmacology.** Pentostatin is an analogue of a normal purine intermediate involved in the conversion of adenosine to inosine. It is an irreversible inhibitor of the enzyme adenosine deaminase (ADA), which is found primarily in lymphoid
cells. Pentostatin-induced inhibition of ADA leads to a build-up of deoxyadenosine and several phosphorylated derivatives that deplete cellular ATP. These metabolic products ultimately inhibit DNA synthesis in lymphatic tumor cells, including chronic lymphocytic leukemia, acute lymphoblastic leukemia, and especially hairy cell leukemia. Some data suggest that the cytotoxic effect is cell-cycle phase specific for G1 phase.116,117

**Adult Dosage.** IV for hairy cell leukemia refractory to interferon alfa 4 mg/m² every other week.

**Dosage Forms.** Inj 10 mg.

**Pharmacokinetics.** Serum levels after doses of 2–10 mg/m² average 1.5–4.7 mmol/L. Vd is 20–23 L/m²; Cl is 3.1 L/hr/kg. The terminal half-life of pentostatin averages 5–10 hr. Up to 90% of a dose is excreted in the urine in an active form, and dosage reduction is indicated in patients with reduced renal function.

**Adverse Reactions.** Renal tubular toxicity and myelosuppression are the major dose-limiting toxicities of pentostatin. Renal toxicity manifested by Cr₅ elevation is much more frequent at doses over 5 mg/m²/day. Adequate hydration and the avoidance of other nephrotoxins can reduce the frequency and severity of pentostatin-induced nephrotoxicity. Lymphocytopenia is frequent, with B- and T-lymphocytes depressed, possibly explaining the relatively frequent, severe systemic infections with organisms that include Gram-negative bacteria, *Candida albicans*, herpes zoster (varicella), and herpes simplex. Neurologic effects are frequent with pentostatin and include lethargy and fatigue; these rarely progress to coma and are more common and severe with high-dose regimens. Mild to moderate nausea and vomiting also occur frequently but are easily controlled with standard antiemetic regimens.

**Pharmacology.** Mercaptopurine (6-MP) and thioguanine (6-TG) are thiolated purines that act as antimetabolites after metabolic activation to the nucleotide forms (phosphorylated ribose sugar attachment). Subsequently, de novo purine biosynthesis is interrupted at a number of enzymatic sites, including the conversion of inosinic acid to adenine- or xanthine-based ribosides. DNA and RNA synthesis is halted in a cell-cycle S-phase-specific fashion.

**Administration and Adult Dosage.** (Mercaptopurine) PO, IV (investigational) 75–100 mg/m²/day.118 (See Drug Interactions.) (Thioguanine) PO, IV (investigational) 2–3 mg/kg/day.

**Special Populations.** Pediatric Dosage. Same as adult dosage.

**Geriatric Dosage.** Same as adult dosage.

**Other Conditions.** Purine antimetabolite toxicities are not consistently increased in patients with renal failure.119,120 (See Precautions.)
Dosage Forms. (Mercaptopurine) Tab 50 mg; Inj (investigational) 500 mg. (Thioguanine) Tab 40 mg; Inj (investigational) 75 mg.

Patient Instructions. (See Antineoplastics Class Instructions.) To maximize absorption, do not take this drug with meals. Nausea and vomiting are uncommon with usual doses.

Pharmacokinetics. Fate. (Mercaptopurine) 12 ± 7% oral bioavailability, increasing to 60% with concurrent allopurinol. The drug is approximately 20–30% plasma protein bound and freely distributed throughout the body including placental transfer; the CSF/serum ratio is 0.19–0.27. Mercaptopurine is metabolized extensively by xanthine oxidase, also methylated to active metabolite and sulfated to inactive thiouric acid. Vₐ is 0.56 ± 0.38 L/kg; Cl is 0.66 ± 0.24 L/hr/kg; 22% excreted unchanged in urine. (Thioguanine) oral bioavailability is unknown. The drug is approximately 20–30% plasma protein bound and freely distributed throughout the body, including placental transfer; the CSF/serum ratio is 0.16. Thioguanine is metabolized predominantly to inactive metabolites.

Adverse Reactions. Emetic potential is low to moderate. The dose-limiting toxicity is myelosuppression (leukopenia and thrombocytopenia). Mild to moderate mucositis occurs with large doses and low daily maintenance doses. Predominantly cholestatic liver toxicities occur frequently with long-term therapy. Marked crystalluria with hematuria has occurred with large IV mercaptopurine doses. Various rashes also have been described with these drugs. Long-term immunosuppressive therapy with any of these agents predisposes patients to carcinogenesis; CNS lymphomas and acute myeloid leukemia are the most frequent malignancies.

Contraindications. Pregnancy; pre-existing severe bone marrow depression.

Precautions. Investigational use of mercaptopurine for inflammatory bowel disease can predispose to pancreatitis.

Drug Interactions. Patients taking allopurinol must receive substantially reduced doses of oral mercaptopurine (25–33% of the normal dose) to avoid life-threatening myelosuppression caused by blocked inactivation. Thioguanine is inactivated primarily by methylation; thus, no dosage reduction is necessary with concomitant allopurinol. Enhanced bone marrow suppression can occur with the combination of trimethoprim/sulfamethoxazole and mercaptopurine.

Parameters to Monitor. WBC and platelet counts and total bilirubin at least monthly.

Notes. There is usually complete cross-resistance between mercaptopurine and thioguanine.

UFT (Investigational—Bristol-Myers Squibb) Orzel

Pharmacology. UFT is a combination containing the fluorouracil prodrug tegafur (formerly floranur) and the ribonucleoside pyrimidine uracil in a fixed molar ratio of 1:4 (tegafur:uracil). Tegafur is gradually converted to the antimetabolite fluorouracil by metabolism in the liver. This approximates a continuous infusion of fluorouracil after oral ingestion of UFT. The uracil component slows the metabolism
of fluorouracil and reduces production of the toxic metabolite, 2-fluoro-β-alanine, resulting in reduced GI and myelosuppressive toxicity with the combination.\textsuperscript{124,125}

**Administration and Adult Dosage.** PO for colorectal cancer 800–900 mg/m\textsuperscript{2} weekly or daily for 5 consecutive days, repeated q 28 days. Alternatively 360–400 mg/m\textsuperscript{2}/day for 28 consecutive days, repeated q 35–42 days. All daily dosages are given in 3 divided doses q 8 hr.\textsuperscript{124}

**Special Populations.**

**Pediatric Dosage.** Safety and efficacy not established.

**Geriatric Dosage.** Same as adult dosage.

**Dosage Forms.** Cap containing tegafur 100 mg and uracil 224 mg.

**Patient Instructions.** (See Antineoplastics Class Instructions.) Take this drug on an empty stomach with 4–8 fluid ounces of water.

**Missed Doses.** If you are taking this drug daily, take a missed dose as soon as possible if you remember within 12 hours. If it is within 2 hours of the next dose, skip the missed dose and do not double the next dose. If you miss 2 or more doses, contact your physician. If you are taking this drug weekly and miss a dose, contact your physician.

**Pharmacokinetics.**

**Serum Levels.** No correlation has been found between the peak level or AUC of any component of UFT and myelotoxicity.

**Fate.** Tegafur and uracil are rapidly absorbed, but levels of tegafur are higher than those of uracil, despite the 4-fold higher uracil dosage. With a daily dosage of 800–900 mg/m\textsuperscript{2}, peak levels of tegafur, uracil, and 5-FU are 24.6, 13.6, and 1.4 mg/L, respectively. Tumor levels of 5-FU and its nucleotide metabolites are higher than in normal tissue. Uracil is quickly metabolized and excreted via non-biliary pathways. Some tegafur metabolites are excreted in bile.\textsuperscript{124}

**Adverse Reactions.** The dose-limiting toxicity in phase I trials using daily doses was GI, including nausea, vomiting, anorexia, and diarrhea. With daily schedules, the GI effects tend to be cumulative, resulting in moderate mucositis and diarrhea. Fatigue also occurs in over one-half of patients treated using the 28-day dosage schedule. With the shorter 5-day schedules, myelosuppression, principally neutropenia, is dose-limiting.

**Notes.** UFT also can be combined with oral leucovorin in the treatment of advanced colorectal cancer.\textsuperscript{125}

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**Cytokines**

**ALDESLEUKIN**

**Pharmacology.** Aldesleukin (interleukin-2, IL-2) is a cytokine produced by activated T-lymphocytes. It binds to T-cell receptors to induce a proliferative response and differentiation into lymphokine activated killer (LAK) cells in the blood and tumor-infiltrating lymphocytes (TIL cells) in specific tumors. The pharmaceutical product is a nonglycosylated molecule produced by recombinant DNA techniques in *Escherichia coli*.\textsuperscript{126,127}

**Administration and Adult Dosage.** IV for metastatic renal cell carcinoma 600,000 IU/kg over 15 min q 8 hr for 14 doses; repeat after 9 days of rest for a
total of 28 doses. **IV infusion** 3–6 million IU/m² infused over 6 hr is commonly used.

**Special Populations.** **Pediatric Dosage.** (<18 yr) safety and efficacy not established.

**Geriatric Dosage.** Same as adult dosage.

**Other Conditions.** Intereponential pharmacokinetic differences are not known; however, withholding dose(s) is required if severe cardiovascular collapse, pulmonary or renal insufficiency, coma, psychosis, or GI toxicity occurs.

**Dosage Forms.** **Inj** 22 million IU (1.3 mg protein). (See Notes.)

**Patient Instructions.** (See Antineoplastics Class Instructions.)

**Pharmacokinetics.** **Fate.** Limited human data indicate that the drug undergoes biphasic elimination after IV administration. The kidney is believed to be the major organ of elimination, and the drug undergoes intrarenal metabolism to inactive fragments.¹²⁸

\[ t_{1/2} \alpha = 14 \pm 7.7 \text{ min}; \beta = 80 \pm 34 \text{ min}. \]

**Adverse Reactions.** Emetic potential is low. Severe cardiovascular toxicities include fluid retention (>10% of body weight) and pulmonary interstitial edema. Hypotension requiring treatment has occurred 2–4 hr after treatment with high-dose bolus or continuous infusion and low-dose SC regimens. Anemia occurs in up to 77% of high-dose bolus IV regimens. Frequently, nausea, vomiting, diarrhea, rash, pruritus, and nasal congestion occur. Abnormal laboratory findings include frequent increased Cr, oliguria, eosinophilia, and thrombocytopenia.¹²⁶ Increased serum transaminases and bilirubin occur occasionally; hepatic dysfunction occurs rarely. Myocardial ischemia also can occur and fatal MI has been reported. Capillary leak syndrome can occur and requires close monitoring of fluid balance.¹²⁶ When combined with adoptive cellular therapy (reinfused LAK cells), immediate fever and chills result; **indomethacin** 50 mg orally or **meperidine** 25–50 mg IM or IV can lessen these symptoms.

**Precautions.** Aldesleukin has produced severe cardiopulmonary toxicity and must be used cautiously in any patient with a history of cardiac insufficiency from any cause. Patients also must be in good general physical condition to tolerate the hypotension and pulmonary edema that can complicate high-dose aldesleukin therapy.

**Drug Interactions.** Glucocorticoids block some aldesleukin actions and usually are reserved for treating severe toxicity.

**Parameters to Monitor.** Monitor blood pressure, cardiac output, and fluid balance closely.

**Notes.** Some studies describe IL-2 activity in different units or by weight. Aldesleukin is labeled in IU (18 million IU = 1.1 mg protein), and doses for other IL-2 products should be converted to IU for proper dosage. Aldesleukin is active in metastatic renal cell carcinoma (MRCC) and metastatic malignant melanoma. In MRCC, response rates are 15% (with some complete remissions), lasting a median of 23 months. Response rates are higher in patients with good performance status and especially those with pulmonary metastases as the main site of disease.
Pharmacology. Alpha interferons are single-chain proteins. The alpha-2 interferons are biosynthetic; alpha-2a has a lysine at position 23, alpha-2b an arginine. Alpha-2b interferon is a multisubspecies form of natural interferons isolated from human leukocytes. Interferons bind to specific membrane receptors and are then taken up intracellularly to affect diverse cellular functions. These include cell membrane alterations (eg, enhanced antigen expression), cell-cycle blockade at the G1–S portion, enhanced antiviral enzyme synthesis (eg, 2',5'-oligo-adenylate synthetase with resultant products, which destroy double- and single-stranded viral RNA), and immunomodulatory activity (eg, increased activity of natural killer [NK] lymphocytes and phagocytic macrophages). General cellular protein synthesis also is decreased, including cytochrome P450 enzymes. Linking the interferon to polyethylene glycol allows once weekly administration.

Administration and Adult Dosage. IM or SC for hairy cell leukemia (alpha-2a or 2b) 2 million IU/m^2 daily or 3 times a week. IM or SC for AIDS-related Kaposi’s sarcoma (alpha-2b) slowly increase dose from 5 million IU/day up to 20–36 million IU/day. Intralesionally for condylomata acuminata (alpha-2b) 1 million IU/wart 3 times weekly for 3 weeks, to a maximum 5 warts a day (use only the 10 million IU vial); (alpha-n3) 250,000 IU (0.05 mL)/wart twice weekly for up to 8 weeks, to a maximum 0.5 mL/day. IM or SC for chronic hepatitis B 5 million IU/day or 10 million IU 3 times a week for 16 weeks. IM or SC for chronic hepatitis C (conventional) 3 million IU 3 times a week for 6 months or SC for chronic hepatitis C (PEGylated) 1µg/kg once weekly. IM and SC for malignant melanoma (alpha-2b) 20 million IU/m^2 IV 5 times a week for 4 weeks, then 10 million IU/m^2 SC 3 times a week for 48 weeks. IM or SC for chronic myeloid leukemia (alpha-2a) 3–6 million IU/day.

Special Populations. Pediatric Dosage. (<18 yr) not recommended. Geriatric Dosage. Same as adult dosage.

Dosage Forms. (Alpha-2a) Inj 3, 6, 10, 36 million IU/mL. (Alpha-2b) Inj (conventional) 3, 5, 10, 18, 25, 50 million IU; (PEGylated) 100, 160, 240, 300 µg/mL. (Alpha-n3) Inj 5 million IU/mL.

Patient Instructions. (Subcutaneous use) Instruct in proper method of aseptic preparation of vials and syringes, proper technique for subcutaneous administration, and proper disposal of syringes and needles. Rotate subcutaneous injection sites. Acetaminophen is recommended to reduce frequent flu-like symptoms, which usually decrease with continued therapy.

Missed Doses. Take this drug at regular intervals. If you miss a dose of this medicine, call your physician for instructions. Do not double the dose or take extra.

Pharmacokinetics. Fate. (Conventional) Alpha-2a and 2b are 100% bioavailable after IM or SC administration, with an absorption half-life of about 6 hr. IM or SC
doses of 10 million IU produce peak serum levels of 100–200 IU/mL within 4 hr; the same dose IV produces peak serum levels of 500–600 IU/mL within 15–30 min. Alfa-n3 is not detectable in serum after intraliesional administration, although a small amount is probably absorbed. Most of a dose is thought to be metabolized, with none filtered or secreted by the kidney.\textsuperscript{132,133}

$\frac{1}{2}$ (Conventional) $\alpha$ phase 0.11 hr; $\beta$ phase (IV or IM) 2 hr, (SC) 3 hr.\textsuperscript{132,133}

**Adverse Reactions.** Emetic potential is negligible. The most frequent reactions are fevers of 38–39°C, chills, arthralgias, headache, malaise, and myalgias (flu-like syndrome). These reactions are more severe with initiation of therapy and ameliorated by acetaminophen or dosage reduction. Anorexia and nausea without vomiting also are frequent. With large doses (generally >1 million IU), hematologic suppression (eg, mild thrombocytopenia, leukopenia) occurs, as does slight elevation of hepatic enzymes (AST, LDH, alkaline phosphatase), and mild hypertension, occasionally associated with tachycardia. Very high doses (>30 million IU) are associated with somnolence, dizziness, and confusion. Mild erythema and pruritus at the injection site also can occur. Interferons are not mutagenic or carcinogenic in standard animal or in vitro models. Alpha interferons can cause or aggravate life-threatening or fatal neuropsychiatric, autoimmune, ischemic and infectious disorders. These usually resolve with drug withdrawal.

**Contraindications.** Severe hypersensitivity; development of a neutralizing serum antibody (precludes the use of alternate recombinant product; switch to natural interferon alfa-n3). (See Notes.)

**Precautions.** Pregnancy. Use with caution in patients with cardiovascular disease, seizure disorder, or hepatic or renal impairment. Proper hydration during therapy may lessen hypotensive reactions. Neutralizing serum antibodies can form after prolonged interferon administration and has been associated with reduced toxicities and antitumor effects.\textsuperscript{134}

**Drug Interactions.** Interferon can worsen the neutropenia of zidovudine in Kaposi’s sarcoma. Interferon can increase theophylline serum levels. Combination with vidarabine can result in increased neurotoxicity.

**Parameters to Monitor.** Monitor periodically for clinical and laboratory signs of life-threatening adverse effects (see Adverse Reactions.)

**Notes.** A clear dose–response relationship is established for toxicity but not for antitumor effectiveness (except for Kaposi’s sarcoma). Alpha interferons have activity in reducing the symptomatology of hairy cell leukemia; hematologic response rates of 80–90% are possible in this disease. Other cancers responsive to interferon alfa are renal cell cancer (10–30% partial response rate), acute leukemias (15–30% response rate), and the nonblastic phase of CML (40–60% response rate). Although not a labeled use, interferon alfa-n3 can be used systemically and is recommended specifically for antibody-positive patients receiving recombinant products. PEGylated forms appear to be more effective against hepatitis C than conventional forms. Patients who fail interferon treatment for hepatitis C can be given interferon alfa-2b plus oral ribavirin (Rebetol). It is available in a combination package (Rebetron).
DNA Intercalating Drugs

ANTHRACYCLINES:

**DAUNORUBICIN HYDROCHLORIDE**

**DOXORUBICIN HYDROCHLORIDE**

**IDARUBICIN HYDROCHLORIDE**

Pharmacology. Daunorubin (daunomycin), doxorubicin (hydroxydaunomycin), and idarubicin (4-demethoxydaunorubicin) are tetracyclic amino sugar-linked antibiotics that are actively taken up by cells and concentrated in the nucleus; intercalation or fitting between DNA base pairs occurs, which impairs DNA synthesis. Other biochemical lesions produced include quinone moiety-generated production of oxygen and hydroxyl free radicals with lipid peroxidation of cellular membranes. The anthracyclines also interfere with the activity of the G2-specific enzyme, topoisomerase-II, which leads to the formation of cleavable complexes between enzyme and DNA, resulting in DNA double-strand breaks. These agents are primarily cell-cycle phase nonspecific, but with slightly greater activity in late S- or G2-phase cells.

Administration and Dosage.

<table>
<thead>
<tr>
<th></th>
<th>DAUNORUBICIN</th>
<th>DOXORUBICIN</th>
<th>IDARUBICIN</th>
</tr>
</thead>
<tbody>
<tr>
<td>Administration</td>
<td>IV push, infusion.</td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>These compounds are extremely toxic</strong> (potent vesicants) if inadvertently extravasated; very careful IV technique is mandatory.</td>
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<tr>
<td>Adult Dosage</td>
<td>IV 30–45 mg/m²/day for 1–3 days; generally not repeated more often than q 3 weeks.</td>
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<tr>
<td></td>
<td>IV 60–90 mg/m² for 1 dose or 20–30 mg/m²/day for 3 days; generally not repeated more often than q 3 weeks. Alternately, 20 mg/m²/week.</td>
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</tr>
<tr>
<td></td>
<td>IV 12 mg/m²/day for 3 days. ¹³⁵</td>
<td></td>
<td></td>
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<tr>
<td>Pediatric Dosage</td>
<td>Same as adult dosage.</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Cumulative Limits</td>
<td>550 mg/m²; up to 850 mg/m².</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Lifetime Dosage</td>
<td>550 mg/m².</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Desogage</td>
<td>400 mg/m² with prior chest irradiation or pre-existing heart disease. ²</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Limits ³</td>
<td>Unknown.</td>
<td></td>
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</tbody>
</table>

²Attainment of maximal cumulative dosage generally precludes continued use, despite evidence of continuing drug response; however, some patients might continue to respond without development of cardiomyopathy. ¹³⁷ Use of dexrazoxane can extend dosage limits in breast cancer. (See Dexrazoxane.) ³Low weekly doses or continuous 96-hr infusion ¹³⁸ appear to be less toxic and might allow attainment of greater cumulative dosages (>550 mg/m²). ¹³⁹ ¹⁴⁰

Special Populations. Other Conditions. Cumulative dosages of all agents must be reduced in patients with prior irradiation of the cardiac chest region, pre-existing heart disease, or prior large cyclophosphamide dosage. Doxorubicin requires no
dosage adjustment for severe renal impairment, whereas with daunorubicin 75% of the dosage is recommended in severe renal impairment. Doxorubicin dosages, however, must be substantially reduced with severe hepatic dysfunction. Idarubicin dosage reductions are indicated for bilirubin of 2.6–5 mg/dL or Cr ≥ 2 mg/dL. For severe mucositis, administration is delayed until mucositis resolves, and then dosage is reduced by 25%.

### Dosage Forms

(Daunorubicin) **Inj** 20, 50 mg. (Doxorubicin) **Inj** 10, 20, 50, 75, 100, 150, 200 mg. (Idarubicin) **Inj** 5, 10, 20 mg.

### Patient Instructions

(See Antineoplastics Class Instructions.) Immediately report any change in sensation (eg, stinging) at the injection site during infusion (this might be an early sign of infiltration). Red-colored urine does not indicate toxicity.

<table>
<thead>
<tr>
<th>SERUM BILIRUBIN (MG/DL)</th>
<th>PERCENTAGE OF DOSE RECOMMENDED</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>DOXORUBICIN</td>
</tr>
<tr>
<td>≤1.2</td>
<td>100 (no reduction)</td>
</tr>
<tr>
<td>1.2–3</td>
<td>50 (50% reduction)</td>
</tr>
<tr>
<td>&gt;3</td>
<td>25 (75% reduction)</td>
</tr>
</tbody>
</table>

### Fate

- **Absorption**: Extensively degraded to inactive aglycone in GI tract.
- **Distribution**: Both drugs enter cells rapidly and concentrate in the nuclei. Tissue concentrations are highest in lung, kidney, small intestine, and liver; trivial amounts found in the CNS. Avid tissue binding is probably responsible for prolonged terminal half-lives and Vd of 500–600 L/m2.
- **Metabolism**: Both drugs are extensively metabolized, initially to less active alcohol metabolites; further metabolized by liver microsomes to inactive aglycones and demethylated glucuronide and sulfate conjugates.
- **Excretion**: Biliary 20–30% of a dose. Primary route of excretion.

### Dosage Forms

- **Daunorubicin**: **Inj** 20, 50 mg.
- **Doxorubicin**: **Inj** 10, 20, 50, 75, 100, 150, 200 mg.
- **Idarubicin**: **Inj** 5, 10, 20 mg.
DNA INTERCALATING DRUGS

<table>
<thead>
<tr>
<th></th>
<th>DAUNORUBICIN</th>
<th>DOXORUBICIN</th>
<th>IDARUBICIN</th>
<th>IDARUBICINOL</th>
</tr>
</thead>
<tbody>
<tr>
<td>Urinary</td>
<td>14–23% as unchanged drug and metabolites (primarily daunorubicinol).</td>
<td>5–10% as metabolites over 5 days. (141)</td>
<td>8% of a dose over 24 hr.</td>
<td>8% of a dose over 24 hr.</td>
</tr>
<tr>
<td>(t_\alpha)</td>
<td>(45) min (30) min (14) min</td>
<td>(18.5) hr (3) hr (19–34) hr. (143,145)</td>
<td>(65.5) hr. (145)</td>
<td>—</td>
</tr>
<tr>
<td>(t_\beta)</td>
<td>(18.5) hr (3) hr</td>
<td>(19–34) hr. (143,145)</td>
<td>—</td>
<td>—</td>
</tr>
<tr>
<td>(t_\gamma)</td>
<td>(17) hr (32) hr. (147)</td>
<td>—</td>
<td>—</td>
<td>—</td>
</tr>
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**Adverse Reactions.** Emetic potential is moderate to moderately high with all three drugs. Stomatitis, nausea, and vomiting are dose dependent and frequent; prophylactic antiemetics are often helpful. Myelosuppression, affecting platelets and neutrophils, is the major acute dose-limiting side effect. Typical nadirs occur at 9–14 days, with recovery nearly complete within 3 weeks of administration. Hemorrhage occurs in up to 10% of induction courses with idarubicin. Excessive lacrimation is reported in about 25% of patients receiving doxorubicin. Alopecia usually occurs; during low-dose adjuvant chemotherapy administration, regional scalp hypothermia might decrease hair loss. \(146\) Severe, protracted ulceration and necrosis can occur with inadvertent perivenuous infiltration; partially effective local treatments are limb elevation, ice packing, and topical DMSO. (See Notes.) Large evolving lesions necessitate early plastic surgery consultation. Long-term anthracycline use can lead to severe and often fatal cardiomyopathy. (See Cumulative Dosage Limits, Notes.) Symptoms such as shortness of breath, edema, and fatigue are nonspecific and indicative of advanced CHF. The frequency is low (overall 2.2%) when total dosage limits are observed and can be lower when monthly doses are given over several days or by continuous 96-hr infusion. \(147\) Late cardiotoxicity is reported in children receiving total dosages of doxorubicin \(<500\) mg/m\(^2\). \(148\) During drug infusion, various nonspecific ECG changes do not imply an increased risk of cardiotoxicity. Graded endomyocardial biopsy and graded radionuclide angiography have proved most effective for assessment of the emergence of severe cardiomyopathy. Other reactions are transient erythema and phlebitis during administration and a radiation–synergy phenomenon involving heightened tissue reactions in concurrently or previously irradiated tissues, especially the esophagus (avoid by spacing weeks apart). Urine remains red for 1–2 days after administration.

**Contraindications.** Pre-existing bone marrow suppression (WBCs \(<3000/\mu\)L; platelets \(<120,000/\mu\)L); MI in previous 6 months; history of CHF. Marrow suppression is not a contraindication in relapsed leukemia patients.

**Precautions.** Careful administration technique is mandatory to avoid extravasation and tissue necrosis. Hepatocellular disease or cirrhosis can slow production of alcohol metabolites.
Drug Interactions. A number of drugs might interact with the anthracyclines: vinca alkaloids (cross-resistance), amphotericin B (increased drug uptake), and cyclosporine and streptozocin (reduced drug clearance and increased toxicity).149 Most of these drug interactions have been studied only in vitro and require clinical confirmation.

Parameters to Monitor. Obtain pretreatment and at least biweekly nadir WBC and platelet counts. Monitor general cardiac status and serial radionuclide scans of the heart in high-risk patients. Add up prior doses to estimate cardiotoxicity dosage limit.

Notes. These drugs are compatible with usual IV solutions but incompatible with heparin, sodium bicarbonate, and fluorouracil. IV push doses are best reconstituted with NS or D5W. These solutions are stable for prolonged periods and can withstand freezing and thawing.150 Doxorubicin is widely effective in numerous solid tumors, such as ovarian, thyroid, and gastric carcinomas, sarcomas, and cancer of the breast, and hematologic malignancies, such as the lymphomas and leukemias. The iron-chelating agent dexrazoxane reduces doxorubicin-induced cardiotoxicity in patients with breast cancer.151 (See Dexrazoxane.) The activity of idarubicin and daunorubicin is limited primarily to AML. Topical DMSO (1.5 mL of a 90% w/v solution q 6 hr for 2 weeks) has been effective at preventing extravasation ulceration in one trial.152

Pharmacology. Daunorubicin is encapsulated in the lipid component of this red emulsion formulation, which consists of distearoylphosphatidylcholine and cholesterol in a fixed lipid:daunorubicin ratio of 1:18.6 (in mg/mL). These liposomes are taken up into tumor and reticuloendothelial system cells, which release prolonged but low serum levels of daunorubicin over time.153 Murine studies suggest selective (enhanced) uptake of liposomal daunorubicin into tumor tissues compared with normal organs.154 Liposomal daunorubicin is used to treat AIDS-related Kaposi’s sarcoma.

Administration and Adult Dosage. IV for Kaposi’s sarcoma 40 mg/m² q 2 weeks.

Special Populations. Pediatric Dosage. Safety and efficacy not established.

Geriatric Dosage. Same as adult dosage.

Other Conditions. Based on studies with daunorubicin, reduce dose by 25% for a serum bilirubin of 1.2–3 mg/dL and 50% for a serum bilirubin or Cr, >3 mg/dL. Do not administer if absolute granulocyte count is under 750/μL.

Dosage Forms. Inj 50 mg.

Patient Instructions. (See Antineoplastics Class Instructions.)

Pharmacokinetics. Fate. Mean peak serum levels (free plus liposomal) after doses of 20, 40, 60, and 80 mg/m² are 8.2, 18.2, 36.2, and 43.6 mg/L, respectively.153 Compared with equivalent doses of the nonliposomal drug, the free drug levels are 100-fold lower and persist for up to 2.5 days after administration. In adults, $V_d$ is
2.9–4.1 L; Cl is 0.4–0.9 L/hr, about 5% of the Cl of the free drug.\(^{153}\) Thus, the AUC is increased, Cl is slowed, but peak levels are low with the liposomal formulation. \(t_{1/2}\) 2.8–5.2 hr (total of liposomal plus free drug).

**Adverse Reactions.** Emetic potential is low to moderate. The most frequent symptoms are mild to moderate fatigue, which occurs in 56% of patients, and low-grade fever in 26% of patients. An acute triad of back pain, flushing, and chest tightness can occur in up to 14% of patients, usually with initial administration. This liposomal-component reaction subsides with interruption of the infusion and typically does not recur when restarting at a slower infusion rate. Neutropenia occurs in 17% of patients; mild anemia and thrombocytopenia occur in 7% and 4% of treatment courses, respectively. Diarrhea occurs in 10% of patients; mild liver enzyme elevation occurs in 4% of patients.\(^{153}\) Cardiac toxicity appears to be less with this formulation than with aqueous daunorubicin.

**Contraindications.** Previous serious allergy to the drug or any component of the formulation. (See Anthracyclines, Daunorubicin.)

**Precautions.** Pregnancy; lactation. Do not administer if absolute granulocyte count is under 750/\(\mu\)L.

**Drug Interactions.** Not well studied with this formulation. (See Anthracyclines.)

**Parameters to Monitor.** Monitor the number of Kaposi’s sarcoma lesions for response or evidence of disease progression (≥10 new lesions or an increase of 25%). Obtain WBC count before administration. Monitor left ventricular ejection fraction at cumulative dosages of 320 and 480 mg/m\(^2\) and \(q\) 240 mg/m\(^2\) thereafter.

**Notes.** Mix only in D5W; do not filter.

**DOXORUBICIN HYDROCHLORIDE, LIPOSOMAL**

**Pharmacology.** Doxorubicin is encapsulated in the aqueous core of small (100-nm) liposomes composed of a phospholipid bilayer with an outer coating of polyethylene glycol (PEG). The small liposome size and PEG coating mask recognition by reticuloendothelial cells, thereby increasing the half-life of the liposomes in vivo. Once the liposomes accumulate in tissues, free doxorubicin is slowly released to exert its antitumor effect. Most toxicities are reduced by the liposomal formulation without compromising efficacy in solid tumors such as Kaposi’s sarcoma. (See Anthracyclines.)

**Administration and Adult Dosage.** IV for AIDS-related Kaposi’s sarcoma 20 mg/m\(^2\) \(q\) 3 weeks.

**Special Populations.** *Pediatric Dosage.* Safety and efficacy not established.

*Geriatric Dosage.* Same as adult dosage.

**Other Conditions.** (Liver dysfunction) Reduce dosage 50% for serum bilirubin 1.2–3 mg/dL; reduce dosage by 75% for bilirubin >3 mg/dL. (Stomatitis) For patients who develop stomatitis, wait 1 week, re-evaluate, and readminister at 100% for grade II severity (painful ulcers but able to eat), 75% for grade III severity (painful ulcers and unable to eat), 50% for grade IV severity (extensive, disabling stomatitis requiring nutritional support). (Hematologic toxicity) Reduce dose and/or delay administration to allow for ANC and platelet count (PC) to return to
at least 1000/µL and 50,000/µL, respectively. Then readminister at 100% of dosage if nadir ANC was 1000–1500/µL and/or PC was 50,000–150,000/µL; 75% of dosage if nadir ANC was 500/µL and/or PC was 25,000–50,000/µL; or 50% of dosage if nadir ANC was <500/µL and/or PC was <25,000/µL, respectively. (Erythrodysesthesia) For grade I erythrodysesthesia (mild swelling or erythema) present 4 weeks after the dose, administer 75% of standard dosage. For grade II erythrodysesthesia (erythema or desquamation not precluding physical activity) present 3 weeks after the dose, delay the dose for 1 week; if present 4 weeks after the dose, reduce the next dose by 50%. For grade III erythrodysesthesia (palmar–plantar [hand/foot] that is severe [diffuse blistering]) 3 weeks after drug administration, hold the next dose for 1 week; if it is still present at 4 weeks, discontinue the drug.

**Dosage Forms.** Inj 2 mg/mL.

**Patient Instructions.** (See Antineoplastics Class Instructions.)

**Pharmacokinetics.** *Fate.* Mean peak serum levels (±SE) after 10 and 20 mg/m² doses are 4.1 ± 0.2 and 8.3 ± 0.5 µg/mL, respectively. Most of this level is liposomally encapsulated drug; the assay does not differentiate. Liposomal doxorubicin has a smaller Vₚ (2.2–4.4 L/m²) than free doxorubicin; Cl is 0.034–0.108 L/hr/m². The AUC for the 10 and 20 mg/m² doses are 277 ± 33 (±SE) and 590 ± 59 (±SE) mg·L/hr. A small amount (0.8–2.6 ng/mL) of the doxorubicinol metabolite is found in serum after a dose. Cl of parent drug is 24–35 L/hr/m². Tissue concentrations of drug can be 19 times higher in Kaposi’s sarcoma lesions than in adjacent normal skin.\(^{155(10,920)}\)

**t₁/₂.** (Liposomal and free drug) α phase 5.2 ± 1.4 hr; β phase 55 ± 4.8 hr.\(^{155(10,955)}\)

**Adverse Reactions.** Similar to free doxorubicin. Myelosuppression, principally neutropenia, occurs in 49% of patients and sepsis in 5%. Opportunistic infections also occur in AIDS patients, especially those with a high tumor burden, low CD4 count, or pre-existing infection. Palmar–plantar erythrodysesthesia is cumulative. It is manifested as painful red soles and palms, which can progress to ulceration and debilitating infection if doses are not reduced and/or delayed. Doxorubicin-induced cumulative dosage cardiomyopathy and inadvertent extravasation necrosis can be lessened, but not entirely eliminated, with the liposomal formulation. Radiation recall soft tissue toxicity has been reported.

**Contraindications.** (See Anthracyclines, Doxorubicin.)

**Precautions.** Sensitization of soft tissues to radiation damage can occur. To lessen frequency of irreversible cardiomyopathy, observe the cumulative anthracycline dosage limit of 500 mg/m². Avoid extravasation and do not give IM or SC.

**Drug Interactions.** (See Anthracyclines.)

**Parameters to Monitor.** Obtain absolute neutrophil count and platelet count, serum bilirubin level, and severity of stomatitis and palmar–plantar erythrodysesthesia before administration.

**Notes.** Do not filter. Overall response rates of 40–60% are reported for patients with AIDS-related Kaposi’s sarcoma;\(^{155,156\textsuperscript{\textregistered}}\) it also might be effective in other solid tumors in HIV-negative patients.
Pharmacology. Dactinomycin (actinomycin D) is a tricyclic, peptide-containing antibiotic that acts as an intercalator of DNA, resulting in decreased mRNA transcription in a phase-nonspecific fashion. It is used in the treatment of sarcomas and choriocarcinoma.157,158

Adult Dosage. IV 2 mg/week or 500 μg/day for up to 5 days, repeated at 3- to 4-week intervals. Reduce dosage in the presence of hepatobiliary dysfunction. Reconstitute dactinomycin with preservative-free diluents. It is bound by cellulose filters, so avoid in-line filtration.

Pediatric Dosage. IV 450 μg/m²/day, to a maximum of 500 μg/day, for up to 5 days; the course is repeated in 3 weeks. (See Adult Dosage.)

Dosage Forms. Inj 0.5 mg.

Pharmacokinetics. About 30% of the drug is recovered from feces and urine after 1 week; there is no CNS penetration, and it is probably concentrated in the bile. The terminal half-life is >36 hr.

Adverse Reactions. Nausea, vomiting, mucositis, diarrhea, and reversible alopecia occur frequently. Dose- and duration-dependent hepatotoxicity and genotoxic effects have been reported. Severe ulceration occurs if the drug is extravasated. The dose-limiting toxicity is myelosuppression with a leukopenic nadir at 7–10 days. Rarely, radiation recall occurs.

Pharmacology. Mitoxantrone is a substituted salt of a planar anthracene. The drug binds to DNA by intercalation and inhibits topoisomerase-II, producing DNA strand breaks; DNA synthesis is impaired in a cell-cycle phase nonspecific fashion.159

Administration and Adult Dosage. IV for solid tumors 12 mg/m² q 4 weeks or 5 mg/m²/week for 3 weeks. IV for leukemia 10–12 mg/m²/day for 3 days. IV for multiple sclerosis 12 mg/m² q 3 months, to a usual lifetime maximum of 140 mg/m². Administer only through a freely flowing IV line.

Special Populations. Pediatric Dosage. IV for leukemia up to 8 mg/m²/week for 3 weeks or up to 18 mg/m² q 4 weeks.160

Geriatric Dosage. Same as adult dosage.

Other Conditions. Reduce doses by approximately 30–50% in patients with abnormal hepatobiliary function and/or appreciable third-space fluid accumulations.161 Reduced doses also are required in patients with poor bone marrow reserve. No dosage alteration is required with renal function impairment.

Dosage Forms. Inj 2 mg/mL.

Patient Instructions. (See Antineoplastics Class Instructions.) This drug might turn urine blue-green for 24 hr after administration because of its dark blue color. Discoloration of the whites of the eyes might occur.

Pharmacokinetics. Fate. The drug is >95% plasma protein bound and exhibits prolonged retention in tissues. Some liver metabolism to glucuronyl and glu-
tathione conjugates occurs. Urinary recovery is <8% of a dose; the majority is eliminated in the bile; fecal recovery averages 18% of a dose over 5 days. Adverse Reactions. Emetic potential is low. Myelosuppression, principally granulocytopenia (nadir at 10–14 days), occurs and is most severe in heavily pretreated or irradiated patients. Mucositis, which is dose limiting, occurs only with weekly regimens. CHF has been reported frequently, most often after prior anthracycline therapy. Cumulative cardiotoxicity limits are not well established but can approach 125 mg/m² with prior anthracyclines and 160 mg/m² without. Alopecia and extravasation necrosis are minimal. Mitoxantrone is not usually a vesicant, although it causes necrosis rarely and usually tints the tissues a blue color. Precautions. Reduce the dosage in patients with poor hepatobiliary function. Dosage reduction might be necessary in patients previously treated with marrow suppressant or cardiotoxic agents. Drug Interactions. None known. Parameters to Monitor. Obtain serum bilirubin before each dose. Assess cardiac function in patients with prior anthracycline therapy or severe pre-existing cardiovascular disease and in multiple sclerosis patients who reach a cumulative dosage of 100 mg/m². Monitor absolute granulocyte count before each dose; nadir counts 7–10 days after the dose are optimal. Pharmacology. Plicamycin (mithramycin) is a complex, polycyclic, sugar-linked antibiotic that acts by DNA binding in a cell-cycle phase nonspecific fashion; it also has a separate calcium-lowering effect. It is used in testicular cancer and to control severe hypercalcemia caused by malignancy. Adult Dosage. IV for testicular tumors 25–30 μg/kg/day to a maximum of 3 mg for up to 5 days, repeat in 4 weeks if toxicity has resolved. IV for hypercalcemia 25 μg/kg/day to a maximum of 3 mg for 3–4 days. Reduce dosage by 25–50% in moderate to severe renal impairment. Note: Dosage is in μg/kg, with no single dose over 3 mg. Dosage Forms. Inj 2.5 mg. Pharmacokinetics. The metabolic fate of the drug is unknown, but the drug penetrates well into the CNS and 40% of radioactivity from a radiolabeled dose appears in the urine. Adverse Reactions. Mild to moderate myelosuppression with a leukopenic nadir at 7–12 days, nausea, and vomiting occur frequently. Dose- and duration-dependent nephrotoxicity (increased Cr and proteinuria) and hepatotoxicity (increased LDH and AST) occur frequently. Sterility, mutagenicity, and teratogenicity have been reported. The dose-limiting toxicity is a hemorrhagic tendency characterized by decreased platelet count and responsiveness and depressed clotting factor synthesis. Rarely, stomatitis, progressive skin thickening, and hyperpigmentation occur. The drug is an irritant, but not a vesicant if extravasated.
drug is contraindicated in patients with pre-existing bleeding diatheses, hypocalcemia, or severe renal or hepatic dysfunction. Use cautiously, if at all, with other drugs affecting platelet function (eg, aspirin).

### Hormonal Drugs and Antagonists

<table>
<thead>
<tr>
<th>ANTI-ANDROGENS:</th>
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<tbody>
<tr>
<td>BICALUTAMIDE</td>
<td>Casodex</td>
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<tr>
<td>FLUTAMIDE</td>
<td>Eulexin</td>
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<tr>
<td>NILUTAMIDE</td>
<td>Nilandron</td>
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**Pharmacology.** These drugs are nonsteroidal antiandrogens that competitively inhibit binding of testosterone at androgen receptors in the testes and prostate gland, reducing androgen-stimulated cell growth. They are used with a luteinizing hormone-releasing hormone (LHRH) analogue (eg, leuprolide or goserelin). Bicalutamide has a longer half-life and 4-fold higher affinity than flutamide for the androgen receptor, which allows once-daily administration.164,165

**Administration and Adult Dosage.** PO for prostatic carcinoma together with an LHRH analogue; (Bicalutamide) 50 mg once daily; (Flutamide) 250 mg q 8 hr; (Nilutamide) 300 mg/day for 30 days, then 150 mg/day.

**Special Populations.** Geriatric Dosage. Same as adult dosage.

**Other Conditions.** If PSA levels rise with clinical disease progression, consider discontinuing the antiandrogen temporarily and continuing the LHRH antagonist to re-establish androgen receptor sensitivity. Renal or hepatic impairment does not appear to alter elimination of either drug.

**Dosage Forms.** (Bicalutamide) Tab 50 mg. (Flutamide) Cap 125 mg. (Nilutamide) Tab 50 mg.

**Patient Instructions.** Take therapy continuously without interruption. Start bicalutamide at the same time as the luteinizing hormone-releasing hormone agonist. Hot flashes and some feminizing side effects (especially breast enlargement or tenderness) can occur during therapy.

**Missed Doses.** Take a missed dose as soon as possible. If you take the drug once daily and it is time for the next dose, take it at the regular time. Do not double the dose. If you take two or more doses daily, and it is about time for the next dose, skip the missed dose. Do not double the dose. If you miss two or more doses contact your physician.

**Pharmacokinetics.** Fate. These agents are well absorbed orally and absorption is unaffected by food, but absolute bioavailability is unknown. (Bicalutamide) With an oral dose of 50 mg/day, bicalutamide attains a peak serum level of 8.9 mg/L (21 μmol/L) 31 hr after a dose at steady state. Cl of (R)-bicalutamide is 0.32 L/hr. The active (R)-enantiomer of bicalutamide is oxidized to an inactive metabolite, which, like the inactive (S)-enantiomer, is glucuronidated and cleared rapidly by elimination in the urine and feces.165 (Flutamide) Flutamide attains peak serum levels of 78 μg/L (283 nmol/L) 2–4 hr after a 250 mg dose at steady state, and its
metabolite (α-hydroxyflutamide) achieves levels of 0.720–1.68 mg/L. Flutamide and its active metabolite α-hydroxyflutamide are bound to plasma proteins. Both drugs are extensively metabolized. The majority of a flutamide dose is excreted in the urine as 2-amino-5-nitro-4-(trifluoromethyl) phenol (inactive) with little parent and active metabolite (4.2% of a dose) excreted in the bile or feces.\textsuperscript{166,167}  

$t_{1/2}$: (Bicalutamide) 5.8 days; (flutamide) 7.8 hr; (nilutamide) 41–49 hr.\textsuperscript{165–167}

**Adverse Reactions.** These agents are relatively well tolerated. When the drugs are combined with an LHRH agonist, the following side effects occur: hot flashes (50%), general pain (25%), back pain (16%), asthenia (16%), pelvic pain (12%), constipation (15%), diarrhea (10–24%, higher with flutamide, possibly because of lactose intolerance),\textsuperscript{168} nausea (11%), nocturia (10%), liver enzyme elevation (6–10%), abdominal pain (8%), and chest pain (5%). Hepatic injury and jaundice occur rarely.

**Contraindications.** None known.

**Precautions.** Discontinue these drugs if LFTs are consistently over twice the upper limits of normal without hepatic metastases.

**Drug Interactions.** Dosage adjustment of warfarin, based on INR, might be necessary when bicalutamide is administered because it can displace warfarin from protein binding sites in vitro.

**Parameters to Monitor.** Monitor PSA levels q 3 months as an index of disease response. Obtain serum transaminases q 3–4 months to rule out drug-induced hepatic injury.

**AROMATASE INHIBITORS:**

<table>
<thead>
<tr>
<th>AMINOGLUTETHIMIDE</th>
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<tr>
<td>ANASTROZOLE</td>
<td>Arimidex</td>
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<tr>
<td>EXEMESTANE</td>
<td>Aromasin</td>
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<td>LETROZOLE</td>
<td>Femara</td>
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**Pharmacology.** Aminogluthethimide, anastrozole, exemestane, and letrozole inhibit the metabolic conversion of androstenedione to estradiol, which is mediated by aromatase, primarily in peripheral adipose tissues. In postmenopausal women, this deprives hormonally sensitive breast cancers of estrogenic stimulation. Aminogluthethimide is less specific and blocks the cholesterol-based biosynthesis of all corticosteroid precursors (eg, hydrocortisone, aldosterone) in the adrenal gland and at peripheral sites.\textsuperscript{169–171} Anastrozole, exemestane, and letrozole are much more specific inhibitors of estrogen synthesis that do not affect synthesis of other steroids. Exemestane’s inhibition is irreversible and lasts for about 72 hr after a dose.

**Administration and Adult Dosage.** (Aminogluthethimide) PO 750 mg–1.5 g/day; (anastrozole) PO 1 mg/day; (exemestane) PO 25 mg/day after a meal; (letrozole) PO 2.5 mg/day. (See Notes.)

**Special Populations.** Pediatric Dosage. (Aminogluthethimide) safety and efficacy not established, but the following has been used: PO for adrenal hyperplasia
and adrenal tumors (>2.5 yr) 0.375–1.5 g/day. (Anastrozole, exemestane, letrozole) safety and efficacy not established.

**Geriatric Dosage.** Same as adult dosage.

**Other Conditions.** (Anastrozole, exemestane) No change required in hepatic or renal impairment. (Letrozole) No dosage adjustment is required with \( Cl_r \geq 10 \text{ mL/min} \).

**Dosage Forms.** (Aminoglutethimide) Tab 250 mg. (Anastrozole) Tab 1 mg. (Exemestane) Tab 25 mg. (Letrozole) Tab 2.5 mg.

**Patient Instructions.** (Aminoglutethimide) If severe stress or trauma occurs, increased hydrocortisone dosage might be needed. Marked drowsiness can occur during therapy. Skin rashes are common, especially at the start of therapy. (Exemestane) Take this drug after a meal. (Letrozole) This drug may be taken with food.

**Missed Doses.** This drug should be taken at regular intervals exactly as prescribed. If a dose is missed, it should be taken as soon as it is remembered. If it is almost time for the next dose, take only that dose and resume the regular dosage schedule. Do not double the dose.

**Pharmacokinetics.**

**Fate.** (Aminoglutethimide) A 1 g oral dose yields serum levels of 9 \( \mu \text{g/mL} \). \( Cl \) averages 5.5 L/hr in adults. About 50% is metabolized in liver to a less active N-acetyl derivative; this and other metabolites are excreted renally.\(^{1,169}\) (Anastrozole) Extensively metabolized and excreted renally (10% as parent, 60% as metabolites).\(^{170}\) (Exemestane) Absorption is increased by 40% when taken with a high-fat meal. Extensively metabolized by CYP3A4 and aldoketoreductases, with unchanged drug accounting for <10% of drug in plasma. Metabolites have less or no inhibitory activity against aromatase. Less than 1% excreted unchanged in urine. (Letrozole) Well absorbed. \( V_d \) is 1.9 L/kg. The drug is metabolized to a glucuronide metabolite, which is excreted in urine. Only 5% is excreted unchanged in urine.

\( t_{1/2} \) (Aminoglutethimide) \( \alpha \) phase 2.5 hr; \( \beta \) phase 13.3 hr. (Anastrozole) 50 hr. (Exemestane) 24 hr. (Letrozole) about 2 days.\(^{1,169–171}\)

**Adverse Reactions.** (Aminoglutethimide) Lethargy and somnolence (80%), skin rashes (50%), visual blurring, dizziness (15–30%, especially in the elderly), nausea, vomiting, and hypotension (15%), hypothyroidism, hematologic suppression (eg, agranulocytosis, pancytopenia) (<1%). (Anastrozole) Asthenia, nausea, headache, hot flashes, back pain, emesis, dizziness, rash, constipation. (Exemestane) Hot flashes, nausea, fatigue, depression, insomnia, anxiety, dyspnea, and GI disturbances occur frequently. About 4% of patients have androgenic side effects such as acne, hair loss, or hypertrichosis. (Letrozole) Musculoskeletal pain, nausea, hot flashes, headache, sweating, hair thinning, and edema are frequent.

**Contraindications.** These drugs should generally not be given to premenopausal women.

**Precautions.** (Aminoglutethimide) Supplemental hydrocortisone 50–100 mg/day and fludrocortisone 0.1 mg/day are required during therapy.
Drug Interactions. (Aminoglutethimide) Several drug interactions can occur because of the drug’s enhancement of CYP3A metabolism; the effects of dexamethasone, digoxin, medroxyprogesterone, tamoxifen, theophylline, and warfarin might be reduced. Aminoglutethimide also induces its own metabolism, which decreases blood levels and half-lives during long-term therapy. (Exemestane) Although metabolized by CYP3A4, ketoconazole does not decrease its metabolism, so CYP3A4 inhibitor interactions are unlikely. (Letrozole) Inhibits CYP2A6 and CYP2C9.

Parameters to Monitor. (Aminoglutethimide) Monitor thyroid function and blood pressure periodically during therapy.

Notes. Letrozole is approved for first-line treatment of breast cancer based on its superiority to tamoxifen. Anastrazole is also considered a first-line therapy for breast cancer.

ESTRAMUSTINE PHOSPHATE

Pharmacology. Estramustine is a conjugate of nor-nitrogen mustard linked by a carbamate bond to the 3 position of the steroidal nucleus of estradiol. Phosphorylation at position 17 adds water solubility. Estramustine originally was thought to act as a hormonally directed alkylating agent, but later studies suggest an alternate effect, impairment of mitotic spindle formation. Dephosphorylated estradiol and estrone metabolites produce typical estrogenic effects.172

Administration and Adult Dosage. PO for prostatic carcinoma 14 mg/kg/day in 3–4 divided doses.

Special Populations. Geriatric Dosage. Same as adult dosage.

Other Conditions. Diabetic and hypertensive patients might require increased doses of insulin or antihypertensives because of estrogenic effects.

Dosage Forms. Cap 140 mg.

Patient Instructions. Take this drug on an empty stomach; particularly avoid taking with milk, milk products, or calcium-containing foods or drugs.

Missed Doses. If you miss a dose, skip the missed dose and go back to your regular dosage regimen. Do not double the dose. If you miss two or more doses, contact your physician.

Pharmacokinetics. Fate. Milk and calcium salts reduce oral bioavailability by forming nonabsorbable calcium complexes. Dephosphorylated during absorption to estradiol and estrone congeners. (See Estradiol, Estrone.)

Adverse Reactions. Emetic potential is low. The major side effects are caused by estrogenic actions such as very frequent gynecomastia, cardiovascular effects (frequent edema, occasional leg cramps, or thrombophlebitis, and rare pulmonary embolism and infarction), and GI effects (frequent nausea without vomiting, diarrhea, and occasional anorexia). Laboratory abnormalities are minimal; there is no consistent hematologic suppression and only mild increases in AST or LDH in about 30% of patients.172

Contraindications. Thrombophlebitis or thromboembolic conditions (except when tumor is the cause).
Precautions. Use with caution in patients with severe underlying cardiovascular diseases. Poorly controlled CHF also can be exacerbated by estrogen-induced fluid retention. Type 1 diabetics and patients on antihypertensive medications can have increased medication requirements for these diseases.

Drug Interactions. Dairy products or calcium salts can reduce estramustine bioavailability.

Parameters to Monitor. Responses in prostate cancer are predominantly subjective, including reduced pain and less urinary retention. Objective responses can be followed with serial acid phosphatase determinations. Attention to cardiovascular or thromboembolic signs and symptoms is important.

Notes. Estramustine phosphate is principally used in the palliative treatment of advanced prostate cancer. Objective partial response rates of 20% are common. The drug can be safely combined with cytotoxic agents.\textsuperscript{172}

GONADOTROPIN-RELEASING HORMONE ANALOGUES:

<table>
<thead>
<tr>
<th>GOSERELIN ACETATE</th>
<th>Zoladex</th>
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<tbody>
<tr>
<td>LEUPROLIDE ACETATE</td>
<td>Lupron, Viadur</td>
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<tr>
<td>TRIPTORELIN PAMOATE</td>
<td>Trelstar</td>
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Pharmacology. These drugs are synthetic peptide analogues of the natural hypothalamic hormone, gonadotropin-releasing hormone (GnRH). This hormone controls the release of pituitary luteinizing hormone (LH) and follicle-stimulating hormone (FSH) to stimulate sex hormone production in the testes (testosterone) and ovaries (estradiol, others). These synthetic agents have D-amino acid and other substitutions to increase stimulatory potency. FSH and LH are initially stimulated, followed by profound inhibition of circulating sex hormones to castration levels. This retards the growth of hormonally dependent organs including the prostate, breast, endometrium, and ovaries.\textsuperscript{173,174}

Administration and Adult Dosage. SC for prostatic carcinoma (goserelin) insert 3.6 mg implant into upper abdominal wall q 28 days; (leuprolide aqueous) 1 mg/day; (leuprolide implant) insert 72 mg implant into inner aspect of upper arm. IM for prostatic carcinoma (leuprolide depot) 7.5 mg of 1-month formulation q 28–33 days or 22.5 mg of the 3-month formulation q 3 months; (triptorelin pamoate) 3.75 mg once monthly or 11.25 mg of the 3-month formulation q 3 months. SC for endometriosis (goserelin) insert 3.6 mg implant into upper abdominal wall q 28 days for 6 months; IM for endometriosis (leuprolide depot) 3.75 mg monthly for 6 months.

Special Populations. Pediatric Dosage. SC for central precocious puberty (CPP) (leuprolide aqueous) 50 μg/kg/day initially, increasing in 10 μg/kg/day increments until total down-regulation is achieved. IM for CPP initial dosage is (≤25 kg) 7.5 mg monthly; (25–37.5 kg) 11.25 mg monthly; (>37.5 kg) 15 mg monthly. Increase in 3.75 mg/month increments until total down-regulation is achieved.

Geriatric Dosage. (Prostatic carcinoma) same as adult dosage.
Dosage Forms. (Goserelin) Implant 3.6, 10.8 mg. (Leuprolide) Inj (aqueous) 5 mg/mL; Inj (depot, 1-month) 3.75, 7.5, 11.25, 15 mg; (depot, 3-month) 11.25, 22.5 mg with 1.5 mL diluent; (depot, 4-month) 30 mg with 1.5 mL diluent. (Note: Do not use a partial dose of the 3-month formulation in place of a 1-month formulation.) Implant 72 mg of leuprolide acetate equivalent to 65 mg of leuprolide. (Triptorelin) Inj (depot, 1-month) 3.75 mg; (depot, 3-month) 11.25 mg.

Patient Instructions. Instruct in proper method of aseptic preparation of vials and syringes, proper technique for subcutaneous administration, and proper disposal of syringes and needles. (Prostate cancer) Disease symptoms such as bone pain and urinary retention might become worse briefly with initiation of therapy. (Endometriosis) Do not become pregnant while on this drug; always use a barrier contraceptive. Notify your physician if regular menstruation continues. Because therapy can cause a loss of bone density, calcium supplementation is recommended. (Pediatric CPP) A slight increase in pubertal signs and symptoms might occur initially. Adherence to therapy is critical; symptoms such as menses or breast or testicular development might indicate inadequate therapy.

Pharmacokinetics. Fate. These drugs are inactive orally. The SC, IM, and IV routes provide comparable bioavailability. The metabolism of these compounds has not been described. (Goserelin) Goserelin is slowly absorbed over the first 8 days. Thereafter, absorption is steady for the remaining 28 days, with no evidence of dose-to-dose accumulation. Goserelin serum levels of about 2.5 μg/L occur on days 15–16 in males with prostate cancer. (Leuprolide) The absorption profile of leuprolide 3-month formulation is similar to the 7.5 mg 1-month formulation. Leuprolide serum levels after a 7.5 mg depot injection are 20 μg/L at 4 hr and 0.36 μg/L at 4 weeks. (Triptorelin) Triptorelin peak levels occur within 1 week and persist for 4 weeks.

Adverse Reactions. Emetic potential is low; nausea occurs in <5% of patients. Prostate cancer symptoms flare initially, causing bone pain or urinary retention. Sexual dysfunction and decreased erections are reported in about 20% of males. Hot flashes initially can occur in up to 80% of patients with endometriosis who also might experience calcium loss and estrogen-deficiency side effects (eg, decreased libido, vaginal discomfort, dizziness, general malaise, emotional lability, depression). Mild injection site reactions are rare, unless the patient is sensitive to benzyl alcohol (leuprolide aqueous only).

Contraindications. Pregnancy, because of an established teratogenic activity in animals. Do not initiate therapy for endometriosis until after negative pregnancy test.

Precautions. Monitor carefully initially in prostate cancer patients. Those with severe metastatic vertebral lesions are subject to spinal cord compression, and those with severe urinary retention might develop renal impairment.

Drug Interactions. None known.

Parameters to Monitor. (Prostate cancer) Monitor serum LH, FSH, estradiol, and testosterone; concentrations should fall to castrate levels with adequate GnRH
analogue therapy. Close initial monitoring of disease symptom severity (bone pain, urinary retention) is required. Serum PSA levels should fall and remain low in patients who respond. (Endometriosis) Monitor pain and menstrual symptoms.

Notes. In prostate cancer, these drugs are often combined with an androgen receptor antagonist (eg, bicalutamide, flutamide) to provide complete hormonal blockade.

**TAMOXIFEN CITRATE**

**Pharmacology.** Tamoxifen is a synthetic, nonsteroidal antiestrogen that binds to cytosol or nuclear estrogen receptor (ER) proteins in hormonally sensitive organs including the breast, prostate, uterus, and ovary. The tamoxifen–receptor complex binds to chromatin in the cell nucleus, thereby stopping estrogen-dependent growth-stimulatory mRNA synthesis.

**Administration and Adult Dosage.** PO for breast cancer usually 20 mg bid in premenopausal patients and 10 mg bid in postmenopausal patients. To rapidly achieve steady-state levels, an initial 2-week course of 40 mg/m² bid followed by the standard maintenance dosage has been recommended.

**Special Populations.** **Geriatric Dosage.** Same as adult dosage.

**Dosage Forms.** Tab 10, 20 mg.

**Patient Instructions.** In premenopausal patients, the chance of becoming pregnant is increased and a barrier contraceptive should be used. You should have regular gynecologic examinations after taking this drug and report any menstrual irregularities, abnormal vaginal discharge or bleeding, or pelvic pain or pressure. Lactation can occur while you are on tamoxifen.

**Missed Doses.** If you miss a dose, skip the missed dose and go back to your regular dosage regimen. Do not double the dose. If you miss two or more doses, contact your physician.

**Pharmacokinetics.** **Onset and Duration.** Therapeutic levels are attained in ≥7 days with 10–20 mg/m²/day but 3 hr after the loading dose regimen of ≥40 mg/m² bid.

**Serum Levels.** There does not appear to be a direct relationship between serum levels and response or time to response, but all responders have tamoxifen levels >180 μg/L (0.48 μmol/L) at the time of remission.

**Fate.** Well absorbed orally, with a peak of 42 μg/L (0.11 μmol/L; 12 μg/L is N-desmethyl metabolite) achieved 3–4 hr after a 20 mg dose. Initially, the N-desmethyl concentration is only 50% of the tamoxifen level, but after 21 days the metabolite level is higher because of its longer half-life. With low-dose continuous therapy, mean steady-state tamoxifen levels of ≥260 μg/L (0.7 μmol/L) are achieved after 16 weeks. Tamoxifen is slowly but extensively metabolized, mainly to N-desmethyltamoxifen, which is equally antiestrogenic to tamoxifen. Neither is readily conjugated, and both undergo hepatic hydroxylation and conjugation followed by elimination into the bile and feces; levels are measurable for up to 6 weeks after drug discontinuation.


**Adverse Reactions.** Emetic potential is moderately low. Well tolerated, producing rare minor myelosuppression (usually in heavily pretreated patients). Menopausal symptomatology, including hot flashes, nausea, and rarely vomiting, is produced in one-third of patients. Menstrual difficulties include irregularity, vaginal bleeding, and pruritus vulvae. A serious disease “flare” occurs occasionally during initial therapy, involving hypercalcemia and an increase in bone or soft tissue pain; the flare often subsides even with continued therapy and might indicate early tumor response. Retinopathy has occurred, most commonly after very large dosages but also with usual dosages. The drug appears to produce estrogen-like effects in the bone; thus, skeletal demineralization is not a problem with long-term therapy. An increased risk of secondary uterine cancer has been reported.

**Precautions.** Pregnancy. Use with caution in patients with pre-existing leukopenia and thrombocytopenia.

**Drug Interactions.** Aminoglutethimide can decrease tamoxifen serum levels. Tamoxifen can attenuate the cytotoxic activities of fluorouracil and doxorubicin.1

**Notes.** The response rate in breast cancer is about 50–70% in ER-positive patients, whereas the rate in ER-negative patients is only about 5–10%.179 Tamoxifen has been used in endometrial, stage D prostatic, and renal cell cancers and melanoma.1 It has been used investigational to decrease the size and pain of gynecomastia.

**Pharmacology.** Toremifene is a chloro derivative of tamoxifen that binds to high-affinity estrogen receptors in hormonally dependent tissues. Like tamoxifen, it has antiestrogenic and estrogenic activities in different tissues. Effects on serum lipids are similar to those of tamoxifen (reduced total and LDL cholesterol), but toremifene slightly increases HDL levels. In ER-positive breast cancer, toremifene is comparable to tamoxifen but has minimal activity in tamoxifen-refractory patients. Unlike tamoxifen, toremifene does not produce DNA/genotoxic effects or hepatocellular carcinoma in animals, suggesting an improved long-term safety profile.166,180,181

**Administration and Adult Dosage.** PO for breast cancer 50 mg/day.

**Special Populations.** Geriatric Dosage. Same as adult dosage.

**Dosage Forms.** Tab 60 mg.

**Patient Instructions.** (See Tamoxifen.)

**Pharmacokinetics.** Fate. Peak serum levels occur 1.5–4.5 hr after a single dose, but the time to reach steady-state with long-term oral administration is 1–5 weeks. Steady-state levels with a dosage of 60 mg/day average 900 μg/L. The drug is metabolized extensively in the liver, primarily by CYP3A4. Major metabolites are N-desmethyl- and 4-hydroxytoremifene, both of which are active antiestrogens. However, only the N-desmethyl derivative is detectable in plasma with a dosage of 60 mg/day.166,180
t_{1/2}. (Toremifene) 5 days; (N-desmethyltoremifene) 6 days.\(^{180}\)

**Adverse Reactions.** Toremifene is generally well tolerated; hot flashes (in 34\%) are the most frequent side effect. Vaginal discharge or bleeding occurs in 13\% of patients, dizziness in 9\%, and edema in 5\%. It causes minimal GI toxicity, consisting of nausea in 14\% and vomiting in 4\% of patients. Sweating, vaginal discharge or bleeding, dizziness, and edema also occur frequently. Acute tumor flare occurs in 16\%, marked by transient increases in bone or musculoskeletal pain, cutaneous erythema, and/or hypercalcemia within 2 weeks of starting therapy. Worsening cataracts occur in 10\%, which is similar to tamoxifen, and mild corneal keratopathies occur in 4\%. All of these ocular effects are reversible with discontinuation. In a comparative trial, mild liver function abnormalities occurred in the toremifene group, although most were related to progressive metastatic breast cancer.

**Drug Interactions.** Drugs that induce CYP3A4 can decrease toremifene levels and those that inhibit CYP3A4 can increase levels. Toremifene can increase PT in patients taking warfarin.

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**Mitotic Inhibitors**

**DOCETAXEL**

**Pharmacology.** Docetaxel is a semisynthetic derivative of a taxane extracted from the needles of the yew tree, *Taxus baccata*. It binds to microtubule tubulin sites distinct from paclitaxel, with the similar result of enhanced microtubule polymerization that causes clumps to form and halts cell division in metaphase. It is active in refractory breast cancer and non–small cell lung cancer.\(^{182}\)

**Administration and Adult Dosage.** IV for breast cancer 60–100 mg/m\(^2\) infused over 1 hr q 3 weeks. IV for non–small cell lung cancer 75 mg/m\(^2\) over 1 hr q 3 weeks. Premedicate all patients with dexamethasone 16 mg/day for 5 days, starting 1 day before administering docetaxel.

**Special Populations.** *Pediatric Dosage.* (<16 yr) Safety and efficacy not established.

*Geriatric Dosage.* Same as adult dosage.

*Other Conditions.* Reduce dosage by 25–50% in patients with elevated hepatic enzymes (and probably elevated serum bilirubin).

**Dosage Forms.** Inj 40 mg/mL.

**Patient Instructions.** Immediately report fever or chills occurring 1–2 weeks after drug administration. This drug can cause swelling of the extremities and tingling sensations.

**Pharmacokinetics.** *Fate.* Peak serum levels average 3.6 \(\mu g/mL\) after a 1-hr IV infusion of 100 mg/m\(^2\). Over 90\% is plasma protein bound. CI averages 40 L/hr in adults; CI is reduced by \(\geq 25\%\) in patients with elevated LFTs (transaminases \(>1.5\) times normal; alkaline phosphatase \(>2.5\) times normal). Most of the drug is metabolized to less active hydroxylated forms and excreted by biliary secretion into the feces; <5\% is excreted in urine.
tₜₘ. α phase 5 min; β phase 38 min; γ phase 12 hr.

**Adverse Reactions.** Emetic potential is moderate. The dose-limiting toxicity is neutropenia, which is more severe with reduced liver function; the onset of febrile neutropenia can be as soon as 5 days after drug administration. Thrombocytopenia also occurs but is less severe. Anemia and alopecia also occur but are not dose limiting. Infusion-associated hypersensitivity symptoms (eg, facial flushing) occur in 50% of patients, whereas dyspnea, chest tightness, and low back pain are rare. A pruritic rash on the forearms, hands, and neck occurs in about 40% of patients. Mucositis, nausea, and vomiting occur in about one-third of patients. Peripheral nerve numbness and paresthesia, fluid retention, and edema are cumulative dose-related toxicities. Weight gain initially involves peripheral edema at cumulative dosages >500 mg/m²; edema can become prominent after 6 cycles (600 mg/m² total dosage) and proceed to pulmonary edema. Pretreatment with oral dexamethasone (8 mg bid for 5 days, starting 24 hr before docetaxel) retards the development of serious fluid retention.

**Contraindications.** Severe hypersensitivity to drugs formulated with polysorbate 80; neutropenia <1500/μL; hepatic transaminase levels >1.5 times the upper limit of normal; hepatic alkaline phosphatase levels >2.5 times the upper limit of normal; severe pre-existing neutropenia, edema, or peripheral neuropathy.

**Precautions.** Febrile neutropenia is frequent, necessitating careful follow-up of infectious signs after administration.

**Drug Interactions.** In vitro, metabolism of docetaxel to its hydroxy metabolites is reduced by inhibitors of CYP3A such as cimetidine, erythromycin, ketoconazole, and troleandomycin. Barbiturates stimulate metabolism of docetaxel. The clinical importance of these findings is not known.

**Parameters to Monitor.** WBC count, peripheral edema, LFTs (ALT, AST, alkaline phosphatase) and signs of infection.

**ETOPOSIDE**

**ETOPOSIDE PHOSPHATE**

**Pharmacology.** Etoposide (VP-16) is a substituted epipodophyllotoxin derivative from the May apple plant. The major cytotoxic activity is cell-cycle phase specific for G₂ and involves the induction of protein-linked DNA strand breaks by inhibiting DNA topoisomerase-II enzymes.

**Administration and Adult Dosage.** IV 200–250 mg/m² q 7 weeks, or 70 mg/m²/day for 5 days. IV of etoposide should be administered over 30–60 min or longer; etoposide phosphate may be administered over 5–210 min. IV continuous infusion 125 mg/m²/day for 5 days. PO for small cell lung cancer 2 times the IV dose, rounded to the nearest 50 mg; alternatively, 50 mg/day for 30 days.

**Special Populations.** **Pediatric Dosage.** Safety and efficacy are not established. However, etoposide has been used in dosages similar to adult body surface area dosages.

**Geriatric Dosage.** Same as adult dosage but adjust for age-related reduction in renal function.
**Other Conditions.** With Cl_r ≤ 20 mL/min, give 75% of standard dose; reduced dosage also is required with severe bone marrow compromise. Dosage reduction might be necessary with altered hepatobiliary function.

**Dosage Forms.** **Inj** (Etoposide) 20 mg/mL; (etoposide phosphate) 100 mg; **Cap** (etoposide) 50 mg.

**Patient Instructions.** (See Antineoplastics Class Instructions.)

**Pharmacokinetics.** **Fate.** Oral bioavailability is 52 ± 17% with inter- and intrapatient variabilities. CSF levels are <10% of serum levels. V_d is 0.36 ± 0.13 L/kg;**10** Cl is 1.1–1.7 L/hr/m^2^ or 0.04 ± 0.014 L/hr/kg.**184,188** Inactive metabolites include the hydroxyacid and cis-lactones. Up to 16% of a dose can be eliminated in bile; 30 ± 5% is eliminated in urine, about 70% of this is unchanged drug. t_1/2. 8.1 ± 4.3 hr, increased in uremia.**184,188**

**Adverse Reactions.** Emetic potential is low. Myelosuppression occurs, with a nadir at 7–10 days (longer with daily regimens), affecting principally the granulocytes but also platelets, with a nadir at 9–16 days. Myelosuppression might be less frequent with the phosphate form. Mild mucositis and alopecia can occur. Diarrhea is more frequent with oral administration. Hypotension occurs rarely with rapid IV bolus injections. There is one report of radiation recall skin injury in 13 of 23 patients with small cell lung cancer. Long-term administration can result in development of acute leukemia.

**Precautions.** Pregnancy; decrease dosage in severe renal dysfunction; avoid rapid IV bolus injection.

**Drug Interactions.** Anaphylaxis and possible synergistic neuropathy with vincristine and/or cardiomyopathy with anthracyclines have been reported. Cyclosporine can increase serum etoposide levels and toxicity. Phenytoin, phenobarbital, and possibly other CYP3A inducers can decrease etoposide serum levels.

**Parameters to Monitor.** Obtain peripheral granulocyte counts immediately before administration on repetitive courses. Nadir counts (1–2 weeks after dose) are optional.

**Notes.** Etoposide is indicated in the combination treatment of small cell carcinoma of the lung and refractory nonseminomatous testicular cancer. The drug is also active in lymphomas and acute leukemias (lymphoblastic and myeloblastic varieties). Etoposide is not a vesicant. Store capsules under refrigeration. Etoposide and cisplatin are compatible for 24 hr in the same container. Concentrated etoposide solutions (>1 mg/mL) can cause cracking of ABS plastic infusion system components and have short stability times of 2 hr. More dilute solutions in NS or D5W of 0.4–0.6 mg/mL have longer stability times of 8 hr and 48 hr, respectively.

**IRINOTECAN**

**Camptosar**

**Pharmacology.** Irinotecan is a water-soluble camptothecin derivative. It is a prodrug for the despiperidine metabolite SN-38, an inhibitor of topoisomerase-I enzymes. This causes single strand breaks in DNA. Irinotecan is approved for first-line treatment of colorectal cancer with 5-FU and leucovorin. The overall
response rate in advanced fluorouracil-refractory colon cancer is about 15%, with a 5.2 month median duration of response.189

**Adult Dosage.** IV as a single agent or in combination with fluorouracil and leucovorin 125 mg/m² once weekly for 4 consecutive weeks administered in 500 mL of D5W over 90 min. Subsequent doses are increased by 25–50 mg/m² if no toxicity occurs; if severe toxicity occurs, dosage is decreased by 25–50 mg/m². IV as a single agent alternatively, 350 mg/m² q 3 weeks, with subsequent doses adjusted in 50 mg/m² increments.

**Dosage Forms.** Inj 40, 100 mg.

**Pharmacokinetics.** The half-lives of irinotecan and the active SN-38 metabolite are 5.7 and 9.8 hr, respectively, with peak SN-38 levels (2–5% of irinotecan) achieved 1 hr after administration. Renal excretion accounts for <10% of a dose as irinotecan and <1% as SN-38; hepatic elimination predominates.

**Adverse Reactions.** Diarrhea occurs in 90% of patients and can be severe, requiring aggressive prophylaxis with fluids and multiple doses of loperamide. Early diarrhea can be prevented or blunted with doses of 0.25-1 mg of atropine IV or SC. Leukopenia occurs in one-third of patients, although moderate to severe myelosuppression occurs in only 15% and 11% of patients, respectively. Other serious adverse reactions include anaphylactoid reactions, orthostatic hypotension, and rarely renal impairment.

**PACLITAXEL**

**Pharmacology.** Paclitaxel is a naturally occurring diterpene taxane obtained from the bark of the Pacific yew tree, Taxus brevifolia. It binds to tubulin proteins, causing abnormal microtubule polymerization and cell-cycle arrest in metaphase.190,191

**Administration and Adult Dosage.** IV 135–175 mg/m² over 3 or 24 hr q 3 weeks. Doses up to 250 mg/m² have been used with hematopoietic colony stimulation factors. Use non-PVC infusion systems.

**Special Populations.** Pediatric Dosage. Safety and efficacy not established.

**Geriatric Dosage.** Same as adult dosage.

**Dosage Forms.** Inj 6 mg/mL.

**Patient Instructions.** (See Antineoplastics Class Instructions.)

**Pharmacokinetics.** Fate. Erratic oral bioavailability precludes oral administration. CI is 18 L/hr/m². It is metabolized to a much less active hydroxylated species by CYP3A. Eliminated 30–40% by hepatobiliary excretion; only 1–5% excreted in urine.192

\[ t_{1/2} \alpha \ 	ext{phase} \ 0.2 \ 	ext{hr} \ ; \ \beta \ 	ext{phase} \ 1.9 \ 	ext{hr} \ (\text{range} \ 0.5–2.8) \ ; \ \gamma \ 	ext{phase} \ 20.7 \ 	ext{hr} \ (\text{range} \ 4–65).192 \]

**Adverse Reactions.** Emetic potential is low. The usual dose-limiting toxicity is neutropenia, with an 8- to 11-day nadir; more severe with prolonged infusion. Dose-limiting toxicity in combination regimens with doxorubicin include neutropenia, inflammation of the cecum (typhlitis), and, with cisplatin, neuropathy.
Peripheral neuropathy (eg, numbness, paresthesias) is cumulative, dose related, and more severe with prior vinca alkaloid or concurrent cisplatin therapy. Alopecia can involve all body hair, with an abrupt onset of 2 weeks. Mucositis is dose dependent. Cardiotoxicity, primarily bradycardia, occurs in 10–30% of patients but rarely requires treatment. Myalgia and arthralgia are common but usually transient. Hypersensitivity reactions, thought to be caused by Cremophor, can occur within the first few minutes of infusion; symptoms include chest pain, hypotension, bronchospasm, urticaria, and flushing and can rapidly progress to anaphylaxis. (See Precautions.)

**Contraindications.** Hypersensitivity to Cremophor vehicle; neutropenia (<1500/μL).

**Precautions.** Recommended premedications are dexamethasone PO 20 mg at 12 and 6 hr before paclitaxel, and diphenhydramine IV 50 mg, plus cimetidine 300 mg, ranitidine 50 mg, or famotidine 20 mg 30 min before paclitaxel. Ensure that emergency resuscitation equipment is available at the start of infusion. Use cautiously in patients with heart rhythm disturbances.

**Drug Interactions.** Ketoconazole can decrease paclitaxel clearance and enhance toxicity. Although their effect is not well studied, use other CYP3A inhibitors with caution. (See also Adverse Reactions.)

**Parameters to Monitor.** Neutrophil count before administration.

**Notes.** Highly effective as a first-line or refractory treatment for ovarian cancer; typically used in platinum-containing regimens; also active in breast cancer, non–small cell lung cancer, lymphoma, and malignant melanoma.191

**TENIPOSIDE**

**Pharmacology.** Teniposide is a semisynthetic podophyllum derivative that has cell-cycle S- and G2-phase–specific cytotoxic activities similar to those of etoposide. Teniposide is active in acute leukemias and in children with relapsed acute leukemia or neuroblastoma.188,193,194

**Pediatric Dosage.** IV for acute leukemias 165–200 mg/m²/week or 165 mg/m² twice weekly.

**Dosage Forms.** Inj 10 mg/mL.

**Pharmacokinetics.** Teniposide is >90% plasma protein bound and eliminated much more slowly than etoposide. Teniposide half-lives are α phase 45 min, β phase 4 hr, and γ phase 11–30 hr (average 20); 40% of a dose is eliminated in the feces; CSF drug levels are high (27% of serum levels).

**Adverse Reactions.** The dose-limiting side effect of teniposide is myelosuppression, with the leukopenic nadir at 10–14 days. Emetic potential is low; nausea and vomiting are typically mild (more severe after oral etoposide). Hypotension is reported with rapid drug infusions. Rarely severe hypersensitivity reactions (including anaphylaxis), alopecia, and chemical phlebitis during infusion occur.
Pharmacology. Topotecan is a topoisomerase-I inhibitor that causes single-strand breaks in DNA. It is a semisynthetic derivative of camptothecin, which is derived from the bark of the Chinese tree, *Camptotheca acuminata*. Topotecan is approved for metastatic ovarian carcinoma and small cell lung cancer after failure of a primary agent and being studied in colon and breast cancers.

**Adult Dosage.** IV 1.5 mg/m²/day administered over 30 min for 5 days, starting on day 1 of a 21-day course of therapy for at least 4 courses. With a Clcr of 20–39 mL/min, the dosage is reduced to 0.75 mg/m²/day; no guidelines exist for Clcr <20 mL/min. If severe neutropenia occurs, reduce further doses by 0.25 mg/m²/day or administer filgrastim with subsequent courses.

**Dosage Forms.** Inj 4 mg.

**Pharmacokinetics.** Topotecan is rapidly hydrolyzed in plasma. About 70% of the drug is excreted renally as metabolites.

**Adverse Reactions.** The primary dose-limiting side effect of topotecan is neutropenia, with a nadir at a mean of 11 days; severe neutropenia occurs in 80% of patients. Severe anemia in 40% of patients and severe thrombocytopenia in 26% also have been reported. Nausea and vomiting occur in most patients; other GI effects are frequent diarrhea, constipation, and abdominal pain. Alopecia occurs in about 60% of patients; fatigue and fever of ≥101°F also frequent. Topotecan is contraindicated in pregnancy, breastfeeding, or severe bone marrow depression.

**VINCA ALKALOIDS:**

**VINBLASTINE SULFATE**

Velban, Various

**VINCRISTINE SULFATE**

Oncovin, Various

**VINORELBINE TARTRATE**

Navelbine

**Pharmacology.** The vinca alkaloids are *Vinca rosea* (periwinkle) plant-derived antimitotic agents; cytotoxic activity is related to specific binding to the microtubule protein tubulin, causing microtubule dissolution. This blocks formation of the mitotic spindle apparatus necessary for cell division. The vincas are lethal to cells at high concentrations; at lower concentrations, dividing cells are arrested in the metaphase portion of mitosis.

**Administration and Dosage.**

<table>
<thead>
<tr>
<th><strong>VINBLASTINE</strong></th>
<th><strong>VINCRISTINE</strong></th>
<th><strong>VINORELBINE</strong></th>
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</thead>
<tbody>
<tr>
<td>Administration</td>
<td>IV push, infusion.</td>
<td>IV push.</td>
</tr>
<tr>
<td>Adult Dosage</td>
<td>IV push 4–12 mg/m² as a single agent at monthly intervals; or 1.5–1.7 mg/m²/day for 5 days as a continuous infusion.</td>
<td>0.4–1.4 mg/m²/week (2.5 mg typical single dose limit).</td>
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Special Populations. Other Conditions. Vinblastine and vinorelbine require substantial dosage reductions in heavily pretreated patients (i.e., drug or radiation therapy). Reduce vinorelbine dosage by 50% in patients in whom >75% of the liver is replaced by tumor or for granulocyte counts on the day of treatment of 1000–1499/µL; do not administer at lower WBC counts. Vinca alkaloids are eliminated extensively in the bile, and the dosages of vinblastine and vincristine must be reduced by approximately 50–75% with severe hepatobiliary dysfunction. Reduce vinorelbine dosages to 15 mg/m²/week for a serum total bilirubin of 2.1–3 mg/dL and 7.5 mg/m²/week for a bilirubin >3 mg/dL.

Dosage Forms. (Vinblastine) Inj 1 mg/mL, 10 mg vial. (Vincristine) Inj 1 mg/mL. (Vinorelbine) Inj 10 mg/mL.

Patient Instructions. (See Antineoplastics Class Instructions.)

Pharmacokinetics. Fate. Pharmacokinetics can be described by a two-compartment open model: an initial short phase with rapid tissue uptake (Vd approximating total body water) and a long terminal phase >1 day with a large Vd reflecting slow drug release from tissue binding sites—see below. Vinca do not effectively penetrate into the CNS or other fatty tissues and achieve their highest levels in the liver, gallbladder, and spleen. Approximately 50% of renally and fecally excreted products are closely related metabolites. An example is the formation of desacetyl vinblastine (which is more active on a weight basis than vinblastine) after vinblastine administration. Vinca alkaloids appear to be eliminated primarily in the bile and feces, some in the urine.

<table>
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<tr>
<th>Pharmacokinetic Parameters</th>
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<tr>
<td><strong>VINBLASTINE</strong></td>
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<tr>
<td>Vc (L/kg)</td>
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<tr>
<td>Vd (L/kg)</td>
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<tr>
<td>Urinary Excretion (cumulative)</td>
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<td>Fecal Excretion (cumulative)</td>
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<td><strong>t1/2α</strong> (min)</td>
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Adverse Reactions. Emetic potential is low (vincristine) to moderate (vinblastine and vinorelbine). Myelosuppression is the dose-limiting toxicity for vinblastine and vinorelbine, with the leukopenic nadir at 4–10 days; unless patients have been heavily pretreated with drugs or radiation, recovery from leukopenia is rather prompt, sometimes facilitating weekly or semimonthly drug administration. The major toxicity of vincristine is peripheral neuropathy manifested by paresthesias, constipation, jaw pain, decreased deep tendon reflexes, and rarely bladder atony or paralytic ileus; gut neurotoxicity occurs rarely with vinorelbine. All of these neurologic symptoms slowly resolve over 1 month and necessitate substantial dosage reduction if present at the time of drug administration. Seizures and ocular toxicity presenting as blurred vision or ptosis occur frequently. Mild laxatives or metoclopramide may be useful for constipation. The vincas are extremely toxic if inadvertently extravasated; hyaluronidase (150 units/mL) is sometimes effective as a local (subcutaneous) antidote. Vinorelbine also causes substantial phlebitis, which can be lessened by a short (6- to 10-min) infusion. Transiently severe pain in tumor masses occurs with vinblastine frequently. Alopecia is frequent with all agents.

Contraindications. Inadvertent intrathecal administration of any vinca alkaloid is fatal. (Vinblastine) Severe bone marrow compromise from prior therapy; uncontrolled infection. (Vincristine) Severe peripheral nervous system effects from prior doses, particularly paralytic ileus, tingling paresthesias, or decreased deep tendon reflexes; demyelinating form of Charcot-Marie-Tooth syndrome. (Vinorelbine) Pretreatment granulocyte count <1000/μL.

Precautions. Pregnancy. Use with caution in patients with neurologic deficiencies or hepatic disease.

Drug Interactions. Vinca administration (especially vincristine) has been associated with increased cellular retention of methotrexate (increased even in CNS tissues). Concurrent use of vincristine with zalcitabine can increase neuropathy.

Parameters to Monitor. (Vinblastine and vinorelbine) Obtain pretreatment and at least monthly WBC and hemoglobin/hematocrit assessments; (vincristine) obtain serial peripheral neurologic assessments; (all drugs) assess biliary function before making dosage adjustments for impaired hepatobiliary status and administering the drugs.

Notes. Protect these drugs from light and store under refrigeration. Place individual vincristine doses in an overwrap (eg, plastic bag) that is labeled, “Do not remove covering until the moment of injection. Fatal if given intrathecally. For intravenous use only.” Useful in hematologic neoplasms (primarily vincristine) and solid tumors, including non–small cell lung cancer in combination with cisplatin (vinorelbine) and refractory breast cancer and Kaposi’s sarcoma (vinblastine).

Monoclonal Antibodies

IODINE I-131 TOSITUMOMAB Bexxar

Pharmacology. This agent is a mouse-derived monoclonal antibody that binds to the CD-20 receptor of normal and malignant B-lymphocytes. The β-particle-emitting isotope $^{131}$I is coupled to the antibody, forming a selective radioimmuno-
Critical conjugates for patients with CD-20–positive non-Hodgkin’s lymphoma refractory to chemotherapy.201,202

**Adult Dosage.** In clinical trials, a two-stage dosage schedule has been used. First, patients are administered unlabeled antibody IV to suppress nonspecific binding sites. Next, patients are given trace-labeled doses of $^{131}$I-labeled antibody (15–20 mg containing 5 mCi IV) to assess antibody distribution. One week later, patients are given a 685 mg dose of unlabeled antibody IV, followed 1 day later by 2 individualized therapeutic (labeled) doses of 135 mg and 685 mg of antibody to deliver up to 75 cGy of whole-body radiation. A total body irradiation dosage of 55 cGy appears to be the maximum tolerated in patients who have undergone bone marrow transplantation. Give diphenhydramine 50 mg and acetaminophen 650 mg before each infusion.

**Adverse Reactions.** Nonhematologic toxicities after infusions are mild, with low-grade fever in 31% of patients and chills or rigors in 1%. Mild fatigue and nausea occur in 6–8% of radiolabeled infusions. The dose-limiting toxicity is hematologic suppression with grade 3 or 4 leukopenia and thrombocytopenia in 66% of patients who receive a whole-body radiation dose of 85 cGy. All patients develop complete or near-complete depletion of CD-19- and CD-20–positive B-cells from peripheral blood, recovering to normal levels in 3 months. Serum immunoglobulin levels are unchanged. Because of the short, single course of therapy, antimouse antibody (HAMA) reactions are uncommon. However, when they occur, they are usually marked by hypotension that can be treated with fluid hydration and vaspressors.

**Pharmacology.** Rituximab is a chimeric murine/human monoclonal antibody that binds to the CD-20 antigen on the surface of normal and malignant B-lymphocytes. This blocks normal CD-20–dependent signaling of cell-cycle initiation and differentiation, leading to apoptosis. After binding to CD-20, the free Fc portion of the antibody can recruit immune effector functions to cause lysis of B-cells of B-lymphocytes.203,204

**Administration and Adult Dosage.** IV infusion (not bolus) for relapsed or refractory low-grade B-cell non-Hodgkin’s lymphoma 375 mg/m$^2$ once weekly for 4 doses (days 1, 8, 15, and 22), diluted in NS or D5W. Premedicate the patient with diphenhydramine and acetaminophen and begin the first infusion at a rate of 50 mg/hr. Increase the rate by 50 mg/hr q 30 min to a maximum of 400 mg/hr if no hypotension or hypersensitivity develops. Subsequent infusions are started at a rate of 100 mg/hr and increased by 100 mg/hr q 30 min to the maximum rate of 400 mg/hr, if tolerated. If severe hypersensitivity reactions occur, stop the infusion and treat with diphenhydramine, acetaminophen, and a corticosteroid; for life-threatening reactions, add saline, epinephrine, and bronchodilators. If the reaction is not life-threatening, the infusion may be restarted at one-half of the earlier rate after symptoms subside.

**Special Populations.** **Pediatric Dosage.** Safety and efficacy not established.

**Geriatric Dosage.** Same as adult dosage.
Dosage Forms. **Inj** 10 mg/mL.

**Patient Instructions.** This drug might cause a flu-like syndrome including fever, chills, and weakness shortly after you receive it.

**Pharmacokinetics.** **Fate.** Peak serum levels are inversely correlated with the number of CD-20–positive B-cells. Levels average about 280 mg/L. Major sites of antibody distribution are to lymphoid cells of the thymus gland, white pulp of the spleen, and B-lymphocytes in peripheral blood and lymph nodes. Cl averages 0.054 L/hr; the antibody is still detectable in serum 3–6 months after the last dose.

$t_1/2$. Half-life is proportional to the dose (range 11–105 hr) with an average of 60 hr after a dosage of 375 mg/m².

**Adverse Reactions.** Most patients experience an infusion-related symptom complex with fever, chills, and rigors on the first infusion. Other frequent, acute infusion-related symptoms are nausea (18%), vomiting (10%), angioedema (13%), urticaria or pruritus (10%), and bronchospasm and rhinitis (8%). Hypotension and other acute effects are moderate or severe in 10% of the first doses. Overall, the frequency and severity of all reactions diminishes with subsequent injections. Most first-dose reactions occur within 30 min to 2 hr and resolve with slowing of the infusion rate for mild to moderate reactions or temporary halting the infusion and treating with supportive medications for severe reactions. Epinephrine is required only occasionally. Myelosuppression (neutropenia and thrombocytopenia) is typically mild and occurs in only 10% of patients, although long-term depletion of B-cells occurs in 70–80%; a minority also have decreased serum immunoglobulins. The frequency of grade 3 infections during the 4-week treatment period is 9% and grade 4 infections generally do not occur. Serious, sometimes fatal, skin reactions (bullous reactions, pemphigus) occur rarely.

**Contraindications.** Patients with a known type I hypersensitivity (or anaphylaxis) to mouse proteins.

**Precautions.** Pregnancy; lactation. Contraception is recommended in women of childbearing potential. The ability to respond immunologically to a vaccination is compromised after therapy; the safety of live virus vaccination is not known. Consider stopping antihypertensive medications on the day of treatment to reduce hypotensive reactions. Cardiac monitoring is recommended only in patients with pre-existing arrhythmias and angina that have worsened during the infusion.

**Drug Interactions.** Additive hypotension can occur in patients on antihypertensive therapy. There is no inhibition of cytotoxic activity in patients being treated for lymphoma with CHOP chemotherapy.

**Parameters to Monitor.** Monitor for allergic reactions and hypotension frequently during the infusion. Monitor CBC after therapy.

**TRASTUZUMAB**

**Herceptin**

**Pharmacology.** Trastuzumab is a humanized monoclonal antibody that binds to the HER2 protein found on the surfaces of some normal cells and plays a role in regulating cell growth. It is used only to treat tumors with an overexpression of HER2 protein. It is used alone in the treatment of metastatic breast cancer in pa-
tients who have been treated with chemotherapy or with paclitaxel in patients who have not had chemotherapy for their metastatic diseases.

**Adult Dosage.** IV for breast cancer in tumors with overexpression of HER2

4 mg/kg over 90 min as a loading dose, then 2 mg/kg weekly. Subsequent doses can be infused over 30 min if the loading dose was well tolerated. Do not administer as an IV push or bolus.

**Dosage Forms.** Inj 440 mg.

**Pharmacokinetics.** With the recommended dosage regimen, steady-state peak and trough concentrations are 123 and 79 mg/L, respectively. Trough serum levels are 1.5 times higher when given with paclitaxel, possibly because of inhibition of metabolism. The drug is distributed primarily in serum, with a $V_d$ of 0.44 L/kg. Pharmacokinetics appear to be dose related: Cl decreases and half-life increases with increasing dosages. Half-life averages 25 days (range 1–32 days) with the recommended regimen. Renal impairment appears not to affect pharmacokinetics.

**Adverse Reactions.** Side effects are frequent but usually not severe. Mild to moderate chills with or without fever occur in 40% of patients during the infusion and can usually be treated with acetaminophen, diphenhydramine, and/or meperidine. Other common side effects are diarrhea, pain, asthenia, nausea, vomiting, flu-like symptoms, cough, dyspnea, rash, edema, anemia, and leukopenia. Occasional serious reactions include anaphylaxis, thrombosis, pancytopenia, convulsions, apnea, hypoxia, and renal failure. Cardiac dysfunction and CHF have occurred. Use the drug with caution in pre-existing cardiac dysfunction; monitor cardiac function during therapy, and consider discontinuation if clinically important CHF develops. Serious infusion-related reactions including hypersensitivity (including anaphylaxis) and pulmonary events occur in about 0.25% of patients.

**Precautions.** Use with caution in patients with symptomatic intrinsic lung disease or extensive tumor involvement in the lungs. Interrupt infusion if the patient experiences dyspnea or clinically significant hypotension. Consider discontinuing therapy in patients who experience anaphylaxis, angioedema, or acute respiratory distress syndrome.

**Miscellaneous Antineoplastics**

**ASPARAGINASE**

Elspar

**PEGASPARGASE**

Oncaspar

**Pharmacology.** Asparaginase is the levo isomer of a macromolecular protein, isolated from *Escherichia coli* and other bacteria, that hydrolyzes the essential amino acid asparagine in the serum, thus depriving susceptible lymphocyte-derived malignancies of a necessary element for protein synthesis. Pegaspargase is a PEG-modified form of asparaginase that can be given to patients allergic to asparaginase. The drug is cell-cycle G-phase specific.

**Administration and Adult Dosage.** IM (preferably) or IV for combination therapy of acute leukemia (asparaginase) 200 IU/kg/day for 28 days,$^{205}$ or
1000–6000 IU/m^2/day for 5 days,\(^{206}\) or 20,000 IU/m^2/week;\(^ {207}\) (pegaspargase) 2500 IU/m^2 q 14 days.

**Special Populations.**

**Pediatric Dosage.** IM (preferably) or IV for combination therapy of acute leukemia (asparaginase) 1000–6000 IU/m^2/day for 5 days;\(^ {206,208}\) up to 20,000 IU/m^2/week; (pegaspargase) 2500 IU/m^2 q 14 days.

**Geriatric Dosage.** Same as adult dosage.

**Dosage Forms.** Inj (Asparaginase) 10,000 IU vial. (Pegaspargase) 750 IU/mL.

**Patient Instructions.** (See Antineoplastics Class Instructions.) Asparaginase often causes allergic reactions that can be life-threatening. This drug also can alter blood glucose levels and might worsen diabetes mellitus. Report any abdominal pain immediately because it might be a sign of pancreatitis.

**Pharmacokinetics.**

**Fate.** (Asparaginase) IV and IM produce equivalent serum levels. There is negligible distribution out of the vascular compartment, with minimal urinary and biliary excretion. Clearance is probably immune mediated. Asparaginase remains detectable in serum 13–22 days after administration.\(^ {209}\) (Pegaspargase) asparaginase is slowly released from pegaspargase and distributed in the body similarly to native asparaginase.

\[ t_{1/2} \]

(Asparaginase) \( \alpha \) phase 4–9 hr; \( \beta \) phase 1.4–1.8 days.\(^ {209}\) (Pegaspargase) 3.2 ± 1.8 days in patients hypersensitive to asparaginase; 5.7 ± 3.3 days in nonsensitive patients.

**Adverse Reactions.** Emetic potential is low. Moderate to severe non-dose-related hypersensitivity reactions occur in about 20–35\% of patients. (IM use might reduce and/or delay allergic complications);\(^ {205}\) a prophylactic antihistamine sometimes can be helpful. (See Precautions.) The drug is usually not myelotoxic. Transient blood glucose lowering followed by a pancreatitis-induced hyperglycemia can occur. Elevated serum cholesterol, severely elevated hepatic enzymes, steatosis, depressed clotting factors (especially profound for fibrinogen), and decreased albumin synthesis occur frequently. Lethargy and somnolence occur and might be more frequent in adults.\(^ {210}\) Fatal hyperthermia has been reported.

**Contraindications.** Anaphylactic reaction to commercial *E. coli* preparation; severe pancreatitis or history of pancreatitis.

**Precautions.** Onset of abdominal pain, serum amylase elevation, any changes in mental status, or severe elevation of prothrombin time require drug discontinuation. Some elevations of LFTs should be anticipated. Anaphylaxis can occur with any dose; ensure that emergency resuscitation equipment is available at the time of each dose. Intradermal scratch tests and desensitization procedures are not reliably predictive or preventive for anaphylaxis.\(^ {205,209}\)

**Drug Interactions.** None known.

**Parameters to Monitor.** Monitor serum hepatic enzymes, amylase, glucose, and prothrombin time routinely, and all vital signs during administration.

**Notes.** Reconstitute with NS or D5W (2 mL maximum for IM use); stable at least 24 hr; do not filter.
Pharmacology. Bleomycin is a mixture of 13 glycopeptide fractions produced by *Streptomyces verticillus*. Antineoplastic effects include single- and double-strand DNA scission, producing excision of thymine bases mediated through binding with ferric iron and subsequent production of highly reactive hydroxyl and superoxide radicals. It is cell-cycle phase specific, with maximal activity in the G2 (premitotic) phase.211

Administration and Adult Dosage. IM test dose 1–2 units may be useful in malignant lymphoma patients to assess exaggerated hyperpyrexia response. If no reaction occurs in 2–4 hr, give regular dose. SC, IM, or IV 10–20 units/m² 1–2 times/week.211 IV continuous infusion 15–20 units/day for 4–5 days.212 Experimental evidence in animals favors continuous administration to lessen pulmonary toxicity and maximize cell kill. A total lifetime dosage limit of 400 units is recommended to avoid pulmonary fibrosis. Intracavitary for malignant effusion 15–240 units (60 units for pleural effusion) in 50–100 mL or NS.213

Special Populations. Pediatric Dosage. SC, IM, or IV 10–20 units/m² 1–2 times a week in combination regimens. IV continuous infusion 15–20 units/m²/day for 4–5 days, usually as a single agent.

Geriatric Dosage. Same as adult dosage but use with caution in patients >70 yr and adjust dosage for age-related reduction in renal function.

Other Conditions. Dosage reduction has been recommended in renal impairment:214

<table>
<thead>
<tr>
<th>SERUM CREATININE (MG/DL)</th>
<th>PERCENTAGE OF DOSE RECOMMENDED</th>
</tr>
</thead>
<tbody>
<tr>
<td>2.5–4</td>
<td>25 (75% reduction)</td>
</tr>
<tr>
<td>4–6</td>
<td>20 (80% reduction)</td>
</tr>
<tr>
<td>6–10</td>
<td>5–10 (90–95% reduction)</td>
</tr>
</tbody>
</table>

Dosage Forms. Inj 15, 30 units.

Patient Instructions. (See Antineoplastics Class Instructions.) Report any coughing, shortness of breath, or wheezing. Skin rashes, shaking chills, or transient high fever can occur after administration. Hyperpigmentation of skin fold areas, scars, pressure areas, or sites of trauma can occur.

Pharmacokinetics. Fate. Poorly absorbed topically; roughly one-half of intracavitary-administered drug is systemically available (use this fraction to calculate lifetime exposure). After an IV dose of about 15 units/m², serum levels of 10–1000 milliunits/L are obtained.212 Steady-state levels during continuous infusion of 20 units/day are 50–200 milliunits/L.215 Vd is 0.27 ± 0.04 L/kg; Cl is 0.066 ± 0.018 L/hr/kg.10 Tissue inactivation is mediated by specific bleomycin-hydrolase, which is low in skin and lung, the two main toxicity targets of the drug.212,215 From 50% to 60% of a dose is recovered in the urine, 68% of this as unchanged drug.10 τ₁/₂α phase 24 min; β phase 3.1 ± 1.7 hr.10,212,215

Adverse Reactions. Emetic potential is moderately low. Alopecia, acute fever, and generalized erythema with edema, eventually leading to hyperpigmentation and skin
thickening, are frequent. The most serious long-term toxicity is pulmonary fibrosis manifested by dry cough, rales, dyspnea, and bilateral infiltrates. Pulmonary function studies show hypoxemia and reduced CO diffusing capacity. Pulmonary toxicity usually does not occur below 150 units/m², but the frequency increases to 55% at doses >283 units/m² and 66% at 360 units/m². Life-threatening pulmonary fibrosis is rare if dosage limits are observed. Prior chest radiotherapy, age >70 yr, and hypoxic ventilation predispose patients to toxicity. About 1% of high-dose bleomycin-treated patients die from pulmonary fibrosis. Low-dose hypersensitivity pneumonitis, which might be responsive to a glucocorticoid, also occurs.

Precautions. Use with extreme caution in patients with renal or pulmonary disease, in those with lymphoma, and in those >70 yr.

Drug Interactions. Inspired oxygen concentrations >35% can cause acute respiratory failure in bleomycin-treated patients.

Parameters to Monitor. Calculate cumulative dosage before and after each treatment. Monitor temperature initially, especially in lymphoma patients. Assess renal function before administering. Pulmonary damage is best monitored with CO diffusing capacity and forced vital capacity; specific serial pulmonary function studies have been suggested before and during therapy. Characteristic x-ray findings include changes suggestive of progressive diffuse bilateral fibrosis.

Notes. One milligram of bleomycin equals 1 unit of activity. Reconstituted solution is stable for 1 month under refrigeration and 2 weeks at room temperature. Incompatible with divalent cations (especially copper), ascorbic acid, and compounds with sulfhydryl groups.

**IMATINIB MESYLATE**

Pharmacology. Imatinib inhibits the abnormal Bcr-Abl tyrosine kinase created by the Philadelphia chromosome abnormality of CML, inhibiting proliferation and inducing apoptosis in leukemic cells with this abnormality. It may also inhibit the tyrosine kinase of platelet-derived growth factor and stem cell factor. It is used in CML after failure of interferon alfa therapy and in chemotherapy-resistant GI stromal carcinoma.

**Adult Dosage.** PO for the chronic phase of CML. 400 mg once daily, increasing to 600 mg once daily if conditions below are met; PO for the accelerated phase of CML or blast crisis 600 mg once daily, increasing to 400 mg bid if conditions below are met. All doses should be taken with a meal and a large glass of water. Dosages may be increased as indicated above in the absence of severe adverse reactions and severe non–leukemia-related neutropenia or thrombocytopenia for disease progression, failure to achieve a satisfactory hematologic response after ≥3 months of therapy, or with loss of a previous hematologic response.

**Dosage Forms.** Cap 100 mg.

**Pharmacokinetics.** Oral bioavailability is 98% with a peak at 2–4 hr. It is 95% plasma protein bound. Metabolism is primarily by CYP3A4 to the N-desmethyl metabolite that has activity similar to the parent drug. Imatinib Cl is 0.14–0.16 L/hr/kg. The drug is eliminated in feces, mostly as metabolites. Elimination half-lives are 18 and 40 hr for the drug and active metabolite, respectively.
Adverse Reactions. Most adverse reactions are mild to moderate and more frequent during the accelerated phase and especially during blast crisis. The most frequent are nausea, vomiting, and periorbital or lower limb edema. Edema is occasionally severe and can be managed by diuretics, supportive measures or imatinib dosage reduction. Muscle cramps and pain, hemorrhage, skin rash, headache, fatigue, abdominal pain, arthralgia and fever are also frequent. Dose-related neutropenia (duration 2–3 weeks) and thrombocytopenia (duration 3–4 weeks) also occur, and respond to dosage reduction or interruption of therapy. Elevated transaminases and bilirubin can occur, sometimes requiring dosage reduction or treatment interruption; one death from hepatotoxicity has been reported.

Drug Interactions. Inhibitors and inducers of CYP3A4 are expected to alter the metabolism of imatinib and should be used with caution. Imatinib decreases the metabolism of simvastatin, apparently through CYP3A4 inhibition. Use other drugs with caution that are metabolized by CYP3A4. Patients requiring anticoagulation should receive heparin or a LMWH rather than warfarin.

Pharmacology. Tretinoin (all-trans-retinoic acid) is a modified form of vitamin A used in the treatment of acute promyelocytic leukemia. It causes immature promyeloblasts to differentiate into mature granulocytes, thereby halting cell division and inducing complete remissions in up to 90% of patients. Resistance rapidly develops during therapy because of accelerated drug catabolism to the 4-oxo metabolite, which is excreted in the urine, increased cellular retinoic acid binding protein (II), and tumors with high levels of a mutated α-retinoic acid receptor.18–20

Pediatric Dosage. PO 45 mg/m²/day as a single dose until remission is obtained.

Dosage Forms. Cap 10 mg.

Pharmacokinetics. Peak serum levels of 294 g/L occur 1–2 hr after a dose; the serum half-life is 0.8 hr.

Adverse Reactions. Hyperleukocytosis and effects typical of hypervitaminosis A (e.g., headache, dry skin and mucosa, cheilitis, bone pain, hypertriglyceridemia) occur. Tolerance to these effects develops rapidly, and skin creams, lip balms, and eye and nasal drops are helpful. Liver and renal function test elevations occur occasionally.

Chemoprotectants

AMIFOSTINE Ethylol

Pharmacology. Amifostine is a phosphorothiol compound metabolized by membrane-bound alkaline phosphatase to an active sulfhydryl form capable of binding electrophilic metabolites from DNA-binding anticancer agents or ionizing radiation. It is used prophylactically to block cisplatin-induced nephrotoxicity and neurotoxicity without altering antitumor efficacy in patients with advanced ovarian cancer.21
Administration and Adult Dosage. IV to block cisplatin toxicity 910 mg/m² in NS over ≤15 min, beginning 30 min before cisplatin.

Special Populations. Pediatric Dosage. Safety and efficacy not established.

Geriatric Dosage. Same as adult dosage, but limited experience exists in patients >70 yr.

Other Conditions. In patients who develop rare symptomatic acute hypocalcemia, reduce dose to 740 mg/m² and extend infusion time. Reduce dosage in patients who developed hypotension (drop of 15–20 mm Hg systolic) with prior courses. When used as a radioprotective agent, the maximally tolerated dosage is 340 mg/m² for 4 days/week.

Dosage Forms. Inj 50 mg/mL.

Patient Instructions. The severity of chemotherapy-induced nausea and vomiting can increase with amifostine. Drink lots of fluids in the hours before receiving this medication to reduce its toxicity.

Pharmacokinetics. Fate. The mean peak serum level is 100 μmol/L after an IV dose of 740 mg/m². Vdss is 6.4 ± 1.5 L; Cl is 2.2 ± 0.4 L/min. Relatively little unchanged drug (1.1% of a dose), actifostine (1.4% of a dose), or disulfide metabolite (4.2% of a dose) is excreted renally. $t_{1/2}$ α phase 0.88 ± 0.12 min; β phase 8.8 ± 2 min.

Adverse Reactions. Toxic effects are all acute and include transient hypotension during or immediately after drug infusion; nausea; and vomiting. Prophylactic antiemetics before administration can reduce nausea and vomiting. Rapid infusion (≤15 min) and aggressive hydration can lessen or eliminate hypotensive toxicity. Stopping the infusion and placing the patient in the Trendelenberg position usually reverses the hypotension. Other less serious but common reactions are sneezing (27%), a flushed sensation (26%), somnolence (10–20%), a sensation of cold hands, or a metallic taste in the mouth (<5% each).

Contraindications. Allergy to aminothiol compounds or mannitol.

Precautions. Do not administer concurrently with or after cisplatin infusion. Use with caution in patients in whom hypotension or nausea might pose a serious risk. There is limited experience in patients with pre-existing cardiac or cardiovascular conditions such as CHF, angina pectoris, history of stroke, or TIAs.

Drug Interactions. Amifostine can reduce the systemic exposure to paclitaxel but not to docetaxel, cisplatin, carboplatin, or cyclophosphamide.

Parameters to Monitor. Monitor blood pressure frequently during drug administration and immediately after infusion.

Notes. Amifostine has been safely combined with ionizing radiation, carboplatin, and cyclophosphamide. Amifostine markedly reduces mucositis when it is combined with carboplatin and radiotherapy in the treatment of head and neck cancer. Amifostine allows dose escalation of paclitaxel and has hematopoietic activity in the investigational treatment of refractory myelodysplastic syndrome.
**DEXRAZOXANE**

**Pharmacology.** Dexrazoxane, a cardioprotectant for anthracyclines, is the water-soluble dextro isomer of razoxane. Dexrazoxane’s two piperazinedione rings open to form sites that chelate intracellular ferrous ions, blocking the formation of doxorubicin-iron complexes capable of forming membrane-damaging oxygen free radicals. Dexrazoxane can extend doxorubicin cumulative dosage in patients with breast cancer. It also reduces the risk of short-term subclinical cardiotoxicity in pediatric sarcoma patients. It does not alter the pharmacokinetics of doxorubicin.228–230

**Administration and Adult Dosage.** IV as a cardioprotectant give in a 10:1 dexrazoxane:doxorubicin mg/m² ratio (eg, 500 mg/m² dexrazoxane:50 mg/m² doxorubicin). Infuse IV over 15 min, beginning not more than 30 min before an IV push dose of doxorubicin.

**Special Populations.**

**Pediatric Dosage.** Safety and efficacy not established, but it has been used in a 20:1 dosage ratio (dexrazoxane:doxorubicin) in children with sarcomas.230

**Geriatric Dosage.** Same as adult dosage.

**Dosage Forms.** Inj 250, 500 mg.

**Patient Instructions.** (See Antineoplastics Class Instructions.)

**Pharmacokinetics.** Fate. The mean peak serum level after a dose of 500 mg/m² given over 15 min is 36.5 mg/L (136 μmol/L). The drug is not protein bound. \( V_d \) is 22 L/m² or approximately body water; Cl averages 7.9 L/hr/m². About 42% is renally eliminated as parent drug and mono- and diacid amide metabolites.231

\( t_{1/2} \)  
\( \alpha \) phase 0.2–0.3 hr; \( \beta \) phase 2.1–2.5 hr.231

**Adverse Reactions.** Dexrazoxane has little toxicity but slightly increases the myelosuppressive and emetogenic toxicities of doxorubicin-containing regimens. Pain on injection also occurs.

**Contraindications.** None known.

**Precautions.** Avoid use with bleomycin. Do not administer after doxorubicin.

**Drug Interactions.** None known.

**Parameters to Monitor.** WBC counts at nadir (7–11 days) after doxorubicin.

**Notes.** Dexrazoxane does not reduce the antitumor activity of fluorouracil, doxorubicin, and cyclophosphamide regimens in advanced breast cancer.228 Effects on other antineoplastics are unknown.

**MESNA**

**Pharmacology.** Mesna (2-mercaptopethanesulfonate) is a sulfhydryl compound that minimizes urotoxicity from the alkylating agents cyclophosphamide (CTX) and ifosfamide (IFX) by binding to the irritant metabolite acrolein in the urinary bladder to prevent hemorrhagic cystitis.35,232

**Administration and Adult Dosage.** IV or PO (ampule contents dissolved in water or juice) in 3 doses as a percentage of the dose of ifosfamide or cyclophos-
phamide. Oral administration is not recommended for patients with poor compliance or those experiencing nausea or vomiting.

<table>
<thead>
<tr>
<th>TIME BEFORE OR AFTER CTX OR IFX</th>
<th>PERCENTAGE OF CTX OR IFX DOSE</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>IV MESNA ROUTE</td>
</tr>
<tr>
<td>15 min before</td>
<td>20</td>
</tr>
<tr>
<td>4 hr after</td>
<td>20</td>
</tr>
<tr>
<td>8 hr after</td>
<td>20</td>
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</table>

CTX = cyclophosphamide; IFX = ifosfamide.

**Special Populations.**

**Pediatric Dosage.** Same as adult dosage.42,234

**Geriatric Dosage.** Same as adult dosage.

**Dosage Forms.** 100 mg/mL.

**Patient Instructions.** This agent does not have antitumor activity but is essential to reduce or prevent permanent bladder damage from chemotherapy.

**Pharmacokinetics.**

**Fate.** About 48% is orally absorbed.235 Vₐ is 0.65 L/kg; Cl is 1.23 L/hr/kg. Mesna is oxidized to the inactive dimer, dimesna, which does not inactivate CTX or IFX metabolites in the serum. About 60% of the dimesna is converted back to mesna in the renal tubule and delivered to the bladder in the active sulfhydryl form. About two-thirds of a dose is excreted in the urine, one-half as mesna and one-half as dimesna.236

\[ t_{1/2} \] (Mesna) 22 min; (dimesna) 1.2 hr.236

**Adverse Reactions.** When administered alone, mesna produces little if any serious toxicity.236 GI effects (eg, diarrhea, nausea, and, rarely, vomiting) of CTX or IFX might be slightly greater when mesna is administered. Other CTX or IFX toxicities such as myelosuppression or alopecia are not altered by mesna. With oral administration, disagreeable sulfur odor might lessen palatability unless the drug is diluted with cola or juice.

**Drug Interactions.** Mesna inhibits the antitumor activity of cisplatin and carboplatin but not of other anticancer agents.

**Notes.** Mesna is compatible with solutions of CTX or IFX and has been administered concurrently as a continuous infusion of both agents at equal doses in the same infusion container.237 It is stable in D5W or NS for at least 96 hr.

**Immunosuppressants**

**General Precautions for Immunosuppressants.** Immunosuppression increases the risk of infectious complications. Serious opportunistic infections can occur during immunosuppressive therapy. Long-term immunosuppression also increases the risk of malignancy or lymphoproliferative disease. Vaccinations might be less effective during immunosuppression. Live or live attenuated vaccines might proliferate excessively in immunosuppressed patients and should be avoided.
Pharmacology. Antilymphocyte immune globulins are polyclonal IgG purified from sera of horses or rabbits immunized with human thymus lymphocytes. These drugs are immunosuppressants that inhibit cell-mediated immunity. The immunosuppressive effects of antilymphocyte immune globulins may be secondary to clearance of alloreactive T-lymphocytes from the plasma. However, the exact pharmacologic mechanism of action has not been elucidated.

Administration and Adult Dosage. Antilymphocyte immune globulins are generally used in conjunction with other immunosuppressants. Intradermal sensitivity testing to identify patients at risk for anaphylaxis is strongly recommended before administration of equine lymphocyte immune globulin. Freshly diluted equine lymphocyte immune globulin (5 μg in 0.1 mL 0.9% NaCl) should be administered intradermally on the anterior aspect of one forearm, with intradermal administration of 0.9% NaCl 0.1 mL on the contralateral forearm as a control. During the hour after administration, the skin test should be observed q 15–20 min for swelling, urticaria, pruritus, and wheel or erythema. A positive skin test is defined as local wheel or erythema formation ≥10 mm in diameter. If the skin test is positive, the risk of serious hypersensitivity or anaphylaxis from drug administration should be weighed carefully against the anticipated benefits of drug administration. A systemic reaction to the skin test generally precludes further administration of equine lymphocyte immune globulin. Administration of equine lymphocyte immune globulin to patients after a positive skin test or systemic reaction to the skin test should be done only in a facility capable of supporting life-threatening allergic reactions. The skin test is not 100% predictive of subsequent hypersensitivity reactions. Allergic reactions and anaphylaxis to equine lymphocyte immune globulin have been reported after a negative skin test. The manufacturer does not recommend a test dose before administration of rabbit antithymocyte globulin.

IV for prevention of renal allograft rejection (equine lymphocyte immune globulin) 15 mg/kg/day for 14 doses, followed by the same dose every other day for an additional 14 days. This regimen administers up to 21 doses of equine lymphocyte immune globulin in 28 days. The first dose of equine lymphocyte immune globulin should be administered within 24 hr before or after surgery; (rabbit antithymocyte globulin) 1.5 mg/kg/day beginning on the day of surgery, for a total of at least 7 doses, has been used for prevention of renal allograft rejection.238 IV for treatment of renal allograft rejection (equine lymphocyte immune globulin) same dosage regimen as above, with administration of the first dose at the diagnosis of the initial rejection episode; (rabbit antithymocyte globulin) 1.5 mg/kg/day for 7–14 days. IV for prevention of rejection after heart transplantation (rabbit antithymocyte globulin) 4 mg/kg/day administered as an IV infusion over 6 hr on postoperative days 1–5.239 IM for prevention of rejection after heart transplantation (rabbit antithymocyte globulin) 1.5 mg/kg/day or 200 mg/day for
3–7 days has been administered. IV for treatment of aplastic anemia (equine lymphocyte immune globulin) 10–30 mg/kg/day for 8–14 days, followed by the same dose every other day for 14 days, has been used. This regimen administers up to 21 doses of equine lymphocyte immune globulin in 28 days. An alternative regimen uses 40 mg/kg q 24–48 hr for 3–4 doses; (rabbit antithymocyte globulin) 3.5 mg/kg/day for 5 days has been administered with cyclosporine and filgrastim for treatment of severe aplastic anemia unresponsive to equine lymphocyte immune globulin. IV for prevention of acute graft-versus-host disease (GVHD) in allogeneic bone marrow transplant recipients (equine lymphocyte immune globulin) 7–15 mg/kg every other day for 6 doses; (rabbit antithymocyte globulin) 2, 3.75, or 5 mg/kg/dose for 4–5 doses before unrelated bone marrow transplantation. IV for treatment of moderate-to-severe steroid-refractory acute GVHD (equine lymphocyte immune globulin) 7–15 mg/kg for 6 doses or as indicated by the patient’s clinical status. IV for skin allograft survival in patients with full-thickness burns (equine lymphocyte immune globulin) 10–15 mg/kg every other day is generally used; however, doses of 5 mg/kg every other day up to 40 mg/kg/day have been given. The duration of therapy is generally 40–60 days, ending when skin allografts cover <20% of the BSA. The maximum tolerated cumulative dosage of antilymphocyte polyclonal immune globulins has not been determined. A total of 50 doses of equine lymphocyte immune globulin has been administered over 4 months, and four 28-day courses of 28 equine lymphocyte immune globulin doses have been administered in renal allograft recipients without changing the frequency, severity, or character of adverse drug reactions. Intravenous administration (equine lymphocyte immune globulin) dilute in 0.45% or 0.9% NaCl to a final concentration ≤4 mg/mL and infuse slowly over 4–8 hr; (rabbit antithymocyte globulin) after reconstitution with the diluent provided, dilute to a final concentration of 0.5 mg/mL in 0.9% NaCl or 5% dextrose injection. Infuse the first dose at 0.25 mg/kg/hr (1.5 mg/kg/6 hr). In the absence of moderate-to-severe adverse effects, infuse subsequent doses over 4 hr. Antilymphocyte polyclonal immune globulins should be infused through an inline filter with pore sizes of 0.22–1 μ. Premedication and as-needed administration of a corticosteroid, acetaminophen, and an antihistamine are common practice intended to reduce infusion-related adverse effects.

Special Populations. Pediatric Dosage. Same as adult dosage.

Geriatric Dosing. Same as adult dosage.

Dosage Forms. Inj (equine lymphocyte immune globulin) 50 mg/mL; (rabbit antithymocyte globulin) 25 mg.

Patient Instructions. (Equine lymphocyte immune globulin) This medicine can cause serious allergic symptoms, especially in people allergic to horses and horse products. You will receive a skin test to check for allergy to this product. (Rabbit antithymocyte globulin, equine lymphocyte immune globulin) You might experience fever, shaking, and chills when this medication is being given. You may be given additional medications to reduce these side effects.

Pharmacokinetics. Fate. (Equine lymphocyte immune globulin) Peak concentrations of 727 μg/L occur with repeated doses of 10 mg/kg. Systemic distribution
of equine immune globulin is not well defined. In vitro studies predict binding to circulating lymphocytes, granulocytes, and platelets. Binding to bone marrow cells, plus thymus and testis cell membranes, occurs in vitro. (Rabbit antithymocyte globulin) IV infusion of 1.25–1.5 mg/kg/day yields a peak concentration of 10–40 μg/L after the first dose and 23–170 μg/L after repeated doses. \( V_d \) is 0.12 L/kg.\(^{243} \)

\( t_{1/2} \). (Equine lymphocyte immune globulin) 3–9 days; (rabbit antithymocyte globulin) 14–45 days.\(^{243} \)

**Adverse Reactions.** Anaphylaxis can occur anytime during therapy. If signs or symptoms of anaphylaxis occur, the infusion must be stopped immediately and appropriate management must be initiated. Serum sickness occurs frequently. The onset of serum sickness is typically 6–18 days after initiation of therapy with antilymphocyte immune globulins. A morbilliform rash generally starts as a truncal distribution of faint macules, with subsequent progression to the extremities. The macules can become confluent. Erythema can spread to involve palms of the hands and soles of the feet. Antihistamines are helpful for pruritus-associated adverse effects. Although not clearly shown to reduce serum sickness-related adverse effects, corticosteroids have been used. Antilymphocyte immune globulins might bind formed elements in the blood other than T-lymphocytes and promote splenic clearance of these blood constituents. Subsequently, patients might experience acute normochromic normocytic anemia, thrombocytopenia, or leukopenia during administration of antilymphocyte immune globulins that is reversible with drug discontinuation. Immunosuppression increases the risk of infectious complications from opportunistic and pathogenic microbes. Rare adverse effects reported with antilymphocyte immune globulins include Epstein-Barr virus infections, lymphoproliferative disorders and (equine lymphocyte immune globulin) periorbital edema, seizures, acute renal failure, headache, hypertension, edema, CHF, bradycardia, adult respiratory distress syndrome, myocarditis, pancytopenia, LFT abnormalities, hyperglycemia, and transient myopia; (rabbit antithymocyte globulin) tachycardia, dyspnea, and dizziness.\(^{244–247} \)

**Contraindications.** (Equine lymphocyte immune globulin) allergy to equine lymphocyte immune globulin, horse serum, or horse products; (rabbit antithymocyte globulin) hypersensitivity or anaphylaxis to rabbit proteins; acute viral illness.

**Precautions.** Pregnancy; lactation.

**Drug Interactions.** None identified.

**Parameters to Monitor.** Observe for anaphylaxis during infusion and serum sickness 6–18 days after initiation of therapy. CBC and platelet count q 1–3 days during therapy.

**Notes.** Equine lymphocyte immune globulin is also known as ATG, antithymocyte globulin, antithymocyte gamma globulin, antithymocyte immunoglobulin, and horse antihuman thymocyte gamma globulin. Rabbit antithymocyte globulin is also known as r-ATG or RATG. Lot-to-lot variation of immunosuppressive potency and avidity for formed blood elements can occur with these products.
AZATHIOPRINE

Pharmacology. Azathioprine is a thiopurine prodrug of 6-mercaptopurine (6-MP). Conversion to 6-MP with subsequent phosphoribosylation yields antimetabolites capable of inhibiting DNA and RNA synthesis. The metabolite 6-methylmercaptopurine ribotide is a potent inhibitor of de novo purine synthesis. T-lymphocytes are sensitive to inhibition of de novo purine synthesis because these cells lack efficient salvage pathways to maintain adequate intracellular stores.

Administration and Adult Dosage. PO or IV for immunosuppression after solid organ transplantation 3–5 mg/kg/day as a single daily dose beginning the day of, or 1–3 days preceding, transplantation. Maintenance dosage is 1–3 mg/kg/day as a single daily dose. PO for rheumatoid arthritis 1 mg/kg/day in 1 or 2 doses. The dosage may be increased after 6–8 weeks if indicated by disease response and patient tolerance. Increase the dosage in increments of 0.5 mg/kg/day q 4 weeks to a maximum of 2.5 mg/kg day. (See Drug Interactions.)

Special Populations. Pediatric Dosage. PO or IV for immunosuppression after renal transplantation same as adult dosage.

Geriatric Dosage. Same as adult dosage.

Dosage Forms. Tab 50 mg; Inj 100 mg.

Patient Instructions. This medication may be taken with food to reduce stomach upset. Notify your physician if any of the following symptoms occur: unusual bleeding or bruising, fever, sore throat, mouth sores, abdominal pain, yellowing of the eyes, pale stools, or dark urine, or if nausea, vomiting, diarrhea, skin rash, or joint pains become severe or persist.

Missed Doses. Take a missed dose as soon as possible. If you take the drug once daily and it is time for the next dose, take it at the regular time. Do not double the dose. If you take two or more doses daily, and it is time for the next dose, take both doses together. If two or more doses are missed, contact your physician.

Pharmacokinetics. Onset and Duration. Onset of immunosuppression occurs within days to weeks. Immunosuppression continues for days to weeks after drug discontinuation.

Serum Levels. No correlation between serum concentrations and efficacy or toxicity has been defined.

Fate. After oral absorption, conversion to 6-MP occurs rapidly.1 (See Mercaptopurine.)

\[ t_{1/2} \] (Azathioprine) 9.6 ± 4.2 min; (6-MP) 0.9 ± 0.37 hr.110

Adverse Reactions. Dose-related bone marrow suppression, which can include leukopenia, thrombocytopenia, and anemia, occurs frequently. Macrocytic anemia, with megaloblastic features, or selective erythrocyte aplasia can occur with long-term azathioprine administration. Skin rash is a common adverse effect. Mouth sores can occur. Dose-related nausea and vomiting are frequent and can be reduced by administration in divided doses. Rare GI hypersensitivity characterized by severe nausea and vomiting, diarrhea, hyperpyrexia, malaise, myalgia, and
LFT abnormalities can occur early in the course of therapy. GI hypersensitivity is reversible with discontinuation of azathioprine and can recur with rechallenge. Hepatic veno-occlusive disease of the liver, secondary lymphomas, and other malignancies can occur with long-term administration. Azathioprine is teratogenic. Rare adverse effects include pancreatitis, constrictive lung disease, renal failure, alopecia, arthralgia, and retinopathy.

**Contraindications.** Pregnancy in patients treated for rheumatoid arthritis.

**Precautions.** Pregnancy; lactation. (See also General Precautions for Immunosuppressants.)

**Drug Interactions.** To reduce the risk of life-threatening myelosuppression, azathioprine dosage must be reduced to 25–33% of the normal dosage in patients receiving allopurinol. Enhanced bone marrow suppression can occur with concurrent use of drugs that inhibit hematopoiesis. Concurrent corticosteroids used for immunosuppression can mask fever.

**Parameters to Monitor.** Monitor CBC and platelet count weekly for 1 month after initiation of therapy or any dosage increase. Then, for patients with stable hemograms, the CBC and platelet count may be monitored twice monthly for 2 months and then monthly for the duration of therapy. Monitor serum transaminases, alkaline phosphatase, and total bilirubin periodically. Observe for signs of infection.

**Notes.** Azathioprine is used in combination with other immunosuppressants as an adjunct in the prevention of renal allograft rejection and for the prevention of solid organ rejection for cardiac and hepatic allografts. It is rarely used for the management of acute or chronic GVHD in allogeneic bone marrow transplant recipients because it markedly increases the risk for infections.

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**CYCLOSPORINE**  
**Gengraf, Neoral, Sandimmune, SangCya, Various**

**Pharmacology.** Cyclosporine is a cyclic polypeptide immunosuppressant produced by the fungus *Tolypocladium inflatum* Gams. The intracellular drug-ligand complex formed by cyclosporine and cyclophilin indirectly blocks T-lymphocyte activation by inhibiting calcineurin-mediated dephosphorylation of transcription factors necessary for IL-2 transcription.

**Administration and Adult Dosage.** PO for prophylaxis of organ rejection or GVHD 8–12 mg/kg/day in 2 divided doses depending on the type of transplant and the other immunosuppressants being given. To hasten achievement of an immunosuppressant blood level, an oral loading dose of cyclosporine 15 mg/kg may be administered. Cyclosporine is usually started 4–12 hr before surgery. Maintenance dosage is based on cyclosporine blood levels, the risk of organ rejection or GVHD, and patient tolerance. PO for rheumatoid arthritis 2.5 mg/kg/day in divided doses bid. As patient tolerance allows, dosage may be increased by 0.5–0.75 mg/kg/day at 8 weeks and again at 12 weeks, to a maximum of 4 mg/kg/day. PO for psoriasis 2.5 mg/kg/day in divided doses bid. After 4 weeks of therapy, as patient tolerance allows, the dosage may be increased by 0.5–0.75 mg/kg/day q 2 weeks, to a maximum of 4 mg/kg/day. An IV loading dose of 3–4 mg/kg might be useful in patients with low cyclosporine levels during...
periods of mild-to-moderate diarrhea with oral maintenance therapy. **IV for patients unable to take oral medication** 2–6 mg/kg/day in 1–2 divided doses. **IV for prevention of GVHD** 3–4 mg/kg/day in 2 divided doses q 12 hr. Cyclosporine is generally started 1–2 days before bone marrow transplantation. Drug-induced mucositis or diarrhea usually necessitate use of IV cyclosporine in allogeneic bone marrow transplant recipients. Dilute IV cyclosporine in a glass container (it might leach plasticizers from PVC containers) with D5W or NS to a concentration of 50 mg/20–100 mL. Doses may be infused over 2–6 hr or given as a continuous infusion. **Conversion from IV to PO administration** the ratio of IV:PO dosage is typically 1:3 to 1:4 for Sandimmune capsules or 1:1 to 1:3 for microemulsion capsules or solution (Gengraf, Neoral, SangCya, various). **Conversion from PO Sandimmune to microemulsion capsules or solution** (Gengraf, Neoral, SangCya, various) give the same daily dosage or reduce microemulsion dose by 30% with prompt dosage adjustment based on subsequent blood levels. A cyclosporine blood level should be drawn 48 hr after dosage form conversion. Because of better bioavailability, maintenance dosages of the microemulsion formulation are usually lower than Sandimmune dosages. **Interchange of various microemulsion capsules or solutions** (Gengraf, Neoral, SangCya, various) give the same daily dosage with prompt dosage adjustment based on subsequent blood levels. A cyclosporine blood level should be drawn 48 hr after interchange. **Discontinuation** cyclosporine may eventually be discontinued in certain renal or allogeneic bone marrow transplant recipients. Cyclosporine dosage must be decreased gradually over time to reduce the risk of reactive immune stimulation and graft rejection or GVHD.

**Special Populations. Pediatric Dosage.** Initial dosage same as adult dose. Adjustment based on blood levels. Children may require higher weight-based maintenance dose.

**Geriatric Dosage.** Same as adult dosage. Age-related loss of renal function and comorbid conditions can predispose geriatric patients to cyclosporine-induced nephrotoxicity or hypertension.

**Other Conditions.** Use IBW to calculate initial dosage in obese patients.

**Dosage Forms.** Cap (Neoral, various) 25, 100 mg; (Sandimmune) 25, 50, 100 mg; Oral Soln 100 mg/mL; Inj (Sandimmune) 50 mg/mL.

**Patient Instructions.** Take this medication on a regular schedule relative to the time of day and meals. Do not discontinue it unless directed to do so. Sandimmune cannot be interchanged with any other brands. The oral solution may taste better if mixed with another liquid. Sandimmune may be mixed with milk, chocolate milk, or orange juice. Neoral may be mixed with orange juice or apple juice. Using a glass container, mix the cyclosporine solution with the milk or juice, stir well, and drink immediately to ensure that the entire cyclosporine dose is swallowed. Do not refrigerate the oral solution. Use the oral solution within 2 months after opening. Grapefruit juice can interact with cyclosporine. Talk to the physician or coordinator who monitors your cyclosporine before drinking grapefruit juice and before starting, stopping, or changing the dose of any medication.
**Missed Doses.** Take a missed dose as soon as possible if you remember within 12 hours. If it is within 2 hours of the next dose, skip the missed dose and do not double the dose. If you miss 2 or more doses, contact the physician or coordinator who monitors your cyclosporine.

**Pharmacokinetics. Serum Levels.** The serum (blood) concentration-response relationship is not completely defined. Trough blood or serum levels are monitored for toxicity. Therapeutic and toxic concentrations vary with assay, biologic fluid, and time post-transplant. Therapeutic serum concentrations are: polyclonal radioimmunoassay (RIA) 100–250 μg/L; monoclonal RIA 50–125 μg/L; high-performance liquid chromatography (HPLC) 50–125 μg/L. Therapeutic whole blood concentrations are: polyclonal RIA 200–800 μg/L; monoclonal RIA 150–400 μg/L; HPLC 150–400 μg/L. In routine clinical practice, spurious serum drug levels can result from in vitro drug redistribution. Artifactual serum or whole blood levels can occur when blood is drawn through the same central venous line used for IV cyclosporine administration.

**Fate.** Oral absorption is formulation dependent (See Notes). Sandimmune absorption is incomplete and variable. The mean bioavailability of Sandimmune is 34%; however, the reported range is 5–90%. Absorption of Sandimmune is improved after a high-fat meal. Peak concentrations occur 2–6 hr after ingestion of Sandimmune capsules or oral solution. Absorption of cyclosporine microemulsion formulations is independent of food intake; peak concentrations occur 1.5–2 hr after ingestion of microemulsion capsules or solution. Factors that can decrease cyclosporine absorption are diarrhea, gastroenteritis, and short small bowel. Absorption may be reduced in allogeneic bone marrow transplant patients because of residual gut damage from intensive chemotherapy, radiation, or GVHD. Bioavailability of AB therapeutic equivalent cyclosporine capsules and liquid can vary by 20–30% for a particular patient or when mixed with various juices. Systemic cyclosporine distributes to erythrocytes (45%), leukocytes (15%), and plasma lipoproteins (35%). Marked elevations of plasma lipoproteins can increase measured cyclosporine levels without proportional changes in therapeutic or toxic effects. $V_{ss}$ (whole blood, HPLC) is $4 \pm 0.8$ L/kg in renal transplant patients and $5.3 \pm 2.9$ L/kg in bone marrow transplant patients. Cl (whole blood, HPLC) is $0.4 \pm 0.2$ L/kg/hr in renal or liver transplant patients and $0.6 \pm 0.4$ L/kg/hr in allogeneic bone marrow transplant recipients. Cyclosporine is extensively metabolized by CYP3A. At least 25 metabolites, some with immunosuppressant activity, have been identified. Cyclosporine and its metabolites are cleared primarily in the bile. About 3% is excreted in the urine as cyclosporine and metabolites. Less than 1% is excreted in the urine as unchanged cyclosporine.

**$t_{1/2}$.** (Whole blood, HPLC) $10 \pm 3.5$ hr, possibly prolonged in hepatic failure.

**Adverse Reactions.** Acute nephrotoxicity, which generally occurs during the first month of treatment, is characterized by Crs increasing by $\geq 0.3$ mg/dL/24 hr or $\geq 30\% / 24$ hr and usually abates with interruption of drug therapy. Dosage reduction may be required for continuation of therapy. Chronic progressive renal toxicity is characterized by a slow continual increase in Crs and BUN, mild proteinuria, and tubular dysfunction. Crs $>2$ mg/dL in adult patients or doubling of Crs is an indication for interruption of therapy or dosage reduction. Electrolyte abnormalities,
including hypomagnesemia, hypokalemia, hyperkalemia, and renal tubular acidosis, are consequences of cyclosporine-induced nephrotoxicity. Concurrent administration of nephrotoxic drugs increases the likelihood of renal dysfunction. Hypertension occurs frequently. Calcium channel blockers and clonidine are suitable agents for cyclosporine-induced hypertension because they do not have deleterious effects on renal blood flow. Fine tremors occur frequently and can persist and worsen after drug discontinuation. Neurotoxicity can also present as seizures, cortical blindness, paresthesias, hyperesthesia, headache, or expressive aphasia. Patients with low serum cholesterol may be at increased risk of neurotoxicity. Anaphylactic reactions to cyclosporine or the solubilizing agent, polyoxyethylated castor oil, can occur. Ethanol is a minor constituent in the intravenous and oral formulations. Cyclosporine-induced cholestasis is dose related and transient. Elevated serum transaminases and hypertriglyceridemia can occur. Additional side effects are hemolytic uremic syndrome, pancreatitis, hirsutism, gingival hyperplasia, nausea, vomiting, acne, and gynecomastia. Leukopenia, anemia, and thrombocytopenia occur rarely.

**Contraindications.** Allergy to cyclosporine or polyoxyethylated castor oil.

**Precautions.** Pregnancy; lactation. Use cautiously in patients with aldehyde dehydrogenase 2 (ALDH2) deficiency. (See also General Precautions for Immunosuppressants.)

**Drug Interactions.** Numerous important drug interactions have been identified. Additive or synergistic renal toxicity can occur with concomitant administration of nephrotoxic drugs. Sirolimus can potentiate cyclosporine nephrotoxicity. Potassium-sparing diuretics can exacerbate hyperkalemia. Concurrent use of the following drugs frequently increases cyclosporine blood levels: corticosteroids, erythromycin and macrolide antibiotics, itraconazole, and ketoconazole. Other drugs that can increase cyclosporine blood levels are acetazolamide, alcohol, allopurinol, calcium-channel blockers, cimetidine, colchicine, oral contraceptives, fluconazole, imipenem/cilastatin, metoclopramide, norfloxacin, and sulindac. Enzyme-inducing drugs such as carbamazepine, phenobarbital, phenytoin and rifampin reduce cyclosporine blood levels. Octreotide can decrease cyclosporine oral absorption. Additional drugs that can reduce cyclosporine blood levels are cotrimoxazole, nafcillin, and sulfonamides. Cyclosporine reduces the clearance of HMG-CoA reductase inhibitors, such as lovastatin and atorvastatin, and increases the risk of drug-induced rhabdomyolysis.

**Parameters to Monitor.** Observe for anaphylaxis with IV administration. Monitor Cr, q 2–7 days and daily in patients at risk of acute renal dysfunction. Monitor blood pressure regularly. Monitor LFTs weekly, triglycerides and amylase monthly. Monitor blood or serum cyclosporine concentrations q 2–3 days when starting therapy. As the patient’s clinical condition and renal function allow, reduce frequency to once or twice weekly and then monthly during the first year of therapy. After the first year of therapy, blood or serum concentration monitoring may be reduced to q 1–2 months in stable patients. Monitor for signs and symptoms of graft rejection or GVHD, especially after dosage reduction.
Notes. Assignment of AB therapeutic equivalency by the FDA requires bioequivalence testing in normal healthy volunteers. Absorption of bioequivalent products can vary in bone marrow or solid organ transplant patients with compromised gut function. Bioequivalence to Neoral in transplant patients has been established for microemulsion Gengraf capsules and SangCya solution.

Cyclosporine is generally used in combination with other immunosuppressant drugs for prevention of graft rejection or GVHD. Cyclosporine is used in the management of various immunologic diseases such as aplastic anemia, psoriasis, atopic dermatitis, acute ocular Behcet’s syndrome, endogenous uveitis, primary biliary cirrhosis, and acute Crohn’s disease. High-dose cyclosporine is administered with certain chemotherapy regimens as a modulator of P-glycoprotein-mediated drug resistance. Optimmune (cyclosporine) ophthalmic (University of Georgia College of Veterinary Medicine) has orphan drug status for treatment of severe keratoconjunctivitis sicca with Sjögren’s syndrome. Sandimmune 2% ophthalmic ointment (Allergan) has orphan drug status for patients at high risk for graft rejection after penetrating keratoplasty and for treatment of corneal melting syndrome.

Comparative trials found greater nephrotoxicity and neurotoxicity with tacrolimus than with cyclosporine. Case reports describe resolution of certain drug-induced toxicities after replacement of cyclosporine with tacrolimus and resolution of cyclosporine-refractory GVHD with initiation of tacrolimus.

Pharmacology. Basiliximab and daclizumab (formerly dacliximab) are immunosuppressive, humanized, recombinant IgG1 monoclonal antibodies that bind specifically to the alpha subunit (p55 alpha, CD25, or Tac subunit) of the human high-affinity IL-2 receptor present on the surface of activated lymphocytes. They act as IL-2 receptor antagonists by preventing IL-2 from binding to lymphocytes, subsequently reducing IL-2–mediated immune activation. Both drugs are indicated for prevention of renal allograft rejection in a regimen that includes cyclosporine and a corticosteroid.

Administration and Adult Dosage. IV for prevention of renal allograft rejection (basiliximab) 20 mg infused over 30 min for 2 doses. The initial dose should be given about 2 hr before transplantation, the second dose is given 4 days after the transplant. Withhold the second dose if severe hypersensitivity or graft loss occurs. (Daclizumab) 1 mg/kg for 5 doses at 14-day intervals. The initial dose should be infused within 24 hr preceding surgery. The remaining 4 doses should be administered at 14-day intervals after surgery. Dilute each dose in 50 mL NS and infuse over 15 min through a peripheral or central venous line. IV for treatment of steroid-refractory GVHD (daclizumab) 1 mg/kg on days 1, 4, 8, 15, and 22, with day 1 representing the 1st day of daclizumab therapy; or 1.5 mg/kg, with repeated administration in 11–48 days for patients with transient improvement.
Special Populations. Pediatric Dosage. (Basiliximab) 12 mg/m² to a maximum of 20 mg/dose, given as adult dosage above. (Daclizumab) same as adult dosage. Information about use in pediatric patients is limited.

Geriatric Dosage. Same as adult dosage. Information about use in patients >65 yr is limited.

Other Conditions. No dosage adjustment is necessary for patients with severe renal dysfunction.

Dosage Forms. Inj (basiliximab) 20 mg; (daclizumab) 5 mg/mL.

Patient Instructions. This drug is being used as part of combination therapy to prevent rejection of your transplanted kidney.

Pharmacokinetics. Onset and Duration. (Basiliximab) receptor saturation is maintained for 36 ± 14 days with the recommended regimen. (Daclizumab) receptor saturation is maintained for about 120 days with the recommended regimen.

Serum Levels. (Basiliximab) >200 μg/L maintains complete binding to IL-2 receptor and maintains effective T-lymphocyte suppression. (Daclizumab) 5–10 mg/L inhibits activated T-lymphocytes.

Fate. (Basiliximab) peak serum levels of 9.3 ± 4.5 mg/L are attained with the recommended dosage regimen. (Daclizumab) after the first dose of 1 mg/kg, a peak of 21 ± 14 mg/L occurs, and after the 5th dose, a peak of 32 ± 22 mg/L results. The trough is 7.6 ± 4 mg/L after repeated doses of 1 mg/kg in adult renal transplant recipients.

t½. (Basiliximab) about 14 days; (daclizumab) 11–38 days.

Adverse Reactions. Both drugs usually are well tolerated. The frequency and type of adverse events were similar between renal transplant patients receiving these drugs or placebo, along with a corticosteroid and cyclosporine.257,258 Cases of severe acute hypersensitivity reactions have occurred with basiliximab, usually within 24 hr of a dose. Discontinue the drug permanently if this occurs. Hypertension and dehydration with daclizumab might be more frequent in children than in adults.

Precautions. Pregnancy; lactation. Use basiliximab with extreme caution in patients who have had previous courses of therapy. (See also General Precautions for Immunosuppressants.)

Drug Interactions. None known.

Parameters to Monitor. Monitor signs and symptoms of infection and graft rejection periodically.

Notes. After preparation, these drugs should be used within 4 hr if stored at room temperature or 24 hr if refrigerated. Although basiliximab and daclizumab have not been directly compared, their efficacies seem to be similar.
duction. Muromonab-CD3 binding to CD3 blocks allograft rejection by inhibition of T-lymphocyte function. Muromonab-CD3 is used to treat acute renal allograft rejection or steroid-resistant heart or liver allograft rejection.

**Administration and Adult Dosage.** **IV for the treatment of allograft rejection** 5 mg/day as an IV push for 10–14 days. Administration of methylprednisolone 8 mg/kg IV 1–4 hr before muromonab is strongly recommended by the manufacturer to reduce the frequency and severity of reactions with the first dose. Acetaminophen and diphenhydramine also are often used to control symptoms. The manufacturer recommends that the patient’s temperature be <37.8°C (<100°F) before infusion of muromonab-CD3.

**Special Populations.** **Pediatric Dosage.** Safety and efficacy are not established, but children have received dosages of ≤5 mg/day.

**Geriatric Dosage.** Same as adult dosage.

**Dosage Forms.** Inj 1 mg/mL.

**Patient Instructions.** This medication can cause shortness of breath, fever, and chills during the initial days of treatment.

**Pharmacokinetics.** **Serum Levels.** Levels ≥0.8 mg/L block cytotoxic T-lymphocyte function in vitro and in vivo.

**Adverse Reactions.** Cytokine release syndrome (CRS) occurs frequently with the initial 2–3 doses; it is related to cytokine release from activated lymphocytes. CRS can present as mild flu-like symptoms or as a life-threatening, shock-like reaction. Onset of CRS is usually 30–60 min after drug administration but can be delayed for hours. CRS might last for hours. Pretreatment and symptomatic treatment (see Administration and Adult Dosage) can reduce the frequency and severity of reactions with the first dose. Common symptoms are fever, headache, rigors, chills, tremor, nausea, vomiting, abdominal pain, myalgia, arthralgia, and rash. CRS can include CNS and cardiovascular adverse effects. CNS side effects are headache, seizures, encephalopathy, and aseptic meningitis. Cardiovascular side effects are angina, acute MI, CHF, hypertension, hypotension, and arrhythmias. Arterial and venous thromboses of allograft and other vascular beds have occurred. Consider coadministration of prophylactic antithrombotic agents in patients with histories of thrombotic events or underlying vascular disease. Pulmonary edema occurs frequently. Additional respiratory side effects are dyspnea, bronchospasm, wheezing, tachypnea, adult respiratory distress syndrome, and respiratory arrest. Hypersensitivity, including anaphylaxis and Stevens-Johnson syndrome, has been reported. Leukopenia, thrombocytopenia, pancytopenia, and lymphopenia also have occurred. Transient elevations of Cr, and serum transaminases can occur 1–3 days after initiation of treatment.

**Contraindications.** Human antimouse antibody titer ≥1:1000.

**Precautions.** Pregnancy; lactation. Use with caution in patients with volume overload or history of thrombotic events or vascular disease. (See also General Precautions for Immunosuppressants.)

**Drug Interactions.** Concurrent use of indomethacin has been associated with encephalopathy and other CNS side effects.
Parameters to Monitor. A chest x-ray obtained ≤24 hr before starting muromonab should be free of evidence of volume overload or heart failure. Obtain human antimouse antibody before initiating treatment. (See Contraindications.) Obtain Cr, q 2 days, AST and ALT q 3 days, CBC including differential and platelet counts q 3 days. Monitor one of the following immunologic tests during therapy: serum muromonab concentrations or quantitative T-lymphocyte surface phenotyping (target: CD3+ T-lymphocytes <25 cells/µL blood).

Notes. Transfer muromonab into a syringe through a 0.2 µ low protein-binding filter. Do not dilute with IV fluids for administration. Flush IV line with NS before and after injection.

**MYCOPHENOLATE MOFETIL**

Pharmacology. Mycophenolate mofetil is an ester prodrug of mycophenolic acid. Mycophenolic acid, which was isolated from the mold *Penicillium glaucum*, inhibits de novo purine synthesis by potent inhibition of inosine monophosphate dehydrogenase. Lymphocyte proliferation and antibody formation are subsequently inhibited by purine deficiency because lymphocytes lack an efficient salvage pathway for biosynthesis of purine bases. Mycophenolate is used in combination with cyclosporine and corticosteroids to prevent renal allograft rejection. Mycophenolate mofetil also has been used in combination with other immunosuppressants for the prevention of heart and liver allograft rejection.²⁵⁹

Administration and Adult Dosage. PO or IV for prophylaxis of kidney or liver transplant rejection 1 g bid beginning within 72 hr of transplantation.²⁶⁰,²⁶¹ PO or IV for prophylaxis of heart transplant rejection 1–1.5 g bid beginning within 72 hr of transplantation. PO or IV for treatment of acute or chronic GVHD after allogeneic bone marrow transplantation 1 g bid as adjunctive therapy for corticosteroid-refractory GVHD or to facilitate use of reduced corticosteroid dosage.²⁶²,²⁶³ A dosage of 3 g/day does not confer a therapeutic advantage for any condition and is associated with more adverse effects.

Special Populations. Pediatric Dosage. PO for renal transplantation 600 mg/m² bid, to a maximum of 2 g/day as the suspension. Alternatively, (1.25–1.5 m² BSA) 750 mg bid as capsules; (>1.5 m² BSA) 1 g bid as capsules or tablets.

Geriatric Dosage. Same as adult dosage.

Other Conditions. With a chronic Clcr <25 mL/min, the dosage should not exceed 1 g bid. This does not apply to the immediate post-transplant period for renal transplant patients.

Dosage Forms. Cap 250; Susp 200 mg/mL; Tab 500 mg; Inj 500 mg.

Patient Instructions. Do not stop this medication without consulting your physician.

Missed Doses. Take a missed dose as soon as possible if you remember within 12 hours. If it is within 2 hours of next dose, skip the missed dose and do not double the next dose. If you miss 2 or more doses, contact your physician.
Pharmacokinetics. Serum Levels. Not established; but one study found that dosage adjustment to a blood level of 2.5–4 mg/L decreased heart transplant rejection rate.264

Fate. Bioavailability is 94% in normal, healthy volunteers. Food decreases the peak serum concentration by 40%, but not bioavailability. Bioavailability is decreased immediately after renal transplantation. Peak serum concentrations after 1 g PO bid are 8.2 ± 4.5 mg/L during the first 40 days post-transplant and 24 ± 12 mg/L, 3 months post-transplant (similar to normal volunteers). The mean time to peak is prolonged to 1.3 ± 0.8 hr during the first 40 days post-transplant compared with 0.9 ± 0.2 hr after 3 months. AUC also is reduced by 42% during the first 40 days post-transplant; AUC is increased approximately 1.5-fold in patients with severe renal impairment. Alcoholic cirrhosis appears not to affect AUC. Mycophenolic acid is 97% bound to albumin. Vd in normal healthy volunteers is 4 ± 1.2 L/kg; Cl is 0.17 ± 0.04 L/hr/kg. Mycophenolate mofetil is rapidly hydrolyzed to mycophenolic acid, which is subsequently glucuronidated to an inactive metabolite. Enterohepatic recirculation can contribute to the mycophenolic acid AUC. Less than 1% of the dose is excreted in the urine as mycophenolic acid. t1/2 16.6 ± 5.8 hr.

Adverse Reactions. Hematologic adverse effects are leukopenia, anemia, thrombocytopenia, and pancytopenia. Adverse effects of mycophenolate rarely necessitate discontinuation of therapy, but the drug should be stopped temporarily if neutropenia (ANC <1300/µL) develops during therapy. GI effects, including nausea, vomiting, dyspepsia, abdominal pain, constipation, and diarrhea, occur frequently. GI side effects can be reduced by giving the drug in 3–4 divided doses.

Contraindications. Allergy to mycophenolate mofetil.

Precautions. Pregnancy; lactation. Use with caution in patients with renal dysfunction. (See also General Precautions for Immunosuppressants.)

Drug Interactions. Concurrent iron or aluminum- or magnesium-containing antacids reduce absorption. Cholestyramine reduces the serum concentration of mycophenolate mofetil. In vitro, salicylate increases the unbound fraction of mycophenolic acid.

Parameters to Monitor. Monitor CBC, including differential and platelet counts, weekly during the first 1–2 months of therapy, q 2 weeks during the 2–4 months of therapy, and monthly thereafter. Monitor for signs and symptoms of infection, graft rejection, and GVHD.

Notes. Mycophenolate mofetil has been used in the treatment of certain dermatologic and immunologic disorders such as atopic dermatitis, inflammatory bowel disease, lupus nephritis, myasthenia gravis, pemphigus, psoriasis, rheumatoid arthritis, Takayasu’s arteritis, uveitis, and Wegener’s granulomatosis.

Pharmacology. Sirolimus is a macrocyclic lactone immunosuppressant isolated from Streptomyces hygroscopicus that is structurally related to tacrolimus. It binds FK binding protein-12 (FKBP-12) and inhibits the cytosolic enzyme target of ra-
pamycin (TOR). Inhibition of TOR restricts differentiation and proliferation of T-lymphocytes and B-lymphocytes subsequent to cytokine stimulation.

**Administration and Adult Dosage.** *PO for prevention of renal allograft rejection* (≥40 kg) 6 mg on first day of therapy, followed by 2 mg daily. Dilute in glass or plastic container with at least 60 mL of water or orange juice. Mix thoroughly and administer immediately. Then fill container with at least 120 mL of liquid, stir vigorously, and administer immediately. Administer with or without food and consistently with respect to meals, oral cyclosporine, and substrates of CYP3A4 or P-glycoprotein.

**Special Populations. Pediatric Dosage.** *PO for prevention of renal allograft rejection* 3 mg/m² on first day of therapy, followed by 1 mg/m² daily.

**Geriatric Dosage.** Same as adult dosage.

**Other Conditions.** For adults <40 kg, same as pediatric dosage. In hepatic failure, reduce maintenance dosage by approximately one-third. In renal dysfunction, no dosage adjustment is necessary.

**Dosage Forms.** *Soll* 1 mg/mL; *Tab* 1 mg.

**Patient Instructions.** Take this medication on a regular schedule relative to the time of day and meals. Do not discontinue it unless directed to do so. If your sirolimus is in a bottle, use the amber syringe provided by the manufacturer to measure and take each dose out of the container. If your sirolimus is in a packet, squeeze contents to the lower part of the pouch and cut it across the top. Dilute sirolimus in a glass or plastic container with at least 2 fluid ounces of water or orange juice. Mix thoroughly and swallow immediately. Then fill container with at least 4 fluid ounces of water or orange juice, stir vigorously, and swallow immediately. Take each dose with or without food but consistently with respect to meals and medications. Refrigerate. Discard bottle 1 month after opening.

**Missed Doses.** Take a missed dose as soon as possible if you remember within 16 hours. If it is within 8 hours of the next dose, skip the missed dose and do not double the missed dose. If you miss 2 or more doses, contact the physician or coordinator who monitors your sirolimus.

**Pharmacokinetics. Serum Levels.** Relationship between whole blood levels and therapeutic or toxic effects is not well defined. Whole blood levels are not monitored routinely, although they may be monitored in pediatric patients or patients with markedly impaired hepatic function. Approximate whole blood trough levels (immunoassay) are 9 μg/L and 17 μg/L in patients receiving sirolimus 2 mg/day and 5 mg/day, respectively. The 24 hr post-dose whole blood concentration correlates with AUC. Marked interpatient variability of whole blood levels occurs.

**Fate.** Oral bioavailability is 14%. Time-to-peak whole blood concentration is 1–2 hr; it is delayed and AUC is increased by 35% when sirolimus is taken after a high-fat meal. P-glycoprotein–mediated countertransport affects absorption. The drug is extensively protein bound in plasma, primarily to albumin, α1-acid glycoprotein, and lipoproteins. There is extensive sequestration in erythrocytes, with a whole blood:plasma ratio of 36:1. Vₐss is 12 ± 7.5 L/kg. Sirolimus is a CYP3A4
substrate. After administration of radiolabeled drug, 2% is recovered in the urine and 91% is recovered in the bile.

\[ t_{1/2} = 62 \pm 16 \text{ hr}. \]

**Adverse Reactions.** Phase I studies of sirolimus included concurrent administration of other immunosuppressants, including corticosteroids, and cyclosporine or tacrolimus. Subsequently, many reported side effects may not be directly attributable to sirolimus. Adverse effects related to use of sirolimus include hypercholesterolemia, hypertriglyceridemia, hypertension, anemia, thrombocytopenia, leukopenia, diarrhea, hypokalemia, arthralgia, rash, and acne. Thrombocytopenia and lipid abnormalities are dose related. Thrombocytopenia generally resolves after drug discontinuation. Additional adverse effects reported in patients taking sirolimus in combination with other immunosuppressants are nausea, emesis, dyspepsia, abdominal pain, diarrhea, constipation; renal and metabolic abnormalities such as increased Cr, hypophosphatemia, hyperkalemia, peripheral edema, and weight gain; respiratory system effects are dyspnea, pharyngitis and upper respiratory tract infection. Fever, headache, asthenia, body pain, arthralgia, insomnia, tremor, and posttransplant lymphoproliferative disorder also have been reported.

**Contraindications.** Hypersensitivity to sirolimus, derivatives of sirolimus, or any component of the formulation.

**Precautions.** Pregnancy; lactation. (See also General Precautions for Immunosuppressants.)

**Drug Interactions.** Concurrent administration of oral cyclosporine microemulsion capsules (Neoral) increases AUC, peak and trough, but administration of oral cyclosporine 4 hr after sirolimus has no effect on sirolimus whole blood levels. Sirolimus can potentiate cyclosporine nephrotoxicity. Diltiazem and ketoconazole increase sirolimus levels. Rifampin decreases sirolimus levels. AUC is unchanged with concurrent administration of acyclovir, glyburide, digoxin, nifedipine, norgestrel, or ethinyl estradiol. AUC can be affected by substrates, inhibitors, or inducers of CYP3A4 or P-glycoprotein.

**Parameters to Monitor.** Monitor WBC, erythrocyte, and platelet counts weekly during the first 2–3 months of therapy and monthly thereafter in stable patients. Monitor serum lipids monthly. Monitor for signs and symptoms of graft rejection, especially after dosage reduction.

**Notes.** Sirolimus has been used in the treatment of psoriasis.

**TACROLIMUS**

**Pharmacology.** Tacrolimus (formerly FK506) is a macrolide antibiotic produced by *Streptomyces tsukubaensis*. The intracellular drug–ligand complex of tacrolimus and FKBP-12 indirectly blocks T-lymphocyte activation. It inhibits calcineurin-mediated dephosphorylation of factors necessary for IL-2 transcription.

**Administration and Adult Dosage.** PO for prophylaxis of organ rejection or GVHD 0.15–0.3 mg/kg initially, depending on the type of transplant and concomitant administration of other immunosuppressants. Tacrolimus is usually started 4–12 hr
before surgery and administered on a bid schedule. **Maintenance dosage** is based on tacrolimus blood concentrations, the magnitude of risk for organ rejection or GVHD, and patient tolerance. **IV for patients unable to take medication orally** 0.03–0.1 mg/kg/day as a continuous infusion diluted in D5W or NS in a glass container (it can leach plasticizers from PVC containers) to a concentration of 2 mg/100–500 mL (final concentration 4–20 mg/L).265 Mucositis generally necessitates initial use of IV tacrolimus for allogeneic bone marrow transplant recipients. Tacrolimus is generally started 1 or 2 days before bone marrow transplantation. **Conversion from cyclosporine** the manufacturer recommends at least 24 hr between the last cyclosporine dose and the first tacrolimus dose. **To convert from IV to PO tacrolimus** the IV:PO dosage ratio is typically 1:3. A tacrolimus blood level should be drawn 48 hr after dosage form conversion. Absorption of oral medications may be reduced in allogeneic bone marrow transplant patients due to residual gut damage from intensive chemotherapy or radiation or GVHD. **Discontinuation** tacrolimus may be eventually discontinued in certain renal or allogeneic bone marrow transplant recipients. However, it must be decreased gradually to reduce the risk of reactive immune stimulation and consequent graft rejection or GVHD.

**Special Populations. Pediatric Dosage.** Same as adult dosage.

**Geriatric Dosage.** Same as adult dosage. Age-related reduction in renal function and comorbid conditions may predispose elderly patients to tacrolimus-induced nephrotoxicity and hypertension.

**Special Populations.** Use IBW to calculate initial dosage in obese patients.

**Dosage Forms.** **Cap** 0.5, 1, 5 mg; **Soln** 100 mg/mL; **Inj** 5 mg/mL.

**Patient Instructions.** Take this medication on a regular schedule in relation to the time of day and meals. Do not discontinue it unless directed to do so. Grapefruit juice can interact with cyclosporine. Talk to your physician before drinking grapefruit juice and before starting, stopping, or changing the dose of any medication.

**Missed Doses.** Take a missed dose as soon as possible if you remember within 12 hours. If it is within 2 hours of next dose, skip the missed dose and do not double the next dose. If you miss 2 or more doses, contact your physician.

**Pharmacokinetics. Serum Levels.** The serum (blood) concentration–response relationship is not completely defined. Trough blood or serum concentrations are monitored to reduce the risk of toxicity. Therapeutic and toxic concentrations vary with assay, biologic fluid, and time post-transplant. Artifically elevated serum concentrations can occur when blood is drawn through the same central venous line used for IV tacrolimus administration. Whole blood trough levels of 10–20 μg/L are often considered therapeutic.

**Fate.** The mean absorption of tacrolimus capsules in normal healthy volunteers is 17 ± 7%. In liver transplant patients absorption is 22 ± 6%. Factors that can decrease absorption are diarrhea, gastroenteritis, and short small bowel. Food does not affect bioavailability but decreases and delays peak serum levels. Whole blood peak concentrations in liver transplant patients after 0.15 mg/kg is 52.4 μg/L fasting and 27.5 μg/L with food. Tacrolimus is extensively bound to erythrocytes and plasma proteins, primarily albumin and α1-acid glycoprotein. $V_d$ (whole
blood) is 0.85 ± 0.3 L/kg in liver transplant patients; Cl (whole blood) is 0.053 ± 0.017 L/hr/kg in liver transplant recipients. Tacrolimus is extensively metabolized by CYP3A. At least 10 metabolites, some with immunosuppressant activity, have been identified. Less than 1% is excreted unchanged in the urine.

\( t_{1/2} \) (Whole blood) 21.2 ± 8.5 hr in normal healthy volunteers; 11.7 ± 3.9 hr in liver transplant patients.

**Adverse Reactions.** Acute nephrotoxicity, which usually occurs within 1 month post-transplant and is characterized by Crs increasing ≥0.3 mg/mL/24 hr, frequently abates with interruption of drug therapy. Dosage reduction might be required for continued administration. Chronic progressive renal toxicity is characterized by a slow, continual increase in Crs and BUN, mild proteinuria, and tubular dysfunction. Cr >2 mg/dL in adult patients, or doubling of Crs, is an indication for interruption of therapy or dosage reduction. Electrolyte abnormalities, including hypomagnesemia, hypokalemia or hyperkalemia, and renal tubular acidosis, are consequences of tacrolimus-induced nephrotoxicity. Concurrent administration of nephrotoxic drugs increases the likelihood of renal dysfunction. Hypertension occurs frequently. Calcium-channel blockers and clonidine are suitable agents for the management of tacrolimus-induced hypertension because these agents do not have deleterious effects on renal blood flow. Fine tremors occur frequently and can persist and worsen after drug discontinuation. Neurotoxicity symptoms are headache, seizures, encephalopathy, confusion, insomnia, cortical blindness, expressive aphasia, paresthesia, hyperesthesia, and myoclonic reactions. Anaphylactoid reactions to tacrolimus or the solubilizing agent, polyoxyethylated castor oil, can occur. After an allergic reaction to IV tacrolimus, patients may receive a trial of oral tacrolimus capsules under close observation. Hyperbilirubinemia, increased \( \gamma \)-glutamyltranspeptidase, serum alkaline phosphatase, and serum transaminases occur frequently. Additional adverse effects are photophobia, rash, hirsutism, pleural effusion, gingival hyperplasia, diarrhea, nausea, vomiting, and hypertriglyceridemia.

**Contraindications.** Allergy to tacrolimus or polyoxyethylated castor oil.

**Precautions.** Pregnancy; lactation. Use with caution in patients at risk for renal dysfunction. (See also General Precautions for Immunosuppressants.)

**Drug Interactions.** Additive or synergistic renal toxicity can occur with concurrent administration of nephrotoxic drugs. Potassium-sparing diuretics can exacerbate hyperkalemia. Because tacrolimus is metabolized by CYP3A, numerous drug interactions are possible with concurrent administration of drugs that affect this enzyme system. The following drugs can increase tacrolimus blood levels: corticosteroids, itraconazole, ketoconazole, erythromycin and other macrolide antibiotics, oral contraceptives, fluconazole, calcium-channel blockers, cimetidine, danazol, and metoclopramide. Enzyme-inducing drugs such as carbamazepine, phenobarbital, phenytoin, rifabutin, and rifampin can decrease tacrolimus blood levels. Tacrolimus can decrease the clearance of HMG-CoA reductase inhibitors and increase the risk of drug-induced rhabdomyolysis.

**Parameters to Monitor.** Observe for anaphylaxis with IV administration. Monitor Cr, q 2–7 days and daily in patients at risk for acute renal dysfunction. Monitor
LFTs weekly, triglycerides monthly, and blood pressure regularly. Monitor blood or serum concentrations q 2–3 days when starting therapy. As the patient’s clinical condition and renal function allow, reduce frequency to 1–2 times weekly and then monthly during the first year of therapy. After the first year of therapy, blood or serum concentration monitoring may be reduced q 1–2 months in stable patients. Monitor for signs and symptoms of graft rejection or GVHD, especially after dosage reduction.

Notes. Comparative clinical trials found greater nephrotoxicity and neurotoxicity with tacrolimus than with cyclosporine. Case reports describe resolution of drug-induced toxicity after replacement of cyclosporine with tacrolimus and resolution of cyclosporine-refractory acute GVHD or chronic GVHD with tacrolimus.

REFERENCES


296 ANTI NEOPLASTICS
**Antiarrhythmic Drugs**

**ADENOSINE**

**Pharmacology.** Adenosine is a purinergic agonist that acts on the purine P<sub>1</sub> and P<sub>2</sub> receptors (although P<sub>1</sub> receptors are more sensitive to adenosine). Pharmacologic effects include coronary and peripheral vasodilation, negative inotropic actions, and depression of sinus node and AV nodal conduction. It is used most frequently for supraventricular tachycardia caused by re-entry (ie, AV nodal re-entry or AV re-entry associated with an extranodal pathway). In these instances, restoration of sinus rhythm occurs in 85–95% of patients. The drug also can be helpful in diagnosing wide-QRS tachycardias believed to be supraventricular in origin.<sup>1–3</sup>

**Adult Dosage.** IV for supraventricular tachycardia administer over 1–2 sec through an IV line with minimal dead space, followed by a saline flush; initial dose is 6 mg (3 mg if administered through a central line); if this is ineffective, 12 mg can be given 2 min later and repeated if necessary. An average effective dose of 1 mg has been reported in patients receiving concurrent dipyridamole.

**Pediatric Dosage.** IV 0.1–0.2 mg/kg increased in increments of 0.05 mg/kg q 2 min prn, to a maximum of 0.25 mg/kg.<sup>4</sup>

**Dosage Forms.** Inj 3 mg/mL.

**Pharmacokinetics.** Adenosine is rapidly metabolized in blood to inactive adenosine monophosphate and inosine; elimination half-life is about 1–10 sec.

**Adverse Reactions.** Frequent, but short-lived, subjective complaints include chest discomfort, dyspnea, flushing, and headache. Postconversion arrhythmias also are frequent but transient and include ventricular ectopy, sinus bradycardia, AV block, atrial fibrillation, and rapid reinitiation of supraventricular tachycardia. Adenosine is contraindicated in patients with pre-existing sinus node dysfunction or second- or third-degree heart block without a functioning pacemaker because of the risk of prolonged sinus arrest or AV block. Also use adenosine with caution in asthmatics because it can precipitate bronchospasm, and in patients with atrial fibrillation with an accessory AV pathway because it can accelerate ventricular response.

**Drug Interactions.** Dipyridamole blocks the cellular uptake of adenosine, enhancing the pharmacologic effect; theophylline, a purine antagonist, inhibits the therapeutic actions of adenosine.
Pharmacology. Amiodarone is a type III antiarrhythmic that prolongs the effective refractory period of atrial and ventricular tissue by blocking potassium conductance. It decreases sinus rate and slows conduction through the AV node by β-adrenergic blockade. Amiodarone also blocks sodium and calcium channels. The antiarrhythmic actions can be caused by interruption of re-entrant substrate or abolition of premature beats that trigger re-entry.

Administration and Adult Dosage. PO loading dosage 800–1600 mg/day in divided doses for 1–2 weeks. Loading dosages are usually toward the lower end of this range for atrial arrhythmias and toward the upper end of the range for ventricular arrhythmias. PO maintenance dosage 100–600 mg/day (usually 300–400 mg/day for recurrent ventricular tachycardia and 100–200 mg/day for supraventricular tachycardias such as atrial fibrillation). Some suggest a 600–800 mg/day priming dosage for 1–2 months after the initial loading period and before maintenance therapy. IV for treatment or prevention of refractory ventricular tachycardia or fibrillation 150 mg over 10 min 360 mg over the next 6 hr, and 540 mg over the next 18 hr. In one study, amiodarone was administered as a 300 mg IV bolus for cardiac arrest. Initiate amiodarone only during hospitalization for the first several days of the loading phase.

Special Populations. Pediatric Dosage. Safety and efficacy not established. PO 10–15 mg/kg/day for 10 days and then 5 mg/kg/day maintenance therapy has been used. IV 5 mg/kg in 1 mg/kg increments over 5–10 min each; an additional 1 to 5 mg/kg may be given in 30 min if needed.

Geriatric Dosage. Same as adult dosage.

Dosage Forms. Tab 200 mg; Inj 50 mg/mL.

Patient Instructions. Report any shortness of breath, tiredness, abdominal discomfort, or visual abnormalities. Avoid intense sunlight; use sunscreen. Divided doses during loading or maintenance dosage phases can reduce intestinal upset.

Missed Doses. Take this drug at regular intervals. If you miss a dose, do not take it. If it is about time for the next dose, take that dose only. Do not double the dose or take extra.

Pharmacokinetics. Onset and Duration. Onset is variable, from several days to a month; full effect might not occur for several months.

Serum Levels. 1–2.5 mg/L (1.6–4 μmol/L) proposed but not well established. Desethylamiodarone accumulates to serum levels similar to or greater than the parent drug.

Fate. Oral absorption is erratic and incomplete; bioavailability is 46 ± 22%. Peak serum concentrations occur in 3–7 hr. The drug is 99.9% plasma protein bound. Vd is 66 ± 44 L/kg; Cl is 0.11 ± 0.024 L/hr/kg. Amiodarone is primarily hepatically eliminated with at least one active metabolite, desethylamiodarone. No unchanged amiodarone or desethylamiodarone is found in urine. t1/2, α phase 4–12 hr; β phase changes with duration of therapy and study sampling. Reported variously as 25 ± 12 days and 53 ± 23 days. Similar for desethylamiodarone.
**Adverse Reactions.** Corneal microdeposits occur in virtually all patients and are no reason for stopping treatment; however, visual disturbances are reported in about 5%.\textsuperscript{11} Neurologic effects occur frequently and include tremor, ataxia, paresthesias, and nightmares, which can be more common during the loading phase.\textsuperscript{11} Anorexia, nausea, vomiting, and/or constipation occur frequently. Transient elevations in hepatic enzymes occur in more than 50% of patients, but clinical hepatitis occurs only occasionally.\textsuperscript{12} Photosensitivity occurs frequently, and a blue-gray skin pigmentation (sometimes irreversible) develops in 2–4% of patients.\textsuperscript{11} Hypothyroidism (low-T\textsubscript{3} syndrome) or hyperthyroidism occurs frequently.\textsuperscript{13} Occasional proximal muscle weakness and myopathy have been reported. Symptomatic pulmonary fibrosis has been reported in 1–6% of patients; it is probably not immunologic in etiology and seems to occur more often in patients with underlying lung disease.\textsuperscript{11,14} Pulmonary symptoms usually improve with drug discontinuation, but up to 10% of cases result in death.\textsuperscript{11,14} Aggravation of ventricular tachycardia and drug-induced torsades de pointes can occur.\textsuperscript{11,15} Occasional severe sinus bradycardia (requiring a pacemaker) or AV block has been reported.

**Contraindications.** Sick sinus syndrome or second- or third-degree heart block in the absence of a ventricular pacemaker; patients in whom bradycardia has caused syncope; long-QT syndrome.

**Precautions.** Electrophysiologic studies may not predict the long-term efficacy of amiodarone.\textsuperscript{16} The benzyl alcohol preservative can be hazardous in infants.

**Drug Interactions.** Amiodarone inhibits a wide array of cytochrome P450 enzymes including CYP1A2, 2C9, 2D6, and 3A4; it also inhibits p-glycoprotein.\textsuperscript{17} Amiodarone increases serum levels of cyclosporine, digoxin, flecainide, phenytoin, procainamide, and quinidine. It potentiates the anticoagulant effects of warfarin; reduce the initial dosage of warfarin by one-third to one-half.

**Parameters to Monitor.** Monitor ECG daily during loading phase for heart rate, PR, QRS, and QT duration. Baseline and periodic thyroid function tests and liver enzymes (especially if symptoms present). Obtain baseline pulmonary function tests; repeat chest x-ray and clinical examination q 3–6 months.\textsuperscript{11,14}

**Notes.** Because of the results of the Cardiac Arrhythmia Suppression Trial (CAST),\textsuperscript{18} many clinicians use type III antiarrhythmics (eg, amiodarone, sotalol) as first-line therapy for supraventricular and ventricular arrhythmias. A noniodinated analogue of amiodarone under clinical investigation is dronedarone.

**BRETYLIUM TOSYLA TE**

**Pharmacology.** Bretylium is a type III antiarrhythmic with actions thought to be caused by an initial catecholamine release and subsequent catecholamine depletion and/or direct effect independent of the adrenergic nervous system. Direct actions can be mediated by blockade of potassium channels. Bretylium causes an initial increase in blood pressure, heart rate, and myocardial contractility (from catecholamine release), followed by hypotension (from neuronal blockade). Its greatest usefulness is in severe ventricular tachyarrhythmias resistant to other antiarrhythmics. Bretylium can be effective for ventricular fibrillation but is usually ineffective against ventricular tachycardia.
Administration and Adult Dosage. IV loading dose 5 mg/kg push with an additional dose of 10 mg/kg if no response. Maintenance dosage IM or IV (over 8 min or more) 5–10 mg/kg q 6 hr or as an IV infusion of 1–2 mg/min.

Special Populations. Pediatric Dosage. Not well established, although the following has been suggested: IV loading dosage for ventricular fibrillation 5 mg/kg, followed by 10 mg/kg at 15- to 30-min intervals, to a maximum total dosage of 30 mg/kg; IV maintenance dosage 5 mg/kg q 6–8 hr.  

Geriatric Dosage. Same as adult dosage.

Other Conditions. In renal impairment, lower dosages might be required. A nomogram for dosage in renal insufficiency has been described.

Dosage Forms. Inj 50 mg/mL.

Pharmacokinetics. Onset and Duration. IV onset usually 5–10 min but can be delayed to 20–60 min; myocardial levels increase gradually over 6–12 hr. Duration is usually 6–12 hr after a single dose. Because of persistent myocardial levels, duration after multiple doses can be much longer.

Fate. 23 ± 9% is orally absorbed. The drug is not bound to plasma proteins. Vd is 5.9 ± 0.8 L/kg; Cl is 0.61 ± 0.11 L/hr/kg. After IV administration, bretylium is primarily cleared renally, with 77 ± 15% excreted in the urine unchanged. Disposition is probably route and concentration dependent.

t¹⁄₂. α phase about 25 min; β phase 8.9 ± 1.8 hr, mean of 33.4 hr in renal insufficiency.

Adverse Reactions. Hypotension (usually orthostatic) via adrenergic blockade occurs in up to 50% of patients. The drop in mean arterial pressure is usually not more than 20 mm Hg, but the drop can be severe, necessitating drug discontinuation. Nausea and vomiting occur frequently after rapid IV administration.

Contraindications. Suspected digitalis-induced ventricular tachycardia (can increase the rate of ventricular tachycardia or the likelihood of ventricular fibrillation).

Precautions. Use with caution if hypotension exists before administration. Keep patient supine until tolerance to hypotension develops. Prolonged effects can occur, and dosage reduction in patients with impaired renal function may be required.

Drug Interactions. Bretylium enhances pressor effects of catecholamines.

Parameters to Monitor. Closely monitor blood pressure and constantly monitor ECG.

DIGOXIN Lanoxin, Various

Pharmacology. Digitalis glycosides exert positive inotropic effects through improved availability of calcium to myocardial contractile elements, thereby increasing cardiac output in CHF. In CHF, digoxin improves the symptoms of CHF but does not alter long-term mortality. Antiarrhythmic actions of digoxin are caused primarily by an increase in AV nodal refractory period via increased vagal tone.
sympathetic withdrawal, and direct mechanisms. Digoxin also exerts a moderate, direct vasoconstrictor action on arterial venous smooth muscle.

**Administration and Adult Dosage.** IV **loading dosage** 10–15 µg/kg in divided doses over 12–24 hr at intervals of 6–8 hr.²⁶ **PO loading dosage** adjust dosage for percent oral absorption. (See Fate.) Usually, 0.5–0.75 mg is given and then 0.125–0.375 mg q 6–8 hr until the desired effect or total digitalizing dosage is achieved. **Maintenance dosage** = (total body stores) × (% lost/day), where total body stores is the original calculated loading dosage and % lost/day is 14 + (Clcr/5). Usual maintenance dosage ranges from 0.125–0.5 mg/day.²⁶ A dosage nomogram has also been described.²⁷ IM not recommended.

**Special Populations. Pediatric Dosage.** Base all dosages on ideal body weight. **Total digitalizing dosage (TDD) PO** (premature newborn) 20 µg/kg; (full-term newborn) 30 µg/kg; (1–24 months) 40–50 µg/kg; (2–10 yr) 30–40 µg/kg; (>10 yr) 10–15 µg/kg. Give ½ TDD initially and then ¼ TDD q 8–18 hr twice. **PO maintenance dosage** (premature newborn) 5 µg/kg/day; (full-term newborn) 8–10 µg/kg/day; (1–24 months) 10–12 µg/kg/day; (2–10 yr) 8–10 µg/kg/day; (>10 yr) 2.5–5 µg/kg/day. In children <10 yr, give in 2 divided doses per day. **IV** (all ages) 75% of PO dosage.⁴

**Geriatric Dosage.** Maintenance dosage can be lower because of age-related decrease in renal function.²⁸

**Other Conditions.** Decrease loading and maintenance dosages with renal impairment. Base dosage on ideal body weight in obese individuals.

**Dosage Forms.** Cap 0.05, 0.1, 0.2 mg; Elxr 50 µg/mL; Tab 0.125, 0.25 mg; Inj 0.1, 0.25 mg/mL.

**Patient Instructions.** Report feelings of tiredness, appetite loss, nausea, abdominal discomfort, or visual disturbances such as hazy vision, light sensitivity, spots, halos, or red–green blindness.

**Missed Doses.** Take this drug at regular intervals. If you miss a dose and it has been less than 12 hours since your dose was due, take it as soon as you remember. If it is about time for the next dose, take that dose only. Do not double the dose or take extra.

**Pharmacokinetics. Onset and Duration.** IV onset 14–30 min; peak 1.5–5 hr; somewhat slower after oral administration.

**Serum Levels.** Therapeutic 0.5–2 µg/L (0.6–2.5 nmol/L); toxic >3 µg/L (3.8 nmol/L). Considerable overlap exists between therapeutic and toxic ranges.²⁹ Signs or symptoms of toxicity can be evident below 3 µg/L, especially if other risk factors are present.²⁹ In CHF there does not seem to be an advantage in maintaining the digoxin level above 1 µg/L.³⁰ Obtain blood samples for digoxin levels at least 4 hr after an IV dose and 6–8 hr after an oral dose to allow central and tissue compartment equilibration. Digoxin concentrations (digitalis-like immunoreactive substance) have been detected in patients with renal failure, neonates, pregnant women, and those with severe liver disease not receiving a digitalis glycoside.³¹
Fate. Oral absorption is 70 ± 13% from tablets; 85% from elixir; 95% from capsules. Enterohepatic recycling of digoxin can be as high as 30%. Protein binding to albumin is 25 ± 5%; V_d is 7–8 L/kg; Cl is 0.16 ± 0.036 L/hr/kg; both depend on renal function. The drug is excreted 60 ± 11% unchanged in the urine in patients with normal renal function. Active metabolites include digitoxigenin, bisdigitoxoside, digoxigenin monodigitoxoside, and dihydriodigoxin. 

\[ t_{1/2} \phi_1 = 0.5–1 \text{ hr}; \quad t_{1/2} \phi_2 = 39 ± 13 \text{ hr}; \quad t_{1/2} \phi_3 = 3.5–4.5 \text{ days} \] in anephric patients.

Adverse Reactions. Arrhythmias, listed by decreasing prevalence, are premature ventricular beats, second- and third-degree heart blocks, AV junctional tachycardia, atrial tachycardia with block, ventricular tachycardia, and SA nodal block. Visual disturbances are related to serum level and occur in up to 25% of patients with digoxin intoxication. They include blurred vision, yellow or green tinting, flickering lights or halos, or red–green color blindness. GI symptoms occur frequently and include abdominal discomfort, anorexia, nausea, and vomiting. CNS side effects occur frequently but are nonspecific, such as weakness, lethargy, disorientation, agitation, and nervousness. Hallucinations and psychosis have been reported. Rare reactions include gynecomastia, hypersensitivity, and thrombocytopenia.

Contraindications. Hypertrophic obstructive cardiomyopathy; suspected digitalis intoxication; second- or third-degree heart block in the absence of mechanical pacing; atrial fibrillation with accessory AV pathway; ventricular fibrillation.

Precautions. Electrolyte abnormalities predisposing to digoxin toxicity include hypokalemia, hypomagnesemia, and hypercalcemia. Hypothyroidism can reduce digoxin requirements because of lower V_d and clearance. Direct current cardioversion carries little risk in the absence of digoxin toxicity. Use with caution in patients with pulmonary disease because hypoxia can sensitize the myocardium to arrhythmias and increase the risk of toxicity. Serious bradyarrhythmias can occur with sick sinus syndrome, but controversy exists concerning the clinical importance of its effects on the SA node. Digoxin can increase infarct size in the nonfailing heart.

Drug Interactions. β-Blockers can worsen CHF or digoxin-induced bradycardia. Potassium loss caused by amphotericin B or diuretics can contribute to digoxin toxicity. Spironolactone can decrease digoxin renal elimination. ACE inhibitors, amiodarone, bepridil, diltiazem, nitrendipine, quinidine, and verapamil can increase digoxin levels. Oral antacids, kaolin-pectin, oral neomycin, and sulfasalazine can reduce digoxin absorption. Penicillamine can decrease serum digoxin levels.

Parameters to Monitor. Obtain serum levels only when compliance, effectiveness, or systemic availability is questioned or toxicity is suspected. (See Serum Levels.) Monitor heart rate, ECG for digoxin-induced arrhythmias, subjective complaints of toxicity, and renal function. Monitor serum electrolytes (especially potassium) frequently initially and then q 1–2 months when stabilized.

Notes. Treatment of severe or life-threatening digoxin toxicity should include IV digoxin immune Fab (Digibind). About 40 mg (one vial) of digoxin-specific Fab
fragments binds 0.6 mg of the glycoside. Exact dosage can be calculated based on estimated total body stores.

**DISOPYRAMIDE PHOSPHATE**

**Pharmacology.** Disopyramide has qualitatively the same electrophysiologic actions as procainamide and quinidine and is effective for ventricular and (unlabeled) supraventricular tachycardia. It increases systemic vascular resistance through vasoconstriction; it also can exert a profound negative inotropic effect and has marked anticholinergic properties systemically and on the heart. The isomers of disopyramide have stereospecific pharmacologic actions.38,39

**Administration and Adult Dosage.** PO loading dosage 300–400 mg. PO maintenance dosage 400–800 mg/day, to a maximum of 1.6 g/day. Give daily dosage in 4 equally divided doses q 6 hr with non-SR Cap or in 2 equally divided doses q 12 hr with SR Cap. Initiate disopyramide during hospitalization.

**Special Populations. Pediatric Dosage.** PO <1 yr, 10–30 mg/kg/day; 1–4 yr, 10–20 mg/kg/day; 4–12 yr, 10–15 mg/kg/day; 12–18 yr, 6–15 mg/kg/day. Daily dosage is divided into 4 equal doses q 6 hr.4 (See Notes.)

**Geriatric Dosage.** Decreased dosage is probably necessary because the elderly might not tolerate anticholinergic side effects.

**Other Conditions.** In patients who weigh less than 50 kg or have hepatic disease or moderate renal insufficiency (Clcr >40 mL/min), load with 150–200 mg and then give 400 mg/day in 2 or 4 divided doses, depending on the dosage form used. Initial daily dosage in patients with hepatic disease is about 4.4 mg/kg/day.40,41 In patients with severe renal insufficiency, give maintenance dosages as follows (non-SR Cap):

<table>
<thead>
<tr>
<th>CREATinine CLEARANCE</th>
<th>DAILY MAINTENANCE DOSAGE</th>
</tr>
</thead>
<tbody>
<tr>
<td>30–40 mL/min</td>
<td>300 mg</td>
</tr>
<tr>
<td>15–30 mL/min</td>
<td>200 mg</td>
</tr>
<tr>
<td>&lt;15 mL/min</td>
<td>100 mg</td>
</tr>
</tbody>
</table>

**Dosage Forms.** Cap 100, 150 mg; SR Cap 100, 150 mg. (See Notes.)

**Patient Instructions.** Report any symptoms such as difficulty in urination, constipation, blurred vision, or dry mouth. Also report shortness of breath, weight gain, or edema. Do not crush or chew sustained-release capsules. A sustained-release capsule core in the stool does not indicate lack of absorption.

**Missed Doses.** Take this drug at regular intervals. If you miss a dose, take it as soon as you remember. If it is about time for the next dose, take that dose only. Leave at least 4 hours between regular capsule doses and 6–8 hours between sustained-release capsule doses. Do not double the dose or take extra.

**Pharmacokinetics.** Onset and Duration. PO onset is within 1 hr. Duration differs with individual differences in drug disposition but is usually 6–12 hr.
Serum Levels. Usual range is 2–5 mg/L (6–15 μmol/L),\textsuperscript{41,42} with toxicity more likely over 4 mg/L. Therapeutic range of unbound drug is 0.5–2 mg/L (1.5–6 μmol/L).\textsuperscript{40} Monitoring unbound concentrations eliminates variability caused by concentration-dependent disposition.\textsuperscript{41,43}

Fate. Oral absorption is rapid; systemic availability is 83 ± 11%.\textsuperscript{41,42} Unbound drug in serum varies from 19–46% over a serum concentration range of 2–8 mg/L and is also age dependent.\textsuperscript{44} \( V_d \) (unbound) is 1.4–1.7 L/kg in normal individuals;\textsuperscript{10,41} \( C_l \) (unbound) is about 0.25 L/hr/kg;\textsuperscript{41} \( C_l \) is stereospecific.\textsuperscript{45} The major metabolite is a mono-\( N \)-dealkylated form that has weak antiarrhythmic but potent anticholinergic activity; 55 ± 6% is excreted unchanged in urine.\textsuperscript{10}

\( t_{\beta} \). \( \alpha \) phase 2–4 min (IV);\textsuperscript{42} \( \beta \) phase is concentration dependent, usually 6 ± 1 hr,\textsuperscript{10} 11–17 hr in renal impairment, depending on severity.\textsuperscript{42}

Adverse Reactions. Nausea or anorexia occur frequently. Dry mouth, urinary retention, blurred vision, and constipation are dose-related anticholinergic effects that can occur in up to 70% of patients and result in drug discontinuation in about 20%.\textsuperscript{46} Through its vagolytic action, disopyramide can cause sinus tachycardia. Severe bradycardia, AV nodal block, or asystole also can occur, especially in patients with SA or AV nodal disease. Exacerbation of CHF is most prevalent (20–40%) in patients with left ventricular systolic dysfunction.\textsuperscript{38} Torsades de pointes, similar to quinidine syncope, has been reported. Rarely, rash, hepatic cholestasis, psychosis, or peripheral neuropathy occur. Hypoglycemia also has been reported.

Contraindications. History of disopyramide-induced heart block or serious ventricular arrhythmias; second- or third-degree heart block without a ventricular pacemaker; long-QT syndrome; cardiogenic shock or severe CHF.

Precautions. In atrial fibrillation or flutter, give digoxin or drugs that slow AV nodal conduction before giving disopyramide. Use very cautiously, if at all, in patients with CHF because of negative inotropic and vasoconstrictive actions.\textsuperscript{38} The drug can worsen sick sinus syndrome or aggravate underlying ventricular arrhythmias. If possible, use other antiarrhythmics in patients with prostatic hypertrophy or pre-existing urinary retention. Disopyramide can exacerbate glaucoma or myasthenia gravis.

Drug Interactions. Erythromycin inhibits disopyramide metabolism. Phenytoin can decrease disopyramide serum levels and increase its anticholinergic effects. Rifampin, barbiturates, and other enzyme inducers can decrease disopyramide serum levels. Concurrent use of disopyramide and quinidine can increase disopyramide serum levels or decrease quinidine serum levels.

Parameters to Monitor. Because of concentration-dependent protein binding, total drug levels unreliably reflect active drug concentration, and monitoring unbound drug concentrations is preferable. Monitor serum levels and symptoms or signs of toxicity closely in patients with altered states of drug disposition such as renal dysfunction. When initiating therapy, observe ECG daily for 3–4 days for QT, QRS, or PR prolongation. Frequently obtain vital signs initially for evidence of adverse hemodynamic effects (eg, CHF) and less frequently when a mainte-
nance dosage is attained. Question the patient about anticholinergic manifestations such as urinary and visual abnormalities.

**Notes.** A 1–10 mg/mL suspension, prepared from capsules, in cherry syrup is stable for 1 month with refrigeration in an amber bottle.

**DOFETILIDE**  
**Pharmacology.** Dofetilide is a class III antiarrhythmic drug that selectively prolongs atrial and ventricular repolarization by blocking the delayed rectifier (rapid component) potassium current. It is indicated for the termination and prevention of atrial fibrillation and flutter.

**Administration and Adult Dosage.** PO 125–500 µg bid, adjusted based on response and QT interval prolongation. Initiate therapy during hospitalization.

**Special Populations.** **Pediatric Dosage.** Safety and efficacy not established.

**Geriatric Dosage.** Same as adult dosage.

**Other Conditions.** Reduce maintenance dosages patients with renal dysfunction: 250 µg bid for Clcr 40–60 mL/min and 125 µg bid for Clcr 20–40 mL/min. Avoid the drug in patients with Clcr <20 mL/min.

**Dosage Forms.** Cap 125, 250, 500 µg.

**Pharmacokinetics.** Oral bioavailability is 96% and peak concentrations occur 2.5 hr after oral administration. About 20% of dofetilide is metabolized hepatically and 80% is eliminated renally as unchanged drug. Elimination half-life is 9.7 ± 2.7 hr with normal renal function.

**Adverse Reactions.** The major side effect is drug-induced torsades de pointes, which occurs in 1–10% of patients; risk increases with higher dosages. Other risk factors include female sex and underlying CHF.

**Contraindications.** Severe renal insufficiency; QT prolongation; hypokalemia; previous history of torsades de pointes; Clcr <20 mL/min.

**Drug Interactions.** Avoid using dofetilide with drugs that interfere with its renal elimination (eg, cimetidine, ketoconazole, trimethoprim and sulfamethoxazole, prochlorperazine, megestrol). Use caution with concurrent use of agents that block CYP3A4 (eg, verapamil, erythromycin). Do not use with other drugs that can prolong the QT interval.

**Parameters to Monitor.** Initiate dofetilide during hospitalization with continuous ECG monitoring. Decrease dosage if QT prolongation occurs; discontinue if excessive. Monitor renal function q 3 months.

**Notes.** When administered properly and monitored closely, dofetilide does not seem to increase mortality in patients with CHF. Azimilide is another agent currently under investigation that blocks potassium channels (both the rapid and slow components of the delayed rectifier). Azimilide is another agent currently under investigation that blocks potassium channels (both the rapid and slow components of the delayed rectifier).

**FLECAINIDE ACETATE**  
**Pharmacology.** Flecainide is a type Ic antiarrhythmic that predominantly slows conduction velocity, with minimal effect on refractoriness. (See Electrophysio-
logic Actions of Antiarrhythmics Comparison Chart.) Compared with type Ia or Ib antiarrhythmics, it binds to and dissociates from the sodium channel very slowly. It can decrease cardiac output by a negative inotropic action.

**Administration and Adult Dosage.** **PO** 50 mg q 12 hr initially, increasing in 50 mg increments q 12 hr q 4–7 days until desired response. **Usual maintenance dosage** is 100 mg PO q 12 hr to a maximum of 300 mg/day. Initiate flecainide during hospitalization.

**Special Populations. Pediatric Dosage.** Safety and efficacy not established. **PO** 100–200 mg/m²/day (average 140 mg/m²/day) in 2 divided doses has been used.52

**Geriatric Dosage.** Same as adult dosage.

**Other Conditions.** Lower maintenance dosage requirements are expected in patients with CHF, liver disease, or renal insufficiency. Start these patients with 50–100 mg q 12–24 hr and cautiously increase dosage as required with the aid of serum levels.41,53

**Dosage Forms.** **Tab** 50, 100, 150 mg.

**Patient Instructions.** Report any symptoms of dizziness, extra or rapid heart beats, or visual disturbances. Report symptoms of worsening shortness of breath or exercise intolerance.

**Missed Doses.** Take this drug at regular intervals. If you miss a dose and it has been less than 4 hours since your dose was due, take it as soon as you remember. If it is about time for the next dose, take that dose only. Do not double the dose or take extra.

**Pharmacokinetics. Onset and Duration.** Onset 1–6 hr (average 3); duration 12–30 hr.53

**Serum Levels.** (Therapeutic trough) 0.2–1 mg/L (0.5-2.5 μmol/L).53,54

**Fate.** Oral bioavailability is 70 ± 11%.10,54 From 37–55% is bound to plasma proteins, but the percentage can be higher (61%) post-MI because of increases in α1-acid glycoprotein.53 **Vₐ** is 8–10 L/kg,53 **Cl** has been reported as 0.34 ± 0.1 L/hr/kg,10 and 0.61 ± 0.23 L/hr/kg;41,53 **Cl** decreases with CHF, renal failure, and liver disease. Flecainide is about 60% stereoselectively metabolized by the liver through the CYP2D6 isozyme55 and about 30% excreted unchanged in urine.

**t½.** **α** phase 3–8 min; **β** phase 14 ± 5 hr. **β** phase is 20 ± 4 hr in patients with ventricular ectopy and 37.8 ± 39.7 hr in those with severe renal dysfunction.56

**Adverse Reactions.** Neurologic side effects, which include dizziness and visual abnormalities, occur frequently. Exacerbation of CHF in patients with underlying left ventricular dysfunction occurs frequently. Nausea, dyspnea, and headache also can occur frequently. Flecainide has proarrhythmic effects that can result in new sustained ventricular tachycardia or aggravation of underlying ventricular arrhythmias. These reactions occur more frequently in patients with left ventricular dysfunction, coronary disease, or ventricular arrhythmias.18,57 Risk can be sustained over time and not limited to the several days after initiation of therapy. Flecainide-induced ventricular tachycardia may be unresponsive to cardioversion.
or pacing but responsive to lidocaine therapy or sodium bicarbonate. Aggravation of underlying conduction disturbances also can occur.

**Contraindications.** Second- or third-degree AV block or bifascicular block without a ventricular pacemaker; severe CHF; history of type Ic–induced arrhythmia.

**Precautions.** Use with caution in patients with sick sinus syndrome and in combination with other negative inotropic drugs such as calcium-channel blockers or β-blockers or after recent therapy with a type Ia antiarrhythmic. Flecainide can increase pacemaker capture threshold.58 (See Notes.)

**Drug Interactions.** Amiodarone and cimetidine can increase flecainide serum concentrations; flecainide slightly elevates serum digoxin levels.

**Parameters to Monitor.** Frequent or continuous (preferred) ECG when therapy is initiated and then periodically on an ambulatory basis. Obtain a baseline evaluation of left ventricular function before starting flecainide. Obtain periodic trough serum levels (particularly in those with renal or liver disease and CHF) once an individual’s effective level is determined. Observe closely for neurologic toxicities and CHF symptoms when initiating therapy.

**Notes.** Because the CAST results showed increased mortality in patients with asymptomatic ventricular arrhythmias post-MI who were given flecainide,18 it should be reserved for individuals with life-threatening ventricular arrhythmias (eg, sustained ventricular tachycardia) refractory to other drugs.

**IBUTILIDE FUMARATE**

**Pharmacology.** Ibutilide is a class III antiarrhythmic that selectively prolongs atrial and ventricular repolarizations by increasing sodium influx (the window current) and blocking the rapid component of the delayed rectifier potassium current. It is indicated for the acute termination of atrial fibrillation or atrial flutter of recent onset. In these arrhythmias, sinus rhythm is restored in about 50% of patients.59,60

**Adult Dosage.** IV for atrial flutter or fibrillation (≥60 kg) 1 mg over 10 min; (<60 kg) 0.01 mg/kg. If the tachycardia is not terminated 10 min after the end of the initial infusion, the dose can be repeated.

**Dosage Forms.** Inj 0.1 mg/mL.

**Pharmacokinetics.** Ibutilide is approximately 40% bound to plasma proteins and has a Vd of 11 ± 4 L/kg.59,60 It is metabolized primarily by the liver. Although many metabolites have been identified, only a hydroxylated form has shown weak class III activity. Less than 10% is excreted unchanged in urine. Elimination half-life is about 6 hr (range 2–12 hr).60

**Adverse Reactions.** The major side effect is drug-induced proarrhythmia; torsades de pointes (sustained or nonsustained) occurs in 4–5% of patients. Risk factors are hypokalemia, underlying left ventricular dysfunction, and female sex.61 Rapid IV bolus administration can increase the risk of torsades de pointes.60 Prior administration of IV MgSO4 can prevent torsades de pointes. Heart block and heart failure have occurred rarely.
Contraindications. Pre-existing hypokalemia or hypomagnesemia; pre-existing long-QT interval; congenital long-QT syndromes; concurrent therapy with other drugs known to delay repolarization.

Precautions. Patients with atrial fibrillation of more than 2 days’ duration must be anticoagulated with warfarin for 3 weeks before the administration of ibutilide.

Parameters to Monitor. Give ibutilide with continuous ECG monitoring. Monitor QT interval and serum electrolytes before and after administration.

Pharmacology. Lidocaine’s electrophysiologic actions differ in healthy and diseased cardiac tissues. (See Electrophysiologic Actions of Antiarrhythmics Comparison Chart.) Most of its antiarrhythmic activity is caused by frequency-dependent blockade of the fast sodium channel in Purkinje fibers. In comparison with other antiarrhythmics, lidocaine binds to and dissociates from the sodium channel very quickly. It is used in the acute treatment of ventricular arrhythmias often associated with MI. Effectiveness in the treatment of supraventricular arrhythmia is limited.

Administration and Adult Dosage. IV loading dose for ventricular tachycardia or fibrillation 100 mg (1–1.5 mg/kg) over 1 min; if ineffective, repeat with 50–100 mg q 5–10 min, to a maximum of 300 mg. IV maintenance 2–4 mg/min infusion. IV for neuropathic pain 5 mg/kg/hr for 60–90 min has been used. (See Notes.)

Special Populations. Pediatric Dosage. IV (or intratracheal) loading dose 1 mg/kg; can repeat q 10–15 min to a maximum of 3–5 mg/kg. IV maintenance dosage 20–50 μg/kg/min infusion.

Geriatric Dosage. Same as adult dosage. The elderly can be at increased risk for toxicity because of decreased clearance.

Other Conditions. In CHF, use one-half of IV loading dose. In liver disease or CHF, initial maintenance infusion is 1 mg/min, to a maximum of 2–3 mg/min. In MI without CHF, maintenance infusion rate might need to be decreased by 30–50% in 24 hr; however, empiric dosage alterations in MI are not recommended because of increases in α1-acid glycoprotein and lidocaine binding.

Dosage Forms. Inj 10, 20, 40, 100, 200 mg/mL. Also available premixed in D5W in concentrations of 2, 4, and 8 mg/mL.

Patient Instructions. Report side effects such as drowsiness, perioral numbness or tingling, dizziness, and nausea during maintenance infusion.

Pharmacokinetics. Onset and Duration. IV onset is immediate; duration after initial IV bolus is 10–20 min. IM onset is 10 min; duration is 3 hr. Serum Levels. Therapeutic (total), 1.5–6 mg/L (7–28 μmol/L); unbound, 0.5–1.5 mg/L (2–7 μmol/L). Toxic reactions are more likely at total concentrations >5 mg/L (22 μmol/L). (See Adverse Reactions.)

Fate. The drug is well absorbed orally, but a large hepatic first-pass effect limits systemic availability to 35 ± 11%. IM absorption half-life is 12–28 min. The drug is 70 ± 5% bound to plasma proteins; Vd is 1.3 ± 0.4 L/kg in normal individ-
uals and 0.9 ± 0.2 L/kg in patients with CHF. Cl is 0.55 ± 0.14 L/hr/kg, decreased in CHF, liver disease, and during long-term infusion. Lidocaine is metabolized primarily in the liver, with 2 ± 1% excreted unchanged in the urine. The major metabolites, monoethylglycinexylidide (MEGX) and glycinexylidide (GX), have neurotoxic and antiarrhythmic actions. Accumulation of these metabolites in renal impairment or during prolonged infusions can contribute to lidocaine toxicity.

The major metabolites, monoethylglycinexylidide (MEGX) and glycinexylidide (GX), have neurotoxic and antiarrhythmic actions. Accumulation of these metabolites in renal impairment or during prolonged infusions can contribute to lidocaine toxicity.

$\alpha$ phase about 8 min; $\beta$ phase 98 ± 24 min. The $\beta$ phases in CHF and liver disease can be prolonged to 4.5 ± 2.4 hr and 6.6 ± 1.1 hr, respectively. Elimination half-life of total lidocaine increases to an average of 3.2 ± 0.5 hr 24 hr after MI without CHF and up to 10.2 ± 2 hr after MI with CHF. In MI, the rise in total lidocaine half-life is greater than that of unbound lidocaine.

**Adverse Reactions.** Serum level–related neurologic side effects including dizziness, nausea, drowsiness, speech disturbances, perioral numbness, muscle twitching, confusion, vertigo, and tinnitus are frequent at total serum levels >5 mg/L. Serious toxicities including psychosis, seizures, and respiratory depression occur at serum levels >9 mg/L. Sinus arrest or severe bradycardia is associated with sinus node disease, toxic drug levels, or concomitant therapy with other antiarrhythmics. Complete AV block can occur, especially in patients with pre-existing bifascicular bundle branch block, AV nodal block, or inferior wall MI.

**Contraindications.** History of hypersensitivity to any amide-type local anesthetic (rare); second- or third-degree heart block unless the site of the block can be localized to the AV node itself or ventricular pacemaker is functional; severe sinus node dysfunction; Stokes–Adams syndrome; atrial fibrillation in association with Wolff–Parkinson–White syndrome.

**Precautions.** Lidocaine administered to prevent ventricular fibrillation in acute MI is no longer recommended. Toxicity during bronchoscopy caused by tracheal lidocaine absorption has been reported.

**Drug Interactions.** Propranolol decreases lidocaine clearance, so close monitoring is necessary with concomitant administration of these drugs. Cimetidine can decrease lidocaine clearance, but empiric dosage reduction with concomitant cimetidine is not recommended. Phenytoin can decrease lidocaine serum levels and increase myocardial depression.

**Parameters to Monitor.** Closely monitor serum levels and signs or symptoms of toxicity in patients with altered drug dispositions such as CHF, hepatic disease, acute MI, or prolonged IV infusion (>24 hr). Monitoring unbound levels is preferable post-MI. Minor subjective and objective toxicities are extremely important because they are often subtle and can forecast more serious toxicities (eg, psychosis or seizures). Continuously observe ECG for therapeutic and/or toxic actions. Frequently monitor vital signs such as blood pressure, heart rate, and respiration.

**Notes.** IV lidocaine has been used to treat pain of peripheral origin such as neuropathies and burns.
Pharmacology. Mexiletine has electrophysiologic actions similar to those of lidocaine and tocainide. Depression of conduction is accentuated in ischemic/hypoxic tissue. It also has a slight negative inotropic action. It is used in the treatment of ventricular arrhythmias; effectiveness in supraventricular tachycardias is limited.

Administration and Adult Dosage. PO loading dose 400 mg once, followed by maintenance dosage in 8 hr; PO maintenance dosage 200–300 mg q 8 hr, to a maximum of 400 mg q 8 hr. PO for neuropathic pain 450 mg/day;76 dosages as high as 10 mg/kg/day have been used to treat the thalamic pain syndrome.77 (See Notes.) Initiate mexiletine during hospitalization.

Special Populations. Pediatric Dosage. Safety and efficacy not established.

Geriatric Dosage. Same as adult dosage.

Other Conditions. Reduce maintenance dosage by 30–50% in patients with hepatic disease or severe CHF.41 Dosage also might need to be decreased with Cl<sub>cr</sub> < 10 mL/min.78

Dosage Forms. Cap 150, 200, 250 mg.

Patient Instructions. Report numbness, drowsiness, dizziness, or tingling. Nausea or loss of appetite can occur and reduced by taking the drug with food. Report any abnormal bruising.

Missed Doses. Take this drug at regular intervals. If you miss a dose and it has been less than 4 hr since your dose was due, take it as soon as you remember. If it is about time for the next dose, take that dose only. Do not double the dose or take extra.

Pharmacokinetics. Onset and Duration. PO onset 1–4 hr (average 2); duration 8–16 hr.

Serum Levels. Between 0.5 and 2 mg/L (3–11 μmol/L), although not well correlated with therapeutic or toxic effects.79

Fate. Oral bioavailability is 87 ± 13%, and, unlike lidocaine, mexiletine undergoes less than 10% first-pass hepatic elimination.79,80 Absorption can be incomplete in MI patients receiving narcotic analgesics.79,80 The drug is 63 ± 3% bound to plasma proteins;10 V<sub>d</sub> is large and has been variably reported as 6.6 ± 0.9 L/kg and 10.8 ± 7.2 L/kg.79,80 Cl is variable, 0.4–0.6 L/hr/kg decreased in CHF and liver disease.79 Mexiletine is metabolized predominantly in the liver, where it undergoes polymorphic metabolism, primarily by the CYP2D6 isozyme;81 10–20% is excreted unchanged in urine, depending on urinary pH.79,80

t<sub>1/2</sub>. α phase 3–12 min, β phase 9.2 ± 2.1 hr and 18.5 hr in poor metabolizers,81 15.7 ± 4.9 hr in severe renal dysfunction,78 and 15 ± 0.6 hr in CHF with or without MI.83,84 and can be prolonged in cirrhosis.

Adverse Reactions. Neurologic toxicities are frequent and include tremor, ataxia, drowsiness, confusion, paresthesias, and occasionally psychosis or seizures. Minor CNS side effects can occur in up to 40% of patients.54 Nausea, vomiting, and anorexia are frequent. Mexiletine can aggravate underlying ventricular arrhythmias or conduction disturbances. Thrombocytopenia has been reported.
rarely. Mexiletine is an ether analogue of lidocaine, so cross-sensitivity between mexiletine and tocainide or lidocaine is not expected.

**Contraindications.** Second- or third-degree AV block without a ventricular pacemaker; cardiogenic shock.

**Precautions.** Sick sinus syndrome can worsen. Mexiletine can increase pacemaker capture threshold and alter the effectiveness of internal defibrillators.

**Drug Interactions.** Mexiletine increases theophylline concentrations by 30–50% by decreasing theophylline metabolism. Phenytoin and rifampin can increase mexiletine metabolism. Quinidine and theophylline occasionally increase serum mexiletine levels.

**Parameters to Monitor.** ECG for 3–5 days when therapy is initiated and then q 3–6 months on an ambulatory basis. Obtain periodic serum levels once an individual’s effective level is determined. Observe closely for neurologic toxicities when initiating therapy.

**Notes.** The efficacy of mexiletine for ventricular tachycardia can be increased by adding a type Ia antiarrhythmic such as quinidine. Mexiletine has been used to treat neuropathic pain such as diabetic neuropathy and for thalamic pain syndrome.

**MORICIZINE HYDROCHLORIDE**

**Pharmacology.** Moricizine is a phenothiazinelike type I (probably Ic) antiarrhythmic that (in normal tissue) slows conduction velocity by blocking sodium channels in a frequency-dependent manner. Its effects appear to be accentuated by ischemia.

**Administration and Adult Dosage.** PO 200 q 8 hr initially, increasing daily dosage q 3 days by 150 mg until desired effect or toxicity occurs, to a usual maintenance dosage of 200–300 mg q 8 hr. Initiate moricizine during hospitalization.

**Special Populations.** Pediatric Dosage. Safety and efficacy not established. PO 200–600 mg/m²/day has been used.

**Geriatric Dosage.** Same as adult dosage.

**Other Conditions.** Not well studied; patients with hepatic disease might require lower dosages.

**Dosage Forms.** Tab 200, 250, 300 mg.

**Patient Instructions.** Report any dizziness, rapid heartbeat, or gastrointestinal upset.

**Missed Doses.** Take this drug at regular intervals. If you miss a dose and it has been less than 4 hours since your dose was due, take it as soon as you remember. If it is about time for the next dose, take that dose only. Do not double the dose or take extra.

**Pharmacokinetics.** Onset and Duration. Onset is variable at 2–20 hr after multiple doses; duration is 12–36 hr after long-term use.

**Serum Levels.** Correlation between serum levels and therapeutic effect is not well established, but 0.2–3.6 mg/L (0.5–8.4 μmol/L) has been suggested.
**Fate.** Moricizine is well absorbed after oral administration, but the large hepatic first-pass metabolism limits bioavailability to 34–38%. About 81–90% is bound to plasma proteins.\(^9^3\) \(V_d\) is 5.9 ± 3.2 and 11.6 ± 6.7 L/kg after 1 and 13 days of therapy, respectively.\(^9^4\) CI is 3.8 ± 1.8 to 4.7 ± 2.3 L/hr/kg, depending on length of therapy.\(^9^5\) Moricizine undergoes extensive hepatic metabolism and appears to induce its own metabolism. More than 40 metabolites appear systemically in small quantities.\(^9^5\) At least two, including moricizine sulfoxide, are active and probably account for some of the drug’s antiarrhythmic activity and for its long duration of action. Less than 1% appears in the urine unchanged.\(^9^4,9^5\)

\[ t_{1/2} \alpha \text{ phase } 4–20 \text{ min} ; \beta \text{ phase } 1.6 \pm 0.2 \text{ hr}. \]

**Adverse Reactions.** Frequent noncardiac side effects include nausea, anorexia, and dizziness. Dizziness can be lessened by administering more frequent, smaller doses. Moricizine has proarrhythmic actions that result in new or worsened ventricular tachycardia in 2–5% of patients. Exacerbation of CHF occurs occasionally. Underlying conduction disturbances such as AV block, ventricular conduction defects, or sick sinus syndrome can worsen. Drug fever has been reported.

**Contraindications.** Second- or third-degree AV block or bifascicular block without a ventricular pacemaker; cardiogenic shock; hypersensitivity to phenothiazines.

**Precautions.** Use with caution in patients with sick sinus syndrome. Because of the final results of CAST II,\(^9^6\) moricizine is indicated only for life-threatening ventricular arrhythmias such as sustained ventricular tachycardia, where there is a clear benefit to therapy.

**Drug Interactions.** Cimetidine can increase moricizine serum levels. Moricizine decreases theophylline levels.

**Parameters to Monitor.** Daily ECG for the first 2–4 days, when therapy is initiated, and then q 3–6 months on an ambulatory basis; watch for PR and QRS lengthening and for GI side effects and dizziness.

**Notes.** Limited data exist on the use of moricizine in supraventricular tachycardias.

**PROCAINAMIDE HYDROCHLORIDE**

**Pharmacology.** Procainamide is a class Ia antiarrhythmic that alters conduction in normal and ischemic tissues by sodium-channel blockade in a fashion similar to that of quinidine. It can decrease systemic blood pressure by causing peripheral ganglionic blockade;\(^9^7\) it also has weak anticholinergic action and a slight negative inotropic action. The active metabolite \(N\)-acetylprocainamide (NAPA) has primarily type III antiarrhythmic activity that predominantly delays repolarization by blocking potassium conductance.

**Administration and Adult Dosage.** PO loading dose (Cap, Tab) 1 g over 2 hr in 2 divided doses. PO maintenance dosage (Cap, Tab) 1–6 g/day in 4–6 divided doses, to a maximum of 9 g/day.\(^9^8\) (SR Tab) can be given q 6–8 hr (Pronestyl-SR) or q 12 hr (Procanbid), to a maximum of 50 mg/kg/day. IV loading dose 1–1.5 g at 20–50 mg/min\(^9^9\) or 15–20 mg/kg. IV maintenance dosage 1.5–5 mg/min.
(20–80 μg/kg/min) infusion.\textsuperscript{97} Intermittent IV or IM 1–6 g/day in 4–6 divided doses, to a maximum of 9 g/day. Initiate procainamide during hospitalization.

**Special Populations. Pediatric Dosage.** Safety and efficacy not established. PO 15–50 mg/kg/day in 4–8 divided doses, to a maximum of 4 g/day; IV loading dose 2–6 mg/kg over 5 min (up to 100 mg/dose); can repeat q 5–10 min, to a maximum of 15 mg/kg. IV maintenance dosage 20–80 μg/kg/min infusion, to a maximum of 2 g/day. IM maintenance dosage 20–30 mg/kg/day in 4–6 divided doses, to a maximum of 4 g/day.\textsuperscript{4}

**Geriatric Dosage.** Same as adult dosage but adjust for age-related decrease in renal function.

**Other Conditions.** Reduce maintenance dosage in liver disease. In renal insufficiency, procainamide and its active metabolite accumulate, necessitating a lower maintenance dosage.\textsuperscript{41} Recent data imply no need for decreasing loading and maintenance dosages in CHF and MI.\textsuperscript{99}

**Dosage Forms.** Cap, Tab 250, 375, 500 mg; SR Tab (6-hr; Pronestyl-SR, various) 250, 500, 750, 1000 mg; (12-hr; Procanbid) 500, 1000 mg; Inj 100, 500 mg/mL.

**Patient Instructions.** Report any symptoms such as nausea, vomiting, fever, sore throat, joint pain, rash, chest or abdominal pain, and shortness of breath. Do not chew, split, or crush SR tablets. A sustained-release tablet shell in the stool does not indicate lack of absorption.

**Missed Doses.** Take this drug at regular intervals. If you miss a dose, take it as soon as you remember. If it is about time for the next dose, take that dose only. Leave at least 2 hours between regular capsule or tablet doses and 4–6 hours between sustained-release tablet doses. Do not double the dose or take extra.

**Pharmacokinetics. Onset and Duration.** IV onset is immediate; PO and IM onsets occur within 1 hr; SR Tab preparations are somewhat slower. Duration is usually 3–6 hr.

**Serum Levels.** Therapeutic range is 4–10 mg/L (17–43 μmol/L);\textsuperscript{42} toxicity is more likely at serum levels above 12 mg/L (51 μmol/L). In some arrhythmias (eg, recurrent ventricular tachycardia), levels of at least 20 mg/L (85 μmol/L) may be required to prevent arrhythmias, with average effective levels of 13 mg/L.\textsuperscript{98} Effective serum levels of NAPA are 15–25 mg/L (53–88 μmol/L), with overlap between the toxic and therapeutic ranges.\textsuperscript{100}

**Fate.** Oral bioavailability is 83 ± 16%;\textsuperscript{10} about 16 ± 5% is bound to plasma proteins; \(V_d\) is 1.9 ± 0.3 L/kg.\textsuperscript{10} The drug is 67 ± 8% excreted in the urine as unchanged; the remainder is metabolized, mostly to active NAPA by the liver, with smaller amounts excreted as para-aminobenzoic acid. Cl is highly variable depending on acetylator status and renal function. The total quantity of NAPA produced depends on liver function and acetylator phenotype.\textsuperscript{10,101}

\(t_{1/2}^\alpha\) (Procainamide) α phase about 6 min; β phase 3 ± 0.6 hr in normal individuals, 5.3–20.7 hr in patients with renal dysfunction, and 12.5 ± 1.4 hr in anephric patients. (NAPA) 7 ± 1 hr, 41.5 ± 7.8 hr in renal failure.\textsuperscript{10,100,101}
**Adverse Reactions.** About 50–80% of patients develop a positive ANA, with 30–50% developing symptoms of SLE; genetically slow acetylators more rapidly develop positive ANA and SLE symptoms. Common SLE symptoms or signs are rash, arthralgias, fever, pericarditis, and pleuritis. Although drug cessation usually reverses these symptoms in about 2 weeks, some patients have prolonged manifestations; for others, the SLE syndrome initially can be life threatening. Hypotension frequently can occur after rapid IV administration. Severe bradycardia, AV nodal block, or asystole has been reported. Procainamide can aggravate underlying ventricular arrhythmias and cause torsades de pointes. GI symptoms occur frequently and include nausea and vomiting; drug fever and dermatologic reactions occasionally occur. Agranulocytosis has been reported occasionally and can be fatal. Whether the SR product carries a higher risk of neutropenia than the fast-release preparation is controversial. Hepatitis has been reported rarely.

**Contraindications.** SLE (including that induced by drugs); second- or third-degree heart block without a ventricular pacemaker; long-QT syndrome; severe sinus node dysfunction or torsades de pointes caused by other type Ia antiarrhythmics.

**Precautions.** In atrial fibrillation or flutter, procainamide paradoxically can increase ventricular rate; administer digoxin or other drugs that slow AV nodal conduction before procainamide. Procainamide can worsen symptoms of sick sinus syndrome and exacerbate myasthenia gravis.

**Drug Interactions.** Amiodarone, trimethoprim, cimetidine, and, to a lesser extent, ranitidine can increase procainamide levels; alcohol can decrease levels.

**Parameters to Monitor.** Monitor serum levels and symptoms or signs of toxicity in patients with suspected altered drug dispositions such as hepatic disease or renal dysfunction. Monitor ECG continuously (with IV) or daily initially (with PO) for QRS, QT, and PR prolongation; monitor oral therapy less frequently once maintenance dosage has been established. Monitor blood pressure frequently when therapy is initiated (especially with IV) and less frequently once a maintenance dosage has been established. Periodically monitor WBC count and signs of infection for development of drug-induced agranulocytosis. Observe closely for symptoms of drug-induced SLE.

**Propafenone Hydrochloride**

**Pharmacology.** Propafenone is a sodium-channel blocker that slows predominantly atrial and ventricular conduction velocities without appreciably prolonging repolarization. It is therefore classified as a type Ic antiarrhythmic, similar to flecainide. Propafenone is administered as a racemate; the enantiomers and the 5-hydroxy metabolite are equipotent sodium-channel blockers. Propafenone, in particular the (S)-enantiomer, and its active 5-hydroxy metabolite also have variable, nonselective β-blocking actions.

**Administration and Adult Dosage.** PO 150 mg q 8 hr initially, increasing q 3–4 days to desired effect or toxicity. PO maintenance 150–200 mg q 8 hr, to a maximum of 1.2 g/day. Initiate propafenone during hospitalization.
Special Populations. Pediatric Dosage. Safety and efficacy not established. PO 10–20 mg/kg/day in 2–3 divided doses has been used.107

Geriatric Dosage. Same as adult dosage. Lower initial dosages have been suggested.108

Other Conditions. Bioavailability and half-life are increased in patients with hepatic disease, and a dosage reduction of 50% has been suggested109 and questioned.106 Lower initial dosages have been suggested for patients with renal dysfunction.106

Dosage Forms. Tab 150, 225, 300 mg.

Patient Instructions. Report any symptoms of dizziness, rapid heartbeat, blurred vision, or shortness of breath.

Missed Doses. Take this drug at regular intervals. If you miss a dose, take it as soon as you remember. If it is about time for the next dose, take that dose only. Leave at least 4 hours between doses. Do not double the dose or take extra.

Pharmacokinetics. Onset and Duration. Onset 2–4 hr; peak 2–6 hr; duration 4–22 hr.106,108

Serum Levels. No established therapeutic range. Levels of parent compound and metabolite are highly variable, depending on genetically determined variations in hepatic metabolisms. Mean minimal effective concentration was 0.2 mg/L (6 μmol/L) in one study.110 Side effects are more frequent when the trough propafenone level exceeds 0.9 mg/L (26 μmol/L).111

Fate. Completely absorbed after oral administration, but large hepatic first-pass metabolism limits bioavailability to 12.1 ± 11%. First-pass elimination appears saturable, so bioavailability is highly variable and increases with larger oral doses and long-term therapy.106,108,111 About 85–95% is bound to plasma proteins, primarily α1-acid glycoprotein.106 Vd is 3.6 ± 2.6 L/kg.112 The parent drug undergoes polymorphic hepatic metabolism via CYP2D6. Extensive metabolizers (EMs; about 90% of patients) form clinically important quantities of the active metabolite 5-hydroxypropafenone; poor metabolizers (PMs; about 10% of patients) form little of this compound.108,111 Another active metabolite, N-desethylpropafenone, is not subject to genetic polymorphism.106,111 Cl is 0.96 ± 1.08 L/hr/kg in EMs and 0.23 ± 0.042 L/hr/kg in PMs.113 Cl is also stereospecific.

\[ t_{1/2} \] α phase 5 min; β phase (EMs) 5.5 ± 2 hr; (PMs) 17 ± 8 hr.111

Adverse Reactions. Frequent noncardiac side effects include metallic or bitter taste in 15–20% of patients, and nausea and CNS toxicity such as dizziness and headache in 10–15% of patients.112 Because of the β-blocking activity of propafenone, worsening of asthma or obstructive lung disease can occur.105,112 Propafenone has proarrhythmic actions (sometimes life-threatening) that can result in new or worsened ventricular tachycardia. This can occur in 5–15% of patients, particularly in those with poor left ventricular function caused by structural heart disease or with underlying ventricular tachycardia. Worsening of existing CHF or underlying conduction disturbances, such as AV block or sick sinus syndrome, can occur. Cholestatic jaundice occurs rarely.114
Contraindications. Second- or third-degree AV block or bifasicular block without a ventricular pacemaker; history of type Ic-induced arrhythmia; bronchospastic disorders; uncontrolled CHF; cardiogenic shock; marked hypotension; sick sinus syndrome; bradycardia; electrolyte imbalance.

Precautions. Propafenone can increase pacemaker capture threshold and affect the efficacy of internal defibrillators. Because of the CAST results (although not studied in this trial), propafenone is indicated only for arrhythmias where there is a clear benefit to therapy.

Drug Interactions. Propafenone inhibits hepatic enzymes and reportedly increases serum concentrations of digoxin, theophylline, warfarin, and β-blockers.

Parameters to Monitor. Daily or continuous (preferred) ECG for 3–4 days initially and then q 3–6 months on an ambulatory basis. Observe closely for CNS symptoms such as dizziness.

Notes. Although not a labeled indication, propafenone can be effective for some supraventricular arrhythmias.

Pharmacology. Quinidine is a class Ia antiarrhythmic that slows conduction velocity, prolongs effective refractory period, and decreases automaticity of normal and diseased fibers. (See Electrophysiologic Actions of Antiarrhythmics Comparison Chart.) The cellular mechanism appears to be frequency-dependent blockade of the fast sodium channel. Quinidine also blocks potassium conductance, particularly at low concentrations. AV nodal conduction can be increased reflexly through vasodilation, attributed to peripheral α-adrenergic blockade or vagolytic action. Slight negative inotropic action might be clinically important in patients with severe CHF.

Administration and Adult Dosage. IM and PO loading doses not recommended. PO maintenance dosage generally 200–400 mg q 6–8 hr; SR products (gluconate) can be given q 12 hr, (polygalacturonate) can be given q 8 hr. IV loading dose (gluconate) 5–8 mg/kg (3.75–6 mg/kg in CHF) at a rate of 0.3 mg/kg/min. (See Notes.) Initiate quinidine during hospitalization.

Special Populations. Pediatric Dosage. PO (gluconate salt) 15–60 mg/kg/day in 4 divided doses. IV and IM not recommended.

Geriatric Dosage. (>60 yr) use lower initial dosages and adjust maintenance dosage based on side effects, therapeutic response, and serum levels.

Other Conditions. In liver disease, CHF, or renal disease, use lower initial dosages and adjust maintenance dosages based on side effects, therapeutic response, and serum levels.

Dosage Forms. Tab (sulfate) 200, 300 mg; (polygalacturonate) 275 mg; SR Tab (gluconate) 324, 330 mg; (sulfate) 300 mg; Inj (gluconate) 80 mg/mL. (See Notes.)
**Patient Instructions.** Report any symptoms such as blurred vision, dizziness, tinnitus, diarrhea, abnormal bleeding or bruising, rash, or fainting episodes. Do not crush or chew sustained-release tablets. A sustained-release tablet shell in the stool does not indicate lack of absorption.

**Missed Doses.** Take this drug at regular intervals. If you miss a dose and it has been less than 2 hours since your dose was due, take it as soon as you remember. If it is about time for the next dose, take that dose only. Do not double the dose or take extra.

**Pharmacokinetics.**

**Onset and Duration.** PO onset of sulfate within 1 hr; SR gluconate and polygalacturonate salts 2–4 hr. IM onset within 1 hr; IV is immediate. Duration (sulfate) 6–8 hr; SR (gluconate) 12 hr; (polygalacturonate) 8–12 hr.

**Serum Levels.** Therapeutic range about 2–6 mg/L (6–18 μmol/L), depending on assay. Toxicity is more likely with serum levels above 6 mg/L.41,42

**Fate.** Oral sulfate and gluconate are 80 ± 15% and 71 ± 17% bioavailable, respectively, with some first-pass elimination; bioavailability is increased in the elderly; IM absorption is incomplete.10,116 The drug is 87 ± 3% bound to plasma proteins.10 Vd is 2.7 ± 1.2 L/kg and 1.8 ± 0.5 L/kg in patients with CHF; Cl is 0.28 ± 0.11 L/hr/kg.10,117 The elderly and patients with liver disease or CHF are likely to have decreased clearance.10 Quinidine is metabolized primarily in the liver to two active metabolites, 3-hydroxyquinidine and 2'-quinidinone, and 18 ± 5% of a dose is excreted unchanged in urine.

\[ t_{1/2} \] phase about 7 min; \[ t_{1/2} \] phase in normal individuals, 6.2 ± 1.8 hr.10 In CHF, Cl and \[ V_d \] are decreased, so elimination half-life remains about the same.117 Half-life in alcoholic cirrhosis is prolonged to 9 ± 1 hr.118

**Adverse Reactions.** Diarrhea occurs in up to 30% of patients receiving quinidine and can be treated with aluminum hydroxide gel or lessened by using the polygalacturonate salt. Nausea or vomiting occurs frequently. Cinchonism can occur with high levels of quinidine; symptom complex includes tinnitus, blurred vision, headache, and nausea; in severe cases it can progress to delirium and psychosis. Hypotension can occur, especially after IV administration. Quinidine can aggravate underlying ventricular arrhythmias or CHF. Non–dose-related syncope, attributed to drug-induced torsades de pointes, can occur in 1–8% of patients, usually during the first week of therapy; it can occur in association with hypokalemia and/or hypomagnesemia.119 Asystole or AV nodal block has been reported. Rare or occasional idiosyncratic reactions include hepatitis, drug fever, anaphylactoid reactions, SLE, thrombocytopenia, and hemolytic anemia. IM use can cause pain and muscle damage.116

**Contraindications.** History of immunologic reaction to quinidine or quinine; previous occurrence of quinidine syncope; second- or third-degree heart block without a ventricular pacemaker; severe sinus node dysfunction or long-QT syndrome; digitalis intoxication; myasthenia gravis.

**Precautions.** In atrial fibrillation or flutter, administer digoxin or other drugs that decrease AV nodal conduction before administering quinidine. Chronic quinidine use in patients with atrial fibrillation is associated with increased mortality,120 which can be caused by torsades de pointes that occurs late in therapy.121
Drug Interactions. Quinidine inhibits CYP2D6 and can alter the disposition of many drugs that undergo genetically determined polymorphic metabolism through this pathway. Use care with concurrent digoxin and quinidine therapy because quinidine increases digoxin serum levels approximately 2-fold by inhibiting P-glycoprotein.\textsuperscript{122} Urinary alkalinization (eg, with acetazolamide or antacids) can decrease quinidine clearance. Phenytoin can increase quinidine metabolism. Amiodarone and cimetidine can reduce quinidine clearance. Quinidine occasionally increases warfarin response and serum levels of tricyclic antidepressant.

Parameters to Monitor. Monitor serum levels and signs or symptoms of toxicity in patients with altered drug dispositions such as CHF or liver disease. With ECG, monitor daily for QT, QRS, or PR prolongation for the first 2–4 days of therapy and then q 3–6 months on an ambulatory basis. Frequently monitor blood pressure (especially with IV) when therapy is initiated. Monitoring can decrease after a maintenance dosage has been determined. Monitor liver enzymes during the first 4–8 weeks of therapy. Monitor other parameters such as platelet count and hematocrit only if idiosyncratic reactions are suspected.

Notes. Adjust dosage when switching from one salt form to another; sulfate salt contains 83% quinidine, gluconate 62%, and polygalacturonate 60%. The gluconate and polygalacturonate forms are slowly dissociating salts of quinidine.

Pharmacology. Sotalol is a type III antiarrhythmic that is commercially available as a racemate: the L-isomer has nonselective β-blocking actions, and the D- and L-isomers delay repolarization by blockade of potassium channels. (See Electrophysiologic Actions of Antiarrhythmics Comparison Chart.) Sotalol is effective for ventricular and (unlabeled) supraventricular arrhythmias.

Administration and Adult Dosage. PO for ventricular arrhythmias (Betapace) 80 mg bid initially, increasing at 2- to 3-day intervals, to a maximum of 640 mg/day in 2 or 3 divided doses. Reserve high dosages (480–640 mg/day) for drug-refractory ventricular arrhythmias. PO for atrial fibrillation or flutter (Betapace AF) 80 mg bid initially, increasing at 3-day intervals, to a maximum of 160 mg bid. Initiate sotalol during hospitalization.

Special Populations. Pediatric Dosage. Safety and efficacy not established. PO 2–8 mg/kg/day in 2 divided doses has been used.\textsuperscript{123} Geriatric Dosage. Same as adult dosage.

Other Conditions. Reduce frequency of administration in patients with renal insufficiency as follows. Ventricular arrhythmias (Betapace) Cl\textsubscript{cr} 30–60 mL/min, q 24 hr; Cl\textsubscript{cr} 10–30 mL/min, q 36–48 hr.\textsuperscript{124} Use with caution, if at all, in patients with Cl\textsubscript{cr} <10 mL/min. Atrial arrhythmias (Betapace AF) Cl\textsubscript{cr} 40–60 mL/min, q 24 hr; Cl\textsubscript{cr} <40 mL/min, contraindicated.

Dosage Forms. Tab (Betapace) 80, 120, 160, 240 mg; (Betapace AF) 80, 120, 160 mg.

Patient Instructions. Report any symptoms of fainting, dizziness, shortness of breath, or fatigue.
**Missed Doses.** Take this drug at regular intervals. If you miss a dose and your next dose is more than 8 hours away, take it as soon as you remember. If it is about time for the next dose, take that dose only. Do not double the dose or take extra.

**Pharmacokinetics.** **Onset and Duration.** PO onset 1–3 hr; duration 12–18 hr.

**Serum Levels.** 1–3 mg/L (3.7–11 μmol/L), although not well correlated with therapeutic effect. Concentrations required to achieve delay in repolarization might be greater than those for β-blockade.123

**Fate.** Bioavailability is 90–100% with negligible first-pass metabolism.124 AUC is decreased 20% by food.124 The drug is not bound to plasma proteins. Vd is 1.2–2.4 L/kg; Cl is 0.13 ± 0.04 L/hr/kg;124–126 80–90% is excreted unchanged in urine.124 The disposition of the d-isomer is similar to that of the racemate.125,126

\[ t_{1/2} \]

\[ t_{1/2} \text{ phase 3–5 min; } t_{1/2} \text{ phase variously reported as 7.5 ± 0.8 to 17.5 ± 9.7 hr.} \]

Half-life is highly dependent on renal function: 22.7 ± 6.4 hr for Clcr 30–80 mL/hr; 64 ± 27 hr for Clcr 10–30 mL/hr; and 98 ± 57 hr for Clcr <10 mL/min.127

**Adverse Reactions.** Fatigue, dyspnea, and bradycardia occur frequently, probably caused by the β-blocking actions of sotalol.128 Exacerbation of CHF (1.7%) and asthma also can occur. (See Propranolol.) Sotalol induces arrhythmias, usually torsades de pointes, in 4.6% of patients.128 Risk factors for torsades de pointes are hypokalemia, hypomagnesemia, concurrent diuretic usage, and high sotalol dosages.129

**Contraindications.** (Betapace and Betapace AF) Asthma; second- and third-degree AV blocks without a ventricular pacemaker; sinus bradycardia; cardiogenic shock; long QT-syndrome; uncontrolled CHF; (Betapace AF, additional) sick sinus syndrome; baseline QT interval > 450 msec; hypokalemia (<4 mEq/L); Clcr <40 mL/min.

**Precautions.** Use with caution in sinus node dysfunction. Because of its β-blocking actions, use sotalol with caution in patients with diabetes, depressed left ventricular function, obstructive pulmonary disease, or peripheral vascular disease. Do not abruptly discontinue the drug in patients with coronary artery disease. Use with caution with electrolyte disorders, other drugs that prolong QT interval, or pre-existing QT prolongation. Escalate dosage only after achieving steady state (2–3 days).130

**Drug Interactions.** Because of its β-blocking actions, observe β-blocker interaction precautions. (See Propranolol.)

**Parameters to Monitor.** Baseline and daily ECG for the first 2–5 days, when therapy is initiated or dosage is adjusted, and then q 3–6 months on an ambulatory basis. QT prolongation to over 550 msec is an indication to discontinue sotalol because of the risk of torsades de pointes.

**Notes.** Based on the CAST results,18 many clinicians use type III antiarrhythmics (eg, amiodarone, sotalol) as first-line therapy in supraventricular and ventricular arrhythmias.
Pharmacology. Tocainide has electrophysiologic actions similar to those of lidocaine and mexiletine. Depression of conduction is accentuated in ischemic/hypoxic tissue. Antiarrhythmic actions are somewhat stereospecific. It also has a slight negative inotropic action. It is used in the treatment of ventricular arrhythmias, but it has limited effectiveness in supraventricular tachycardias. There appears to be a concordance of response (and nonresponse) between tocainide and lidocaine.131

Administration and Adult Dosage. PO 400 mg q 8 hr initially; usual maintenance dosage 1.2–1.8 g/day, to a maximum of 2.4 g/day in 2–3 divided doses. PO during lidocaine to tocainide conversion 600 mg q 6 hr for 3 doses, then 600 mg q 12 hr; discontinue lidocaine infusion at the time of the second oral dose of tocainide.132 Initiate tocainide during hospitalization.

Special Populations. Pediatric Dosage. Safety and efficacy not established.

Geriatric Dosage. Same as adult dosage.

Other Conditions. Reduce initial maintenance dosage by 50% in severe liver disease, by 25% in patients with Clcr 10–30 mL/min, and by 50% in patients with Clcr <10 mL/min.133 Dosages might have to be reduced slightly in CHF, but more data are needed.134

Dosage Forms. Tab 400, 600 mg.

Patient Instructions. Report any symptoms of numbness, drowsiness, dizziness, or tingling. Nausea or loss of appetite can occur and reduced by taking the drug with food. Report sore throat, mouth sores, fever, or abnormal bruising.

Missed Doses. Take this drug at regular intervals. If you miss a dose, take it as soon as you remember. If it is about time for the next dose, take that dose only. Leave at least 4–6 hr between doses. Do not double the dose or take extra.

Pharmacokinetics. Onset and Duration. PO onset 1–2 hr (delayed by food); duration 12–24 hr.

Serum Levels. 3–10 mg/L (16–52 μmol/L), although not well correlated with therapeutic or toxic effects.42,135

Fate. Oral bioavailability is 89 ± 5% with negligible first-pass metabolism.133–135 The drug is 10 ± 15% bound to plasma proteins.10,136 Vd is 3 ± 0.2 L/kg but slightly lower in CHF;133,134 Cl is 0.16 ± 0.03 L/hr/kg;10 38 ± 7% of the drug is excreted unchanged in urine, and 50–60% is hepatically eliminated.133,135 Renal clearance depends on urine pH; hepatic metabolism is stereospecific, with the (S)-enantiomer eliminated more quickly.137

t½α phase 5–10 min;136 β phase 13.5 ± 2.3 hr, 14–19 hr with ventricular arrhythmia or CHF;135 and 22 ± 3.1 hr in severe renal insufficiency.138

Adverse Reactions. Neurologic toxicities, which include dizziness, tremor, ataxia, drowsiness, confusion, and paresthesias, are frequent (30–50%); psychosis and seizures occur occasionally. The neurologic toxicities of lidocaine and tocainide can be additive.139 Nausea, vomiting, and anorexia occur frequently. Tocainide can exacerbate underlying ventricular arrhythmias or conduction distur-
bances. Agranulocytosis and other forms of bone marrow depression have been reported in up to 0.18% of patients.\textsuperscript{135} Pulmonary fibrosis or interstitial pneumonitis occurs in 0.03–0.11% of patients.\textsuperscript{132} Rash and fever occur occasionally, and cross-sensitivity between lidocaine and tocainide is possible.\textsuperscript{86}

**Contraindications.** Second- or third-degree AV block without a ventricular pacemaker.

**Precautions.** Sick sinus syndrome and CHF can worsen.

**Drug Interactions.** Cimetidine can decrease tocainide serum levels; rifampin can decrease levels.

**Parameters to Monitor.** Monitor ECG daily for 2–4 days when therapy is initiated and then q 3–6 months on an ambulatory basis. Obtain periodic serum levels once an individual’s effective level is determined. Monitor closely for neurologic toxicities when initiating therapy. Monitor WBC counts frequently, particularly during the first 3 months of therapy.\textsuperscript{135} Obtain baseline chest x-ray; repeat if pulmonary symptoms arise.

**Notes.** Because of reports of bone marrow toxicity, pulmonary fibrosis, and hypersensitivity reactions, the indications for tocainide are restricted to patients with life-threatening ventricular arrhythmias.
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<tr>
<td>Tocainide</td>
<td>0</td>
<td>↓</td>
<td>↓</td>
<td>0</td>
</tr>
<tr>
<td>IC (SLOW ON-OFF SODIUM-CHANNEL BLOCKERS)</td>
<td></td>
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<tr>
<td>Flecainide</td>
<td>↓↓↓</td>
<td>0</td>
<td>↓</td>
<td>0</td>
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<tr>
<td>Moricizine</td>
<td>↓↓↓</td>
<td>↓/0</td>
<td>↓</td>
<td>0</td>
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<tr>
<td>Propafenone</td>
<td>↓↓↓</td>
<td>0</td>
<td>↓</td>
<td>0</td>
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<tr>
<td>II (β-BLOCKERS)</td>
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<tr>
<td>Propranolol</td>
<td>↓</td>
<td>↓ (acute)</td>
<td>↓</td>
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<td></td>
<td>↑ (chronic)</td>
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(continued)
### ELECTROPHYSIOLOGIC ACTIONS OF ANTIARRHYTHMICS COMPARISON CHART (continued)

<table>
<thead>
<tr>
<th>CLASS AND DRUG</th>
<th>CONDUCTION VELOCITY</th>
<th>REFRACTORY PERIOD</th>
<th>AUTOMATICITY</th>
<th>AV NODAL CONDUCTION</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>III (POTASSIUM-CHANNEL BLOCKERS)</strong></td>
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<tr>
<td>Amiodarone&lt;sup&gt;d,f&lt;/sup&gt;</td>
<td>0/↓</td>
<td>↑↑</td>
<td>0</td>
<td>↓</td>
</tr>
<tr>
<td>Bretylium</td>
<td>0</td>
<td>↑↑</td>
<td>↑/0</td>
<td>↑/0</td>
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<tr>
<td>Dofetilide</td>
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<tr>
<td>Ibutilide</td>
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<tr>
<td>Sotalol&lt;sup&gt;g&lt;/sup&gt;</td>
<td>0</td>
<td>↑↑</td>
<td>0</td>
<td>↓</td>
</tr>
<tr>
<td><strong>IV (CALCIUM-CHANNEL BLOCKERS)</strong></td>
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<tr>
<td>Diltiazem</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>↓</td>
</tr>
<tr>
<td>Verapamil</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>↓↓</td>
</tr>
</tbody>
</table>

↑ = increase, ↓ = decrease, 0 = minimal or no effect, ↑/↓ = variable.

<sup>a</sup>Classification system from references 140 and 141.

<sup>b</sup>Classification of moricizine is controversial; it also has type Ia characteristics.

<sup>c</sup>Amiodarone, propafenone, and sotalol also have type II or β-adrenergic-blocking activity.

<sup>d</sup>Most investigational antiarrhythmics are potassium-channel blockers, many of which are analogues of sotalol.

<sup>e</sup>Most investigational antiarrhythmics are potassium-channel blockers, many of which are analogues of sotalol.

<sup>f</sup>Amiodarone also has type Ib sodium-channel-blocking activity.

<sup>g</sup>Amiodarone also has type Ib sodium-channel-blocking activity.

<sup>h</sup>Amiodarone also has type Ib sodium-channel-blocking activity.
Antihypertensive Drugs

Class Instructions. Antihypertensives. This medication can control but not cure hypertension. Long-term treatment is necessary to control hypertension and prevent damage to several body systems. Do not start or stop taking medications or change the dosage without medical supervision and avoid running out of medications. Some prescription and nonprescription medications can interact with medications for hypertension; make sure that your physician and pharmacist know the names of any other medications that you are taking.

Missed Doses. Take this drug at regular intervals. If you miss a dose, take it as soon as you remember. If it is about time for the next dose, take that dose only. Do not double the dose or take extra.

Pharmacology. Doxazosin, prazosin, and terazosin are closely related quinazoline derivatives that selectively block postsynaptic $\alpha_1$-adrenergic receptors. Total peripheral resistance is reduced through arterial and venous dilatations. Reflex tachycardia that occurs with other vasodilators is infrequent because there is no presynaptic $\alpha_2$-receptor blockade. The drugs also decrease total cholesterol, increase HDL-c, and may improve glucose tolerance and reduce left ventricular mass during long-term therapy. They increase urine flow in BPH by relaxing smooth muscle tone in the bladder neck and prostate.$^{142,143}$

Administration and Adult Dosage. Give the initial dose and the first dose of all increased dosage regimens at bedtime and observe the patient closely for syncope.

PO for hypertension (doxazosin) 1 mg/day initially and then double the dose at 1- to 2-week intervals to a maximum of 16 mg/day in a single dose, although dosages over 4 mg/day are more likely to cause postural side effects. (Prazosin) 1 mg bid or tid initially, increasing the dosage slowly, based on response, to the usual dosage of 6–15 mg/day; although the maximum effective dosage is usually 20 mg/day, dosages up to 40 mg/day can be effective in some patients who fail to respond to lower dosages. (Terazosin) 1 mg/day initially, increasing to 2, 5, or 10 mg/day in 1–2 doses to a maximum of 20 mg/day. PO for benign prostatic hypertrophy (doxazosin) 1 mg/day initially, doubling the dose at 1- to 2-week intervals to a maximum of 8 mg/day. (Terazosin) 1 mg/day initially increasing to 2, 5, and 10 mg once daily.

Special Populations. Pediatric Dosage. PO for hypertension (prazosin) 0.05–0.4 mg/kg/day in 2–3 divided doses. Do not exceed single doses of 7 mg and a total daily dosage of 15 mg.$^{144}$ (Doxazosin, terazosin) safety and efficacy not established.

Geriatric Dosage. Same as adult dosage.
Dosage Forms. (Doxazosin) Tab 1, 2, 4, 8 mg. (Prazosin) Cap 1, 2, 5 mg; Cap 1, 2, 5 mg with polythiazide 0.5 mg (Minizide). (Terazosin) Cap 1, 2, 5, 10 mg.

Patient Instructions. (See Antihypertensives Class Instructions.) Take the initial dose of this drug at bedtime. Dizziness or drowsiness can occur with this medication, especially after the first dose or when the dosage is being increased. Do not arise suddenly, stand for long periods, or exercise too vigorously, especially in hot weather. Alcohol can worsen these effects.

Pharmacokinetics. Onset and Duration. (Doxazosin) onset 1–2 hr, duration 24 hr for hypertension; full effect for BPH might not occur for 1–2 weeks. (Prazosin) onset 1–2 hr, duration about 6–12 hr, up to 4–6 weeks might be required for full antihypertensive effect. (Terazosin) onset 15 min, duration 24 hr, but up to 6–8 weeks might be required for full antihypertensive effect. In BPH, at least 4–6 weeks might be required to fully evaluate response to a 10 mg/day dosage. (See α₅-Adrenergic–Blocking Drugs Comparison Chart.)

Serum Levels. No correlation between serum levels and clinical effect has been established.¹⁰,¹⁴³

Fate. (Doxazosin) oral bioavailability is 63 ± 14%; absorption is slowed, but bioavailability is not affected by food; 98–99% is bound to plasma proteins. Vₐ is 1.5 ± 0.3 L/kg; Cl is 0.1 ± 0.024 L/kg/hr. Extensively metabolized and excreted primarily in the feces, with only about 9% excreted in urine as unchanged drug and metabolites.¹⁰,¹⁴³ (Prazosin) bioavailability is 68 ± 17%, 48 ± 16% in the elderly; food can delay but not affect the extent of absorption. About 95% is bound to plasma proteins, decreased in cirrhosis and uremia. Vₐ is 0.63 ± 0.14 L/kg and Cl is 0.24 ± 0.04 L/hr/kg in young patients; Vₐ is 0.89 ± 0.26 L/kg and Cl is 0.21 ± 0.06 L/hr/kg in the elderly; Cl is lower in CHF and pregnancy. Prazosin is metabolized in the liver by demethylation and conjugation; metabolites have about 20% of the activity of the drug. It is excreted renally as metabolites and 3.4% or less as unchanged drug.¹⁴⁵ (Terazosin) pharmacokinetics do not appear to be affected by uremia, CHF, or aging. Oral bioavailability is about 90%; 90–94% is bound to plasma proteins. Vₐ is 0.8 ± 0.18 L/kg, and Cl is 0.066 ± 0.012 L/hr/kg.¹⁰ It is extensively metabolized in the liver, with 18% excreted unchanged in feces, 10% unchanged in urine, and the remainder excreted as metabolites.¹⁴⁶

t₁/₂. (Doxazosin) 10.5 ± 2.4 hr in young adults, 11.9 ± 4.7 hr in the elderly. (Prazosin) 2.1 ± 0.3 hr in young adults, 3.2 ± 0.6 hr in the elderly; also prolonged in CHF and pregnancy. (Terazosin) 13.5 ± 3.5 hr in young adults, 16.2 ± 2.2 hr in the elderly.¹⁴⁷

Adverse Reactions. The most important adverse effect is first-dose syncope, which is more likely in patients being treated with other antihypertensive drugs, especially diuretics. During long-term treatment, the most frequent reactions are dizziness, headache, drowsiness, lack of energy or weakness, palpitations, or nausea, all of which occur in 5–20% of patients. Occasionally reported are rash, vomiting, diarrhea, edema, orthostatic hypotension, syncope, dyspnea, blurred vision, nasal congestion, or urinary frequency. Rarely, allergic reactions, priapism, or impotence occur.¹⁴⁸

Contraindications. Allergy to a quinazoline derivative.
**Precautions.** Syncope can occur after the first dose (doxazosin, 2–6 hr; prazosin, 30–90 min; terazosin, 1–2 hr) and during rapid upward dosage titration or when adding an additional antihypertensive drug. Hold doses of diuretics for 1 day before starting an $\alpha_1$-blocker. Increase dosage gradually, reduce dosage when adding another antihypertensive, and then retitrate dosage. Use doxazosin with caution in patients with hepatic impairment.

**Drug Interactions.** $\beta$-Blockers and verapamil can enhance postural effects of prazosin; NSAIDs can decrease the hypotensive effect of prazosin. The $\alpha_1$-blockers can decrease the hypotensive effect of clonidine.

**Parameters to Monitor.** Monitor blood pressure regularly.

**Notes.** $\alpha_1$-Antagonists can be particularly useful for hypertension in men with BPH, in those with hyperlipidemia or renal disease, in diabetics, in physically active young patients (no decrease in cardiac output), and in the elderly. However, drugs in this class have not been shown to decrease long-term mortality of hypertension. The doxazosin arm of the ALLHAT study was terminated prematurely because of inferior efficacy in reducing cardiovascular events compared with chlorthalidone. Tamsulosin (Flomax) is a selective $\alpha_{1a}$-receptor blocker, specific for adrenoreceptors in the prostate. Tamsulosin is not indicated for hypertension but rather for signs and symptoms of BPH. The initial oral dosage is 0.4 mg/day, increasing as needed up to 0.8 mg/day.

**CAPTOPRIL**

**Pharmacology.** Captopril is an ACE inhibitor pharmacologically similar to enalapril. Captopril’s rapid onset and short duration of action are advantageous initially to assess patient tolerance to ACE inhibitors but inconvenient during long-term use. (See ACE Inhibitors Comparison Chart.)

**Adult Dosage.** PO for hypertension 12.5–25 mg bid–tid initially, increasing after 1–2 weeks to 50 mg bid–tid, to a maximum of 450 mg/day. PO for CHF 6.25–25 mg tid initially, increasing over several days based on the patient’s tolerance to a dosage of 50 mg tid. Delay further dosage increases, if possible, for at least 2 weeks to evaluate response. Most patients respond to 50–100 mg tid. For hypertension or CHF use initial dosages of 6.25–12.5 mg bid–tid and increase slowly in patients on diuretic therapy, with sodium restriction, or with renal impairment. PO for left ventricular dysfunction post-MI 6.25 mg once at 3 or more days post-MI and then 12.5 mg tid; increase to 25 mg tid over several days to a target of 50 mg tid over several weeks as tolerated. PO for diabetic nephropathy 25 mg tid.

**Pediatric Dosage.** PO for hypertension (neonates) 0.01 mg/kg bid–tid initially; (children) up to 0.3 mg/kg tid initially.

**Dosage Forms.** Tab 12.5, 25, 50, and 100 mg, and 25 or 50 mg in combination with hydrochlorothiazide 15 or 25 mg (Capozide, various).

**Pharmacokinetics.** Oral bioavailability is about 65%; food decreases absorption, so the drug should be taken on an empty stomach. About 30% is bound to plasma proteins and its $V_d$ is 0.8 ± 0.2 L/kg, higher in CHF; Cl is 0.72 ± 0.08 L/hr/kg decreased with renal dysfunction. Approximately 50% of a dose is metabolized,
primarily to captopril disulfide, which can be converted back to active captopril in vivo. Urinary excretion of unchanged captopril is 24–38% over 24 hr. Its half-life is 2.2 ± 0.05 hr in healthy subjects and is prolonged in renal dysfunction or CHF.10

Adverse Reactions. Adverse reactions are similar to those of enalapril, although skin rashes and taste impairment can be more prevalent and cough less prevalent.

**CLONIDINE HYDROCHLORIDE**

**Pharmacology.** Clonidine stimulates postsynaptic α2-adrenergic receptors in the CNS by activating inhibitory neurons to decrease sympathetic outflow. Clonidine is not a complete agonist, so some of its effects might result from antagonist actions at presynaptic α-receptors.150 These actions reduce peripheral vascular resistance, renal vascular resistance, heart rate, and blood pressure.

**Administration and Adult Dosage.** PO for hypertension 0.1 mg bid initially, increasing weekly in increments of 0.1 mg/day until the desired response is achieved. Maintenance dosage for monotherapy is usually 0.2–0.6 mg/day, to a maximum of 2.4 mg/day. If rapid lowering of blood pressure is desired (eg, hypertensive urgency), give 0.1–0.2 mg initially and then 0.1 mg q 1 hr until the desired response is achieved or a total of 0.8 mg has been given. SR patch for hypertension initially apply one #1 (0.1 mg/24 hr) patch weekly; dosage can be increased at 1- to 2-week intervals up to a #3 patch that delivers 0.3 mg/24 hr. Dosages in excess of two #3 patches/week do not add efficacy. PO for opiate withdrawal 1.25–1.5 mg/day in 3–4 divided doses and then decreasing over 14 days by 0.1–0.2 mg/day.151 PO for smoking cessation 0.15–0.675 mg/day in divided doses. SR patch for smoking cessation apply one #1 (0.1 mg/24 hr) patch weekly.152 (See Notes.)

**Special Populations.** Pediatric Dosage. PO for hypertension 0.05–0.4 mg bid.

Geriatric Dosage. Lower oral dosages might be required, but decreased skin permeability might require higher transdermal dosages.153

Other Conditions. In renal impairment, lower oral dosages might be required, but decreased skin permeability might require higher transdermal dosages.153

**Dosage Forms.** Tab 0.1, 0.2, 0.3 mg; Tab 0.1, 0.2, 0.3 mg with chlorthalidone 15 mg (Combipres, various); SR Patch 0.1, 0.2, 0.3 mg/24 hr.

**Patient Instructions.** (See Antihypertensives Class Instructions.) Do not abruptly discontinue this drug or interrupt therapy unless under medical supervision. Apply transdermal patches weekly to a clean, hairless area of the upper arm or torso that is free of irritation, abrasions, or scars. Do not touch the adhesive surface. Apply patch to a different location with each application. If the system loosens during the 7 days, apply the adhesive overlay directly over the system. If a generalized rash or moderate to severe redness or vesicles appear at the site of application, notify the prescriber. Dispose of the patch by folding the sides together and placing it in a disposal site inaccessible to children.

**Missed Doses.** Take this drug at regular intervals. If you miss a dose, take it as soon as you remember. If it is about time for your next dose, take that dose only. Do not double the dose or take extra. Contact your physician if you miss two or
more doses or if you are late in changing the transdermal system by 3 or more days.

**Pharmacokinetics. Onset and Duration.** (Hypertension) PO onset 30–60 min; peak 2–5 hr; duration 6–8 hr but can increase to 12–24 hr with long-term use. Transdermally, maximal reduction in blood pressure occurs in 2–3 days and persists throughout the 7-day application period. After removal, blood pressure rapidly increases toward baseline, followed by a slower rate of increase, and returns to pretreatment levels over several days.

**Serum Levels.** (Hypotensive effect) 0.2–2 μg/L (0.9–9 nmol/L); (dry mouth, sedation) 1.5–2 μg/L.

**Fate.** Oral bioavailability is 75–95%. Transdermally, maximum serum levels are reached in 3–4 days and remain constant throughout the 7-day application period. Rate of release is a zero-order process and primarily controlled by the delivery system. Serum concentrations remain constant when a patch is removed and another is immediately applied to a different site. Clonidine is 20% bound to plasma proteins; $V_d$ is 2.1 ± 0.4 L/kg; $Cl$ is 0.186 ± 0.072 L/hr/kg. It is metabolized in the liver, with drug and metabolites excreted primarily in urine; remaining drug may undergo enterohepatic recycling. About 62% is excreted unchanged in urine.

$t_{1/2}$. (PO) $\alpha$ phase 10.8 ± 4.7 min; $\beta$ phase 12 ± 7 hr. (Transdermal) 14 hr but can be up to 26 hr, reflecting continued absorption from a skin depot.

**Adverse Reactions.** Frequent adverse reactions include dry mouth (40%), drowsiness (33%), dizziness (16%), constipation (10%), weakness (10%), sedation (10%), nausea or vomiting (5%), nervousness and agitation (3%), orthostatic hypotension (3%), and sexual dysfunction (3%). Occasionally rash, weight gain, anorexia, transient abnormalities in liver function tests, insomnia or vivid dreams, palpitations, tachycardia or bradycardia, or urinary retention occur. Rarely hepatitis, thrombocytopenia, parotitis, elevations of blood glucose or CPK, or cardiac conduction disturbances occur. Allergic contact dermatitis occurs in up to 50% of patients treated with patches. Abrupt withdrawal of oral therapy can result in a withdrawal reaction characterized by rapid reversal of the antihypertensive effect within 24–48 hr up to or above pretreatment levels, a rise of blood pressure above 40 mm Hg systolic or 25 mm Hg diastolic, or blood pressure above 225/125 mm Hg. Subjective symptoms of sweating, palpitations, anxiety, and insomnia also can occur, even without marked blood pressure changes. The frequency and severity of symptoms appear to be greater in patients treated with high dosages for more than 3 months and in those with more severe hypertension.

**Precautions.** Use with caution in patients with severe coronary insufficiency, conduction disturbances, recent MI, cerebrovascular disease, or chronic renal failure. Patients who develop rashes from the transdermal system can develop generalized skin rashes if oral clonidine is substituted. Inadvertent person-to-person transfer of the patches has been reported; check the application site frequently and dispose of the patch by folding adhesive sides together and placing it in a container inaccessible to children.
Drug Interactions. Tricyclic antidepressants can decrease the hypotensive effect of clonidine. Clonidine can inhibit the antiparkinson effect of levodopa. Clonidine use with propranolol can cause hypertension, especially if clonidine is abruptly discontinued. Direct-acting sympathomimetics can have an exaggerated effect during clonidine use. Prazosin can decrease the effects of clonidine. Synergistic hypotension and conduction disturbances can occur with verapamil.

Parameters to Monitor. Monitor blood pressure regularly; check patient compliance.

Notes. Clonidine has been used to suppress symptoms of withdrawal of opiates and to reduce craving and other symptoms in alcohol and tobacco withdrawal. It has also been used in a variety of psychiatric applications including treatment of mania, anxiety, panic disorders, schizophrenia, and antipsychotic-induced tardive dyskinesia. As an aid in the diagnosis of pheochromocytoma, a single 0.3 mg dose has been administered after determination of baseline catecholamine levels, followed by three subsequent determinations at hourly intervals. Other conditions for which it can be effective include diabetic diarrhea (0.1–0.6 mg q 12 hr), menopausal flushing (0.05–0.15 mg/day in divided doses), and premenstrual syndrome.

Pharmacology. Diazoxide is a nondiuretic thiazide that reduces total peripheral resistance by direct relaxation of arteriolar smooth muscle. It also increases heart rate, cardiac output, and renal blood flow. Diazoxide increases blood glucose by inhibiting insulin release and peripheral utilization.

Adult Dosage. IV for severe hypertension 1–3 mg/kg, to a maximum single dose of 150 mg administered undiluted over less than 30 sec q 5–15 min, until adequate blood pressure reduction is achieved. Repeat q 4–24 hr as needed to maintain blood pressure control to a maximum of 10 days. PO for hypoglycemia 3–8 mg/kg/day in 2–3 equal doses q 8–12 hr, titrated to response.

Pediatric Dosage. IV for severe hypertension same as adult dosage. PO for hypoglycemia (neonates and infants) 8–15 mg/kg/day in 2–3 divided doses q 8–12 hr, titrated to response; (children) same as adult dosage.

Dosage Forms. Inj 15 mg/mL (Hyperstat I.V.); Cap 50 mg (Proglycem); Susp 50 mg/mL (Proglycem).

Pharmacokinetics. Antihypertensive onset is 1–4 min; peak within 5 min; duration is 3–12 hr. Hyperglycemia onset within 1 hr; duration 8 hr. Oral bioavailability is 86–96%; 94 ± 14% is bound to plasma proteins at typical concentrations, decreased at higher concentrations and in uremia. \( V_d \) is 0.21 ± 0.02 L/kg with normal renal function; \( C_l \) is 0.0036 ± 0.0012 L/hr/kg. The drug is metabolized by oxidation and sulfate conjugation and excreted slowly in urine as unchanged drug (20–50%) and metabolites. Half-life is 48 ± 12 hr, prolonged in renal failure in proportion to \( C_l \).

Adverse Reactions. (Hypertension) hypotension, nausea and vomiting, dizziness, and weakness are the most frequent reactions. Sodium and water retention and hyperglycemia can occur, especially with repeated administration. (Hypoglycemia)
frequent reactions include sodium and fluid retention; hyperglycemia or glycosuria, which might require dosage reduction; hirsutism; tachycardia; palpitations; increases in uric acid; thrombocytopenia with or without purpura, which requires discontinuation of the drug. Rarely, diabetic ketoacidosis or hyperosmolar, nonketotic coma can develop rapidly.

**Contraindications.** Hypersensitivity to thiazides or other sulfonamide derivatives; compensatory hypertension, such as that seen secondary to coarctation of the aorta or arteriovenous shunts; functional hypoglycemia; dissecting aortic aneurysm.

**Precautions.** Use with caution with impaired cerebral or cardiac circulation. Avoid extravasation of the IV drug. Recent or co-administration of other antihypertensive drugs can produce excessive blood pressure reduction with the IV route.

**Drug Interactions.** Diazoxide and hydantoins can be mutually antagonistic. Use with a thiazide diuretic can potentiate hyperuricemia and hypotensive effects. Phenothiazines can potentiate the effects of oral diazoxide. Diazoxide can antagonize the effects of sulfonyleureas.

**Parameters to Monitor.** (Hypertension) Obtain blood pressure frequently until stable and then hourly; monitor blood glucose and uric acid with repeated doses. Monitor for signs of cerebral or myocardial ischemia. (Hypoglycemia) Obtain frequent blood glucose and urine glucose and ketones initially, when dosage adjustments or dosage form changes are made, and then regularly during stabilization.

**ENALAPRIL MALEATE**

Vasotec

**ENALAPRILAT**

Vasotec I.V.

**Pharmacology.** Enalapril is a prodrug that is rapidly converted to its active metabolite, enalaprilat, by ester hydrolysis in the liver. Enalaprilat is a competitive ACE inhibitor. It also reduces serum aldosterone, leading to decreased sodium retention, potentiates the vasodilator kallikrein–kinin system, and can alter prostanoid metabolism, inhibit the sympathetic nervous system, and inhibit the tissue renin–angiotensin system. The net effect is reduction in total peripheral resistance and blood pressure in hypertensive patients, especially those with high pre-treatment plasma renin activity and increased renal plasma flow, and reduction of elevated afterload in patients with CHF.

**Administration and Adult Dosage.** PO for hypertension 5 mg/day initially. Usual maintenance dosage is 10–40 mg/day in 1–2 doses. If the patient has recently been receiving a diuretic, discontinue the diuretic for 2–3 days or start with a lower initial enalapril dose of 2.5 mg; bid administration might be necessary in some individuals to achieve adequate 24-hr blood pressure control. A diuretic can be added if blood pressure control is inadequate with enalapril monotherapy. PO for CHF 2.5 mg daily or bid initially, using the lower dosage for patients taking a diuretic. Usual maintenance dosage is 5–20 mg/day, to a maximum of 40 mg; bid administration is preferred. IV for hypertension 1.25 mg (0.625 mg initially if patient is taking a diuretic) over 5 min q 6 hr. Dosages as high as 5 mg q 6 hr can be tolerated for up to 36 hr, but there is inadequate experience with dosages...
over 20 mg/day. For patients converting from PO to IV, 5 mg/day PO is about equivalent to 1.25 mg IV q 6 hr.

**Special Populations. Pediatric Dosage.** Safety and efficacy not established.

**Geriatric Dosage.** No change necessarily required but observe cautions for impaired renal function.

**Other Conditions.** For patients with Clcr $\leq 30$ mL/min, Cr $>3$ mg/dL, or CHF with serum sodium <130 mEq/L, use lower initial doses (2.5 mg PO; 0.625 mg IV). For patients on dialysis, the initial dose should be no greater than 0.625 mg IV q 6 hr or 2.5 mg PO on dialysis days.

**Dosage Forms.** **Tab** 2.5, 5, 10, 20 mg (Enalapril); **Tab** 5 mg with hydrochlorothiazide 12.5 mg, 10 mg with hydrochlorothiazide 25 mg (Vaseretic); **SR Tab** 5 mg with diltiazem 180 mg (Teczem), 5 mg with 2.5 or 5 mg felodipine (Lexxel); **Inj** 1.25 mg/mL (enalaprilat).

**Patient Instructions.** **(See Antihypertensives Class Instructions.)** Use potassium supplements or salt substitutes only under medical supervision. Report any signs or symptoms of the following: infection (eg, sore throat or fever), angioedema (eg, swelling of face, eyes, lips, tongue, larynx, extremities, or hoarseness or difficulty in swallowing), or excessive fluid loss (eg, vomiting, diarrhea, or excessive perspiration). Report any skin rash, taste disturbance, or persistent, dry cough. If you become pregnant while taking this drug, contact your prescriber immediately.

**Pharmacokinetics. Onset and Duration.** PO onset is 1 hr; peak in 4–6 hr; duration is up to 24 hr.$^{164}$ The onset of action and maximal hemodynamic response correspond to the appearance of enalaprilat in serum.$^{10}$ IV onset is 15–30 min; peak is within 1 hr; duration is usually 4–6 hr with recommended doses but can be as long as 12 hr in some patients.$^{165}$

**Serum Levels.** (Enalaprilat) 5–20 μg/L (13–52 nmol/L) is the EC$_{50}$ for ACE inhibition; 40 μg/L (104 nmol/L) produces a mean blood pressure reduction of 12 mm Hg.$^{10,166}$

**Fate.** Oral bioavailability is 41 ± 15%; it is not altered by meals but is decreased in cirrhosis.$^{10}$ Peak enalapril and enalaprilat serum levels after a 10 mg oral dose occur at about 1 and 4 hr, with ranges of 40–50 μg/L (104–130 nmol/L) and 30–40 μg/L (78–104 nmol/L), respectively.$^{166}$ About 60% of a dose is converted to enalaprilat; conversion can be reduced in patients with cirrhosis.$^{166}$ Enalapril and enalaprilat levels are increased in renal dysfunction. Less than 50% of enalaprilat is bound to plasma protein.$^{166}$ $V_{d0}$ is 1.7 ± 0.7 L/kg; $Cl$ is 0.294 ± 0.09 L/hr/kg.$^{167}$ Cl is decreased in uremia, CHF, the elderly, and neonates.$^{10}$ After IV administration, 88% is excreted unchanged in urine,$^{10}$ after oral administration, 33% of the dose is recovered in the feces (6% as enalapril, 27% as enalaprilat) and 61% in the urine (18% as enalapril, 43% as enalaprilat).$^{166,168}$ Enalapril can be actively secreted into the urine; fecal recovery can indicate unabsorbed drug or biliary excretion.$^{166}$

$t_{1/2}$. (Enalapril) estimated to be 11 hr; (enalaprilat) about 30–35 hr in normals, increased in CHF, renal dysfunction, cirrhosis, and uremia.$^{10,166}$
Adverse Reactions. ACE inhibitors have a common side effect profile. Most adverse effects are related to dosage and renal function. A dry, nonproductive cough occurs in 1–3% or more (up to 20% in some surveys) of treated patients, most frequently in women and nonsmokers.169 The cough is caused by potentiation of tissue kinins or prostaglandins in the lung. It can be more frequent with longer-acting drugs but is usually not resolved by switching to another ACE inhibitor. Taste disturbances occur in 2–7% but can resolve despite continued therapy.169,170 Skin rashes occur in 1–7%, usually within a few days to weeks after starting.170 Rashes often resolve with continued therapy and do not appear to cross-react among ACE inhibitors.169 Angioedema is an occasional, serious, potentially fatal reaction, possibly more frequent with longer-acting ACE inhibitors and possibly in blacks.163,169 Hypotension can occur, especially with the first dose, in vigorously diuresed patients, those who are hypovolemic or hypovolemic, those with severe hypertension, and the elderly. In salt-restricted patients with CHF receiving ACE inhibitors and continuous diuretic therapy, up to one-third can experience worsening of renal function that can improve when sodium is replenished.170 Hyperkalemia occurs in 1–4% of patients, most often in those with diabetes mellitus or renal dysfunction. Proteinuria occurs occasionally with normal renal function and frequently with pre-existing renal disease,163 although patients with progressive renal insufficiency tolerate the drug well and many experience a reduction in proteinuria despite transient reductions in renal function.171 Neutropenia can occur, usually in the first 3 months of therapy; it is rare in normal patients but more frequent with high doses or in renal impairment.169 Cholestatic hepatotoxicity is reported rarely and it can cross-react among ACE inhibitors;169 it is reversible with drug discontinuation, but fatalities have been reported. Serious fetal harm, including renal failure, face or skull abnormalities, and increased risk of miscarriage, occurs with ACE-inhibitor use during the second and third trimesters of pregnancy.169

Contraindications. Angioedema caused by any ACE inhibitor.

Precautions. Pregnancy. It is best to avoid ACE inhibitors in women of childbearing potential who are not actively avoiding pregnancy. Monitor patients on dietary salt restriction, diuretic therapy, or dialysis (salt or volume depletion) for hypotensive episodes after the initial dose. If possible, discontinue these therapies before treatment. Titrate dosage slowly to the minimal effective dosage in patients with impaired renal function or collagen vascular disorders or in patients receiving drugs altering WBC count or immune function.171,172 Patients with aortic stenosis can develop decreased coronary perfusion when treated with afterload reducers such as ACE inhibitors. Elevations in Cr, and BUN might require dosage reduction or drug discontinuation. Patients with unilateral or bilateral renal artery stenosis might be more prone to increases in Cr, and BUN. Hypotension responsive to volume expansion can occur during surgical procedures.

Drug Interactions. Hyperkalemia can develop with concomitant use of potassium-sparing diuretics, potassium supplements, or potassium-containing salt substitutes, particularly with pre-existing renal impairment.169 Sodium and volume depletion because of a loop diuretic can cause postural hypotension when an ACE inhibitor is begun. ACE inhibitors can increase lithium levels. ACE inhibitors can
potentiate oral hypoglycemic drugs and increase neutropenia caused by azathioprine and hypotensive reactions when used with IV plasma protein solutions. NSAIDs can antagonize the hypotensive effect of ACE inhibitors. Phenothiazines can increase the effects of ACE inhibitors and rifampin can decrease the effects of enalapril. ACE inhibitors can increase serum digoxin concentrations.

**Parameters to Monitor.** Monitor blood pressure regularly. Obtain baseline Cr, and BUN to assess the potential for adverse effects and titrate dosages accordingly; then monitor periodically. Obtain WBC count with differential q 2 weeks for the first 3 months and then periodically in renally impaired patients or if signs of infection occur. Obtain baseline serum potassium and then monitor periodically, especially in patients receiving potassium-sparing diuretics, potassium supplements, or salt substitutes. Obtain periodic urinary protein estimates (morning urines) by dipstick in patients with renal impairment.

**Notes.** ACE inhibitors are considered first-line drugs, along with diuretics, β-blockers, and calcium-channel blockers, for the treatment of hypertension. They are also first-line treatments for CHF in combination with digoxin and a diuretic because their use is associated with prolonged survival. Regression or attenuation of left ventricular hypertrophy occurs in patients with hypertension and in post-MI patients. Additional advantages of ACE inhibitors are their renal protective effects and improved insulin sensitivity in type 1 diabetics, their lack of adverse effects on serum lipid profile, an improvement in quality of life in hypertensive patients (with one study favoring captopril over enalapril), and possibly prevention of structural changes in the heart, systemic vasculature, and kidneys. ACE inhibitors with greater tissue ACE inhibition (eg, benazepril, quinapril, ramipril) might be more effective in this latter regard, but studies are lacking. Ramipril reduces mortality and cardiovascular morbidity in patients without CHF who are at high risk for cardiovascular events. (See ACE Inhibitors Comparison Chart.)

**FENOLDOPAM**

**Pharmacology.** Fenoldopam is a dopamine D1-receptor agonist that dilates renal and mesenteric vascular beds, thereby reducing total peripheral resistance and increasing renal blood flow and sodium excretion. Stimulation of postsynaptic D1-receptors leads to smooth muscle relaxation through activation of adenylate cyclase and a subsequent increase in intracellular cyclic AMP. Unlike dopamine, fenoldopam has no α- or β-adrenergic receptor activity, stimulation of which causes increases in blood pressure or heart rate, respectively.

**Administration and Adult Dosage.** IV for the in-hospital, short-term (up to 48 hr) management of severe hypertension 0.03–0.1 μg/kg/min initially, increasing in increments of 0.05–0.1 μg/kg/min at intervals of >20 min to a maximum of 1.7 μg/kg/min. Do not use bolus injections. Lower initial infusion rates and slower titration result in less reflex tachycardia. When the desired effect is achieved, the infusion can be stopped gradually or abruptly because rebound elevation of blood pressure has not been observed.

**Special Populations.** Pediatric Dosage. Safety and efficacy not established.
**Geriatric Dosage.** Same as adult dosage.

**Other Conditions.** Dosage adjustments are not necessary for renal or hepatic disease or continuous ambulatory peritoneal dialysis. The effects of hemodialysis have not been evaluated.

**Dosage Forms.** **Inj** 10 mg/mL.

**Pharmacokinetics.** **Onset and Duration.** Onset <15 min; peak 2–6 hr. Blood pressure returns to baseline 2 hr after infusion discontinuation.\(^{178}\)

**Serum Levels.** Plasma fenoldopam concentrations of 3.5 \(\mu g/L\) are required for demonstrable reduction in blood pressure. Each 1 \(\mu g/L\) increase in plasma fenoldopam concentration causes a 0.8% decrease in diastolic blood pressure. A concentration of 18 \(\mu g/L\) is required for each 10 mm Hg reduction in diastolic blood pressure.\(^{177}\)

**Fate.** Fenoldopam has nonlinear increases in \(V_d\) with increases in dosage. \(V_d\) is 0.23, 0.66, and 0.67 L/kg at infusion rates of 0.025, 0.25, and 0.5 \(\mu g/kg/min\), respectively.\(^{177}\) Cl is dose dependent, increasing from 1.49 L/hr/kg at an infusion rate of 0.025 \(\mu g/kg/min\) to 2.29 L/hr/kg at a rate of 0.5 \(\mu g/kg/min\).\(^{179}\) Fenoldopam is about 88% bound to plasma proteins. Elimination is due primarily to conjugation to inactive metabolites. About 90% is excreted in the urine (4% unchanged), 10% in feces.

\(t_{1/2}\), 5–10 min.

**Adverse Reactions.** Fenoldopam causes dose-related reduction in blood pressure and reflex tachycardia; excessive decreases in blood pressure and vasodilation are responsible for most adverse effects. Frequent adverse effects are headache (11–36%), flushing (7–11%), nausea (about 20%), asymptomatic ST-segment abnormalities (6–33%), and hypotension (>5%).\(^{178}\) Most adverse events occur during the first 24 hr of therapy. Hypokalemia, elevated BUN, serum glucose, transaminase, and LDH have been reported in 0.5–5% of patients. Fenoldopam can cause reversible, dose-related increases in intraocular pressure.\(^{180}\)

**Contraindications.** None known.

**Precautions.** Use with caution in patients with glaucoma or intraocular hypertension. Fenoldopam causes hypotension and reflex tachycardia, which can lead to increased myocardial oxygen demand and possibly ischemia. Closely monitor patients with low serum potassium concentrations, especially during the first 6 hr of fenoldopam therapy.

**Drug Interactions.** IV allopurinol can attenuate fenoldopam-induced increases in renal blood flow. If possible, avoid concomitant use of \(\beta\)-blockers, which can cause excessive hypotension and inhibition of reflex responses to fenoldopam.

**Parameters to Monitor.** Monitor blood pressure and heart rate at least q 15 min because of the rapid onset and termination of effects. Monitor serum potassium frequently during fenoldopam therapy, especially during the first 24 hr.

**Notes.** Fenoldopam reduces blood pressure similar to nitroprusside.\(^{181–183}\) Fenoldopam might be preferred to nitroprusside in patients with renal dysfunction or requiring prolonged therapy due to the accumulation of thiocyanate with nitro-
Prepare the infusion solution with NS or D5W. It is stable for 24 hr under normal light and temperature conditions.

**Pharmacology.** Hydralazine is a vasodilator that reduces total peripheral resistance by direct action on vascular smooth muscle, with an effect greater on arterioles than on veins. (See Notes.)

**Administration and Adult Dosage.** PO for hypertension 10 mg qid for the first 2–4 days and increase to 25 mg qid for the remainder of the first week; after the first week, the dosage can be increased to 50 mg qid, to a maximum of 300 mg/day; bid administration can be as effective as qid. **PO for CHF** 50–75 mg bid–qid initially. **Usual maintenance dosage** 200–600 mg/day, but dosages as high as 3 g/day have been used.**IM or IV for hypertension and CHF** 10–40 mg prn.

**Special Populations. Pediatric Dosage.** PO for hypertension and CHF 0.75 mg/kg/day or 25 mg/m²/day initially in 4 divided doses; the initial dose should not exceed 25 mg. Increase gradually over 3–4 weeks, to a maximum of (infants) to 7.5 (children) mg/kg/day or 200 mg/day. **IM or IV for hypertension and CHF** 0.1–0.2 mg/kg q 4–6 hr prn; initial parenteral dosage should not exceed 20 mg.

**Geriatric Dosage.** Lower dosage and slower titration are desirable because of longer half-life in the elderly.

**Dosage Forms.** **Inj** 20 mg/mL; **Tab** 10, 25, 50, 100 mg; **Cap** 25 mg with hydrochlorothiazide 25 mg, 50 mg with hydrochlorothiazide 50 mg, 100 mg with hydrochlorothiazide 50 mg (Apresazide).

**Patient Instructions.** (See Antihypertensives Class Instructions.) This drug can cause headache, dizziness, or palpitations; report if these symptoms are persistent. Report symptoms of drug-induced SLE such as fever, joint pains, dermatitis, pleuritic chest pain, and generalized malaise.

**Pharmacokinetics. Onset and Duration.** PO onset in 1 hr; after 300 mg/day, a minimum of 30 hr is required for MAP to return to 50% of baseline value. IV onset is in 10–20 min, peak in 10–80 min; IM onset is in 10–30 min; duration for IV and IM is 3–8 hr.

**Serum Levels.** 100 µg/L reduces MAP by 10–20 mm Hg.**Fate.** Bioavailability is a function of acetylator phenotype and averages 35 ± 4% for slow acetylators and 16 ± 6% for rapid acetylators. Food can enhance the bioavailability; the first-pass effect might be saturable. Plasma protein binding is 87%. Vₐ is 1.5 ± 1 L/kg; Cl is 3.36 ± 0.78 L/hr/kg, reduced in CHF. The drug is metabolized extensively by acetylation to multiple metabolites, principally hydrazones, at a rate that is genetically determined; only 1–15% of unchanged drug, as well as metabolites, is excreted in the urine. t½. β phase 0.96 ± 0.28 hr, longer in CHF.

**Adverse Reactions.** Frequently, headache, anorexia, nausea, vomiting, diarrhea, palpitations, tachycardia, and angina occur. Occasionally, hypotension, edema, peripheral neuritis, dizziness, tremors, muscle cramps, urinary retention, nasal prusside.
congestion, and flushing occur. A syndrome similar to SLE with joint pain and skin rash (only rarely with cerebritis and nephritis) has been reported at an overall frequency of 6.7% in 281 patients over 51 months; daily dosage affects the frequency, with none at 50 mg/day, 5.4% at 100 mg/day, and 10.4% at 200 mg/day. Women had a higher overall frequency than men (11.6 and 2.8%, respectively), and women taking 200 mg/day had a 19.4% rate; slow acetylator phenotype also can increase the risk; the syndrome is reversible with drug discontinuation, although residual effects can be detected years later. An immune complex glomerulonephritis has been reported in patients with hydralazine-induced SLE.

Contraindications. Coronary artery disease, mitral valvular rheumatic disease.

Precautions. Reflex tachycardia can precipitate anginal attacks or ECG evidence of myocardial ischemia.

Drug Interactions. NSAIDs can antagonize the hypotensive effect of hydralazine.

Parameters to Monitor. Blood pressure and heart rate regularly. Baseline and periodic CBC. ANA titers can become positive after several months of therapy; routine monitoring is generally not warranted because the symptoms of hydralazine-induced SLE are characteristic and reversible with drug discontinuation.

Notes. Reflex increases in heart rate, cardiac output, and stroke volume and increases in plasma renin activity and retention of sodium and water can attenuate the antihypertensive action of hydralazine; therefore, long-term regimens for hypertension should include a diuretic and a sympatholytic drug. When hydralazine is used as an afterload-reducing drug in the treatment of CHF in patients on maintenance diuretics, the increase in cardiac output usually prevents the development of reflex tachycardia; likewise, hypotension is usually prevented by the increased cardiac output but can occur if myocardial reserves are inadequate or if the heart cannot respond by increasing output (eg, severe cardiomyopathy or aortic stenosis).

Pharmacology. Labetalol is an adrenergic receptor blocking drug that has selective $\alpha_1$- and nonselective $\beta$-adrenergic receptor blocking actions. Although its pharmacologic profile resembles that of other $\beta$-blockers and the postsynaptic $\alpha_1$-adrenergic blocking action of prazosin, its $\beta$-blocking activity is approximately 3 times greater than the $\alpha$-blocking activity after oral administration and 7 times greater after IV administration. During long-term treatment, $\alpha$-blocking activity is reduced even more.

Administration and Adult Dosage. PO for hypertension 100 mg bid initially, increasing at 2- to 3-day intervals in 100 mg bid increments until blood pressure is controlled. Usual maintenance dosage is 200–400 mg bid, to a maximum of 1.2–2.4 g/day for severe hypertension. IV for hypertension 20 mg by slow (2 min) injection, followed by 40–80 mg at 10-min intervals until blood pressure is controlled or to a total of 300 mg. Alternatively, administer a dilute solution by continuous infusion at a rate of 2 mg/min, to a maximum total dosage of 300 mg; the usual effective cumulative dosage is 50–200 mg; the infusion can be repeated q 6–8 hr.
**Special Populations. Pediatric Dosage.** Safety and efficacy not established, but the following has been used: IV for hypertension 0.2–1 (average 0.55) mg/kg initially, followed by a continuous infusion of 0.25–1.5 (average 0.8) mg/kg/hr.\(^\text{191}\)

**Geriatric Dosage.** PO Initiate therapy with 50 mg bid.\(^\text{189}\)

**Other Conditions.** Titrate dosage to blood pressure control. No dosage adjustment is required in renal impairment. Patients with hepatic dysfunction might require lower than usual dosages.

**Dosage Forms.** Tab 100, 200, 300 mg; Inj 5 mg/mL.

**Patient Instructions.** (See Antihypertensives Class Instructions.) Do not discontinue medication abruptly except under medical supervision. Do not sit up or stand for 3 hours after intravenous administration.

**Pharmacokinetics. Onset and Duration.** PO onset is within 2 hr, peak in 3 hr, and duration of 8–12 hr; can be longer with higher dosages. IV injection onset <10 min, peak in 5–15 min, duration 3–6 hr.\(^\text{173,190}\)

**Fate.** Almost completely absorbed, but bioavailability is only 18 ± 5% because of extensive first-pass metabolism, with the higher values reported in the elderly and patients with cirrhosis.\(^\text{10,192}\) Peak serum levels occur within 1–2 hr after oral administration; food delays the time to peak but can increase bioavailability. Plasma protein binding averages 50%. There is little distribution into the brain because of low lipid solubility. \(V_d\) is 9.4 ± 3.4 L/kg; \(Cl\) is 1.5 ± 0.6 L/hr/kg, lower in young hypertensive patients and the elderly and unchanged in cirrhosis. The drug is metabolized extensively primarily in the liver and possibly gut wall to inactive compounds. Unchanged drug (<5%) and metabolites are excreted in urine and feces.\(^\text{10,189,190,192}\)

\(t_{1/2}\) \(\beta\) phase 4.9 ± 2 hr, independent of route of administration; increased in the elderly.\(^\text{10,189,192}\)

**Adverse Reactions.** These are generally related to \(\alpha\)- and \(\beta\)-adrenergic blockade and usually occur during the first few weeks of therapy. Frequently, dizziness, fatigue, headache, scalp tingling, nausea, dyspepsia, and nasal congestion occur. Occasionally, postural hypotension, edema, taste disturbance, impotence, rash, and blurred vision occur. IV administration causes ventricular arrhythmias rarely.

**Contraindications.** Bronchial asthma; overt cardiac failure; greater than first-degree heart block; cardiogenic shock; bradycardia.

**Precautions.** Lower dosages might be required in patients with impaired hepatic function.

**Drug Interactions.** Cimetidine can increase the bioavailability of oral labetalol. Glutethimide can decrease the effect of labetalol by inducing hepatic enzymes. Concurrent use with halothane can produce myocardial depression. Labetalol decreases the reflex tachycardia induced by nitroglycerin and the bronchodilator effects of \(\beta_2\)-agonist bronchodilators.

**Parameters to Monitor.** Monitor blood pressure regularly and hepatic and renal function as indicated.
Notes. Labetalol injection is incompatible with 5% sodium bicarbonate, furosemide, or other alkaline products.

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Losartan is a selective, reversible, nonpeptide, competitive antagonist of the angiotensin II receptor (AT₁), which is responsible for the physiologic effects of angiotensin II including vasoconstriction, aldosterone secretion, sympathetic outflow, and stimulation of renal sodium reabsorption. Losartan and other angiotensin II receptor antagonists are highly selective for the AT₁ receptor over the AT₂ receptor, whose physiologic function is unknown. Angiotensin II receptor antagonists have no inhibitory effects on ACE and therefore decrease blood pressure with no appreciable effect on kinin metabolism.¹⁹³

Administration and Adult Dosage. PO for hypertension 50 mg/day initially; 25 mg/day in patients on diuretics or volume depleted. The usual dosage is 25–100 mg/day given without regard to meals once daily; may increase to bid in patients not adequately controlled with once-daily administrations. Most patients respond to 50 mg/day, although further reductions in blood pressure are possible with 100 mg/day.¹⁹⁴ Patients who do not respond to 50 mg/day might benefit more with the addition of hydrochlorothiazide than an increased dosage. Dosages above 100 mg/day offer little added benefit.¹⁹³

Special Populations. Pediatric Dosage. Safety and efficacy not established.

Geriatric Dosage. Same as adult dosage.

Other Conditions. No dosage adjustment is necessary in patients with renal impairment or on dialysis. Patients with hepatic insufficiency might require lower doses (eg, starting dose of 25 mg/day) because of decreased losartan clearance.

Dosage Forms. Tab 25, 50, 100 mg (Cozaar); Tab 50 mg with 12.5 mg hydrochlorothiazide, 100 mg with 25 mg hydrochlorothiazide (Hyzaar).

Patient Instructions. (See Antihypertensives Class Instructions.) This medication can cause dizziness, especially with the first few doses; do not drive or operate dangerous machinery until you know how you will react to this medicine. Do not use this medicine if you are pregnant or planning to become pregnant. If you become pregnant while taking this medicine, contact your prescriber immediately. Report any skin rash or signs or symptoms of angioedema (eg, swelling of face, eyes, lips, tongue, larynx, extremities, or hoarseness or difficulty in swallowing) immediately to your prescriber.

Pharmacokinetics. Onset and Duration. PO onset <2 hr; peak 6 hr; duration >24 hr, can be less with doses ≤25 mg/day.¹⁹⁵ Maximum antihypertensive effect occurs after 1 week in most patients but can take 3–6 weeks.

Serum Levels. Large interindividual variability, with IC₅₀ for AT₁ inhibition occurring at losartan concentrations of 1.4–200 nmol/L.¹⁹⁶

Fate. Oral absorption is rapid, but extensive first-pass metabolism results in a bioavailability of 33%, which might be doubled in hepatic insufficiency. About 14% of an oral dose is converted to an active carboxylic acid metabolite. Peak concentrations of losartan occur in 1 hr and those of its metabolite in 3–4 hr. The
metabolite is approximately 10–40 times more potent than the parent compound and is believed to be responsible for most of the antihypertensive effects of losartan. Losartan and its metabolite are about 99% bound to proteins, mainly to albumin. Vₘₕ of losartan and its active metabolite are 34 and 12 L, respectively. Metabolism of losartan occurs through CYP2C9 and CYP3A4 to the active carboxylic acid metabolite and several inactive metabolites. Cl is about 36 L/hr for losartan (12–15% renal Cl) and 3 L/hr for the active metabolite (50% renal Cl). Losartan Cl can be 50% less with hepatic insufficiency. About 4% of an oral losartan dose is excreted unchanged in the urine and 6% of the dose as active metabolite. After oral administration, 60% of a losartan dose is excreted in the feces.

\( t_{1/2} \) (Losartan) 2 hr; (metabolite) 6–9 hr.

**Adverse Reactions.** Angiotensin II receptor antagonists are generally well tolerated, with adverse reactions occurring at frequencies similar to those of placebo; adverse events are not related to dose. The most frequent reactions are headache (10–20%) and upper respiratory tract infection (1–12%). Nasal congestion, cough, and fatigue occur in fewer than 6% of patients. Unlike ACE inhibitors, angiotensin II receptor antagonists induce cough about as frequently as placebo, probably because bradykinin concentrations are not elevated as they are with ACE inhibitors. Angiotensin II receptor antagonists are effective alternatives in patients who experience cough with ACE inhibitors. Like ACE inhibitors, angiotensin II receptor antagonists can induce reversible renal dysfunction as a consequence of affecting the renin–angiotensin–aldosterone system. Increases in Crₗ and BUN also can occur in patients with unilateral or bilateral renal artery stenosis. Hypersensitivity reactions (eg, angioedema, rash) have been reported in patients receiving losartan or valsartan. Angiotensin II receptor antagonists can decrease hemoglobin and hematocrit and increase serum bilirubin, but these changes are rarely of clinical importance. Neutropenia has been reported in 1.8% of patients taking valsartan (0.9% for placebo). Hyperkalemia has been reported in 1.5% of losartan-treated patients (1.3% for ACE inhibitor) and 4.4% of valsartan-treated patients (2.9% for placebo).

**Contraindications.** Hypersensitivity to any product components.

**Precautions.** Use of drugs affecting the renin–angiotensin–aldosterone system can cause injury and even death to the developing fetus if used in the second or third trimester of pregnancy. Increase dosage slowly in patients with liver dysfunction because of reduced drug clearance (losartan, valsartan) in these patients. Patients taking angiotensin II receptor antagonists whose renal function is dependent on the renin–angiotensin–aldosterone system (eg, CHF patients) can experience oliguria, progressive azotemia, and (rarely) acute renal failure or death. Reversible increases in Crₗ and/or BUN can occur in patients with unilateral or bilateral renal artery stenosis.

**Drug Interactions.** Inhibitors of the CYP3A4 or 2C9 isoenzymes (eg, ketoconazole) can impair the conversion of losartan to the active metabolite. Telmisartan can increase digoxin serum concentrations. No important interactions have been reported with other drugs in this class.
Parameters to Monitor. Monitor for hypersensitivity reactions (e.g., flushing, dyspnea, facial swelling, rash) at the start of therapy. Monitor blood pressure regularly. Monitor patients on dietary salt restriction, diuretic therapy, or dialysis (salt or volume depletion) for hypotensive episodes after the initial dose. Obtain baseline and periodic Cr, and BUN to assess the potential for adverse effects. Obtain baseline serum potassium, WBC count, hemoglobin, and hematocrit. Monitor periodically for hyperkalemia, neutropenia, and anemia.

Notes. Although the guidelines of the sixth report by the Joint National Committee on Prevention, Detection, Evaluation, and Treatment do not promote this, many clinicians consider AT1 antagonists first-line therapy for hypertension because of their efficacy, safety, and ease of administration. Losartan is a uricosuric, which can lower plasma uric acid concentration and increase the risk of acute uric acid nephropathy or acute gout. Losartan has been shown to improve cardiac output and reduce peripheral vascular resistance and pulmonary capillary wedge pressure in patients with CHF. The ELITE II study found losartan to be comparable but not superior to captopril in improving survival in elderly patients with CHF, although this study was not designed to test equivalence. (See Angiotensin II Receptor Antagonists Comparison Chart.)

METHYLDOPA  Aldomet, Various
METHYLDOPATE HYDROCHLORIDE  Aldomet, Various

Pharmacology. The action of methyldopa is thought to be mediated through stimulation of central α-adrenergic receptors in a manner similar to that of clonidine. Stimulation is caused primarily by the metabolite α-methylnorepinephrine.

Administration and Adult Dosage. PO for hypertension 250 mg bid–tid initially, increasing at intervals of no less than 48 hr to the usual daily dosage of 500 mg–2 g/day in 2–4 divided doses. IV for hypertension usual dosage is 250–500 mg over 30–60 min in 100 mL D5W q 6 hr, to a maximum of 1 g q 6 hr.

Special Populations. Pediatric Dosage. PO 10 mg/kg/day in 2–4 doses initially, to a maximum of 65 mg/kg/day or 3 g/day, whichever is less. IV 20–40 mg/kg/day in divided doses q 6 hr, to a maximum of 65 mg/kg/day or 3 g/day, whichever is less.

Geriatric Dosage. Use lower dosages to avoid causing syncope.

Other Conditions. Patients with renal failure might respond to smaller dosages of methyldopa.

Dosage Forms. Tab 125, 250, 500 mg; Tab 250 mg with chlorothiazide 150 or 250 mg (Aldoclor); Tab 250 mg with hydrochlorothiazide 15, 25 mg, 500 mg with hydrochlorothiazide 30, 50 mg (Aldoril, various); Susp 50 mg/mL; IV 50 mg/mL.

Patient Instructions. (See Antihypertensives Class Instructions.) Report changes in mood (depression), loss of appetite, yellowing of eyes or skin, abdominal pain, or unexplained fever or joint pains. This drug can cause your urine to darken if it is exposed to air after voiding.
**Pharmacokinetics.** **Onset and Duration.** PO onset 2 hr, peak within 4–6 hr, duration 12–24 hr. IV onset 4–6 hr, duration 10–16 hr.

**Serum Levels.** No correlation between serum levels and therapeutic effect.

**Fate.** Oral bioavailability is 42 ± 16%. Peak serum levels occur in 2–4 hr but correlate poorly with the hypotensive effect. IV bioavailability is similar to oral, apparently because a large portion of methyldopate ester is not hydrolyzed to methyldopa. From 10 to 15% is bound to plasma proteins. Vd is 0.46 ± 0.15 L/kg; Cl is 0.22 ± 0.06 L/hr/kg and is decreased in uremia. The drug is excreted in the urine as metabolites, sulfate conjugate, and unchanged drug. About 49% (IV) and 70% (PO) of a dose are excreted in urine as sulfate conjugate and unchanged drug.

$\frac{t_1}{2}$: α phase 0.21 hr (range 0.16–0.26); β phase 1.8 ± 0.6 hr, increased in uremia and in neonates.

**Adverse Reactions.** Frequently, drowsiness, headache, weight gain, nasal stuffiness, postural hypotension, or dry mouth occur. A positive Coombs’ test develops in 10–20% of patients, usually between 6 and 12 months of therapy; hemolytic anemia is rare. Occasionally, depression, sexual dysfunction, diarrhea, or nightmares occur. Rarely, hepatitis, drug fever, lupus-like syndrome, leukopenia, thrombocytopenia, or granulocytopenia occur.

**Contraindications.** Active hepatic disease such as acute hepatitis and active cirrhosis or liver dysfunction associated with previous methyldopa therapy; concurrent MAOI therapy.

**Precautions.** Use with caution in patients with histories of liver disease. A previously positive Coombs’ test does not preclude methyldopa use, but early recognition of hemolytic anemia can be more difficult in such patients.

**Drug Interactions.** Methyldopa can potentiate the effect of tolbutamide and lithium. It also can cause confusion or disorientation when used with haloperidol. An increase in the pressor response of norepinephrine can occur with concurrent use. Iron products reduce methyldopa absorption. Amphetamines and heterocyclic antidepressants can decrease the efficacy of methyldopa. Levodopa and methyldopa can enhance each other’s effects.

**Parameters to Monitor.** Obtain direct Coombs’ test initially and at 6 and 12 months. Obtain baseline and periodic CBC and liver function tests to monitor for hemolytic anemia, blood dyscrasias, and hepatic dysfunction.

**Notes.** Methyldopa is not a first-line drug because of its frequent side effects, but it can be useful in those with ischemic heart disease or diastolic dysfunction because it reduces left ventricular mass.

**Minoxidil**

**Pharmacology.** Minoxidil is a potent vasodilator that acts by direct relaxation of arteriolar smooth muscle, thereby reducing total peripheral resistance. The vasodilation and associated reduction in blood pressure lead to reflex sympathetic activation, vagal inhibition, and altered renal homeostatic mechanisms manifested as increases in heart rate and cardiac output, increase in renin secretion, and salt and...
water retention. Because these responses can attenuate the hypotensive actions, give minoxidil with a sympatholytic drug and a diuretic. Topically, minoxidil stimulates vertex hair growth by an unknown mechanism.

**Administration and Adult Dosage.** PO for hypertension 5 mg/day initially as a single daily dose, increasing to 10, 20, and then 40 mg/day q 3 days in single or divided doses based on blood pressure response, to a maximum of 100 mg/day; usual dosage is 10–40 mg/day. If a single dose reduces supine diastolic blood pressure by more than 30 mmHg, divide the total daily dosage into 2 equal doses. **Top for male pattern baldness or female androgenetica** 1 mL to affected areas bid.

**Special Populations.** Pediatric Dosage. PO for hypertension 0.2 mg/kg as a single daily dose, increasing in 50–100% increments q 3 days until optimum blood pressure control or a total daily dosage of 50 mg is achieved; usual dosage is 0.25–1 mg/kg/day.

**Geriatric Dosage.** Same as adult dosage.

**Other Conditions.** In renal impairment, lower dosages might be required.

**Dosage Forms.** Tab 2.5, 10 mg (Loniten, various); Top 20, 50 mg/mL (2, 5%) (Rogaine, various).

**Patient Instructions.** (See Antihypertensives Class Instructions.) If a dose is missed, wait until the next regularly scheduled dose and continue with your regular dose; do not double the next dose. Report any of the following: increase in resting heart rate of greater than 20 beats per minute, rapid weight gain of more than 5 pounds, or the development of edema, increased difficulty in breathing, new or worsening angina, dizziness, lightheadedness, or fainting.

**Pharmacokinetics.** Onset and Duration. PO single dose onset 30 min; peak 2–3 hr; duration up to 75 hr with a gradual return to baseline at a rate of about 30% per day. Time to maximum effect with repeated administration is a function of dose and averages 7 days at 10 mg/day, 5 days at 20 mg/day, and 3 days at 40 mg/day. Top onset 4 or more months; relapse can occur 3–4 months after drug discontinuation.

**Serum Levels.** No correlation between serum levels and effects.

**Fate.** Oral absorption is at least 90%, but bioavailability is probably lower. Protein binding is negligible. Vd is 2.7 ± 0.7 L/kg; Cl is 1.4 ± 0.4 L/hr/kg. The drug is primarily metabolized and renally excreted, with about 20% unchanged drug in the urine. The major metabolite, a glucuronide conjugate, is active and might contribute to the drug’s effect. t1/2 3.1 ± 0.6 hr.

**Adverse Reactions.** Frequently, hypertrichosis (elongation, thickening, and enhanced pigmentation) (80%), transient ECG T-wave changes (60%), temporary edema (7%), or tachycardia occur. Occasionally, pericardial effusion with or without tamponade (3%), CHF, or angina occur. Rarely, breast tenderness and rashes (including Stevens–Johnson syndrome) occur. Minor dermatologic reactions occur occasionally after topical application.

**Contraindications.** (Oral) pheochromocytoma, caused by possible stimulation of catecholamine release from the tumor; acute MI; dissecting aortic aneurysm.
Precautions. For hypertension, minoxidil must usually be administered with a diuretic to prevent fluid retention; a loop diuretic is almost always required. Drugs or regimens that provide around-the-clock sympathetic suppression are usually required to prevent tachycardia, which can precipitate or worsen existing angina. Degenerative myocardial lesions reported in animal studies have yet to be confirmed in humans.

Drug Interactions. Concomitant therapy with guanethidine can result in profound orthostatic hypotension; discontinue guanethidine 1–3 weeks before initiation of oral minoxidil therapy or initiate therapy in the hospital.


Notes. Minoxidil is reserved for use in severe hypertension in combination with other drugs, usually a diuretic and a sympatholytic drug (eg, β-blocker).

Nitroprusside Sodium

Pharmacology. Nitroprusside is a potent vasodilator that has direct action on vascular smooth muscle to reduce arterial pressure and produce a slight increase in heart rate, a mild decrease in cardiac output, and a moderate reduction in total peripheral resistance. The decrease in total peripheral resistance suggests arteriolar dilation (afterload reduction), whereas the reduction in cardiac output might be caused by peripheral pooling of blood (preload reduction). Nitroprusside is somewhat more active on veins than on arteries. The active component of sodium nitroprusside is the free nitroso (NO\(^{-}\)) group.

Administration and Adult Dosage. IV 0.3 \(\mu\)g/kg/min by continuous infusion initially, increasing to an average rate of 3 \(\mu\)g/kg/min based on blood pressure response with a range of 0.5–10 \(\mu\)g/kg/min. Infusion at the maximum rate should never exceed 10 min. Patients receiving other antihypertensives can usually be controlled with smaller dosages. Control administration rates carefully with a microdrip regulator or an infusion pump; avoid too rapid reduction in blood pressure. Infusion rates greater than 2 \(\mu\)g/kg/min generate more cyanide ion (CN\(^{-}\)) than the body can metabolize or eliminate. Maintain infusions at the lowest possible dosage for the shortest possible duration to avoid toxicity.\(^\text{207}\) (See Adverse Reactions.)

Special Populations. Pediatric Dosage. IV same as adult dosage.

Geriatric Dosage. Initiate therapy with low infusion rates and carefully titrate the rate and degree of lowering blood pressure to avoid coronary and cerebral hypoperfusion.

Other Conditions. Patients with CHF, stroke, or receiving other antihypertensive drugs might be particularly sensitive to the blood-pressure–lowering effects of nitroprusside sodium; initiate therapy with low infusion rates and carefully titrate the rate and degree of lowering blood pressure to avoid coronary and cerebral hypoperfusions. Limit the total dosage in renal failure to avoid accumulation of thiocyanate. Use caution in hepatic insufficiency.

Dosage Forms. Inj 50 mg.
Pharmacokinetics. Onset and Duration. Onset within 1 min; peak 1–2 min; blood pressure usually returns to pretreatment levels in 2–10 min.\textsuperscript{173}

Serum Levels. Therapeutic and toxic levels are not established for nitroprusside because of rapid metabolism to cyanide and thiocyanate. Thiocyanate levels >60 mg/L (1 mmol/L) are associated with toxicity.

Fate. Nitroprusside is distributed in a volume that approximates the extravascular space, from which it is rapidly metabolized by a reaction with hemoglobin, yielding cyanmethemoglobin and an unstable intermediate that dissociates, releasing cyanide ion. Cyanide is converted to thiocyanate by the enzyme thiosulfate–cyanide sulfur transferase (rhodanese) in the liver and the kidney. The rate of conversion is determined principally by the availability of sulfur, usually as thiosulfate. Thiocyanate is excreted largely by the kidneys and can accumulate with high infusion rates for prolonged periods or renal dysfunction.

\[ t_{1/2} \] (Nitroprusside) 2 min; (thiocyanate) 2.7 days, up to 9 days in patients with renal dysfunction.\textsuperscript{208}

Adverse Reactions. Most adverse reactions are related to excessive or too rapid reduction of blood pressure and include nausea, retching, diaphoresis, apprehension, restlessness, headache, retrosternal discomfort, palpitations, dizziness, and abdominal pain, all of which resolve when the infusion rate is reduced or the infusion is temporarily discontinued. Thiocyanate is not particularly toxic and usually accumulates to toxic levels only with prolonged (>48 hr) or high-dosage (>10 μg/kg/min) infusions, when cyanide elimination is increased by the administration of thiosulfate, or in the presence of renal dysfunction. To limit the risk of thiocyanate toxicity, infuse at <3 μg/kg/min. Manifestations of thiocyanate toxicity include fatigue, anorexia, nausea, disorientation, toxic psychosis, and hallucinations. Cyanide toxicity usually occurs only when large dosages (>10 μg/kg/min) are infused rapidly or for longer than 1 hr. An early manifestation of cyanide toxicity can be apparent nitroprusside resistance, so increasing dosage requirements to achieve the same level of blood pressure control is an indication to look for metabolic acidosis, an indicator of cyanide toxicity, that might not be evident for more than 1 hr after accumulation of dangerous cyanide levels. Other symptoms of cyanide toxicity include dyspnea, vomiting, dizziness, loss of consciousness, weak pulse, distant heart sounds, areflexia, dilated pupils, shallow breathing, convulsions, and the occasional smell of bitter almonds on the breath. Hydroxocobalamin (25 mg/hr by continuous infusion) can facilitate the conversion of cyanide to cyanocobalamin,\textsuperscript{209} but an appropriate hydroxocobalamin dosage form is unavailable. Concurrent sodium thiosulfate administration also can prevent cyanide toxicity, but thiocyanate levels can increase.\textsuperscript{210} Management of cyanide toxicity includes immediate discontinuation of nitroprusside and the administration of sodium nitrite (0.2 mL/kg of a 3% solution IV over 2–4 min), followed by 12.5 g of sodium thiosulfate infused over 10 min. Methemoglobinemia can develop in patients congenitally unable to convert nitroprusside-induced methemoglobin back to hemoglobin. Management consists of IV administration of methylene blue 1–2 mg/kg over several minutes.

Contraindications. Compensatory hypertension (eg, arteriovenous shunt or coarctation of the aorta); controlled hypotension during surgery in patients with
inadequate cerebral circulation; congenital (Leber’s) optic atrophy; use of sildenafil. (See Drug Interactions.)

**Precautions.** If an adequate hypotensive response is not achieved after the maximum recommended infusion rate of 10 μg/kg/min for a maximum of 10 min, stop the infusion because these dosages increase the risk of toxicity. Use with caution in renal, hepatic, or thyroid disease, and in vitamin B12 deficiency or elevated intracranial pressure.

**Drug Interactions.** Use during general anesthesia can impair the capacity to compensate for hypovolemia and anemia and cause abnormal perfusion:ventilation ratio. Use in patients taking sildenafil can result in profound hypotension with serious consequences, including death.

**Parameters to Monitor.** Monitor blood pressure frequently (ie, every few minutes) because of the rapid onset and offset of effects. Monitor thiocyanate levels after 24–48 hr in patients with normal renal function and daily in patients with impaired renal function or receiving large dosages. However, these levels are of no value in detecting cyanide toxicity. Monitoring of serum cyanide concentrations has been recommended, but the assay is technically difficult and not readily interpretable if fluids other than packed RBCs are analyzed. Frequent monitoring of acid–base balance, particularly in patients with hepatic dysfunction, is considered adequate by most clinicians.

**Notes.** Protect from light and discard solution after 24 hr or if the color changes from the usual faint brownish tint to blue, green, or dark red. Do not administer IV push medications through the same line or use the solution for the simultaneous administration of any other drug.

### Pharmacology.

**Omapatrilat** (Investigational—Bristol-Myers Squibb) Vanlev

**Pharmacology.** Omapatrilat is the first of a new class of drugs called vasopeptidase inhibitors. Omapatrilat inhibits ACE and neutral endopeptidase, leading to blockades of the formation of angiotensin II and the breakdown of vasodilatory hormones such as natriuretic peptides, bradykinin, and adrenomedullin. This results in vasodilation, natriuresis, and diuresis.211

**Adult Dosage.** Not established.

**Pharmacokinetics.** Oral absorption is rapid, with peak plasma concentrations occurring 0.5–2 hr postdose. Biotransformation of the thiol group produces inactive metabolites; half-life is 14–19 hr; dosage adjustments are not necessary in renal dysfunction.211

**Adverse Reactions.** Omapatrilat is well tolerated, with an adverse event profile similar to that of placebo. The most commonly reported adverse reactions are hypotension (11%) and cough (about 10%). Flushing and syncope (about 1%) also have been reported, and angioedema is rare.211

**Notes.** Omapatrilat produces greater blood pressure reductions than lisinopril in hypertensive patients and in one study reduced morbidity and mortality (not the primary endpoint) to a greater extent than lisinopril in patients with CHF.312,313
### ACE INHIBITORS COMPARISON CHART

<table>
<thead>
<tr>
<th>DRUG</th>
<th>DOSAGE FORMS</th>
<th>DAILY ADULT DOSAGE (MG)</th>
<th>INDICATED FOR CHF</th>
<th>PEAK EFFECT (HR)</th>
<th>DURATION (HR)</th>
<th>HALF-LIFE (HR)</th>
<th>ELIMINATION ROUTES</th>
</tr>
</thead>
<tbody>
<tr>
<td>Benazepril</td>
<td>Tab 5, 10, 20, 40 mg.</td>
<td>20–40</td>
<td>No</td>
<td>2–4</td>
<td>24+</td>
<td>10–11</td>
<td>Renal, Hepatic.</td>
</tr>
<tr>
<td>Lotensin</td>
<td></td>
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</tr>
<tr>
<td>Captopril</td>
<td>Tab 12.5, 25, 50, 100 mg.</td>
<td>50–150</td>
<td>Yes</td>
<td>1</td>
<td>6–10</td>
<td>2.2</td>
<td>Renal</td>
</tr>
<tr>
<td>Capoten</td>
<td></td>
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<tr>
<td>Various</td>
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</tr>
<tr>
<td>Enalapril</td>
<td>Tab 2.5, 5, 10, 20 mg.</td>
<td>PO 10–40; IV 1.25 mg q 6 hr.</td>
<td>Yes</td>
<td>4–6 (PO)</td>
<td>24 (PO)</td>
<td>11</td>
<td>Renal.</td>
</tr>
<tr>
<td>Vasotec</td>
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</tr>
<tr>
<td>Fosinopril</td>
<td>Tab 10, 20, 40 mg.</td>
<td>20–40</td>
<td>Yes</td>
<td>3–6</td>
<td>24</td>
<td>12–15</td>
<td>Hepatic, Renal.</td>
</tr>
<tr>
<td>Monopril</td>
<td></td>
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</tr>
<tr>
<td>Lisinopril</td>
<td>Tab 2.5, 5, 10, 20, 30, 40 mg.</td>
<td>10–40</td>
<td>Yes</td>
<td>6</td>
<td>24</td>
<td>12</td>
<td>Renal.</td>
</tr>
<tr>
<td>Prinivil</td>
<td></td>
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<tr>
<td>Zestril</td>
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</tr>
<tr>
<td>Moexipril</td>
<td>Tab 7.5, 15 mg.</td>
<td>7.5–30</td>
<td>No</td>
<td>3–8</td>
<td>24</td>
<td>2–9</td>
<td>Hepatic, Renal.</td>
</tr>
<tr>
<td>Univasc</td>
<td></td>
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</tr>
<tr>
<td>Perindopril</td>
<td>Tab 2, 4, 8 mg.</td>
<td>4–8</td>
<td>No</td>
<td>3–7</td>
<td>24+</td>
<td>3–10</td>
<td>Renal.</td>
</tr>
<tr>
<td>Aceon</td>
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</tr>
<tr>
<td>Quinapril</td>
<td>Tab 5, 10, 20, 40 mg.</td>
<td>20–80</td>
<td>Yes</td>
<td>2–4</td>
<td>24+</td>
<td>2–3</td>
<td>Renal.</td>
</tr>
</tbody>
</table>

(continued)
## ACE INHIBITORS COMPARISON CHART (continued)

<table>
<thead>
<tr>
<th>DRUG</th>
<th>DOSAGE FORMS</th>
<th>DAILY ADULT DOSAGE (MG)⁴</th>
<th>INDICATED FOR CHF</th>
<th>PEAK EFFECT (HR)</th>
<th>DURATION (HR)</th>
<th>HALF-LIFE (HR)</th>
<th>ELIMINATION ROUTES</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ramipril</td>
<td>Cap 1.25, 2.5, 5, 10 mg.</td>
<td>2.5–20</td>
<td>Yes⁵</td>
<td>3–8</td>
<td>24+</td>
<td>13–17b</td>
<td>Renal, Hepatic.</td>
</tr>
<tr>
<td>Altace</td>
<td>Tab 1, 2, 4 mg.</td>
<td>2–4</td>
<td>Yes⁵</td>
<td>6–8</td>
<td>24+</td>
<td>10⁹</td>
<td>Hepatic, Renal.</td>
</tr>
<tr>
<td>Trandolapril</td>
<td>Mavik</td>
<td></td>
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<td></td>
</tr>
</tbody>
</table>

⁴Usual maintenance dosage range for hypertension. Initial dosage is often lower, and higher dosages are sometimes effective.

⁵Half-life of active drug.

⁶Indicated for CHF post-MI.

From references 214 and 215 and product information.
<table>
<thead>
<tr>
<th>DRUG</th>
<th>DOSAGE FORMS</th>
<th>USUAL DAILY ADULT DOSAGE (MG)</th>
<th>PEAK EFFECT (HR)</th>
<th>DURATION (HR)</th>
<th>HALF-LIFE (HR)</th>
<th>ELIMINATION ROUTES</th>
</tr>
</thead>
<tbody>
<tr>
<td>Candesartan</td>
<td>Tab 4, 8, 16, 32 mg.</td>
<td>8–32&lt;sup&gt;a&lt;/sup&gt;</td>
<td>3–4</td>
<td>24+</td>
<td>9</td>
<td>Hepatic, Renal.</td>
</tr>
<tr>
<td>Atacand</td>
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<tr>
<td>Eprosartan</td>
<td>Tab 400, 600 mg.</td>
<td>400–800&lt;sup&gt;a&lt;/sup&gt;</td>
<td>1–3</td>
<td>24+</td>
<td>5–9</td>
<td>Hepatic, Renal.</td>
</tr>
<tr>
<td>Tevoten</td>
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</tr>
<tr>
<td>Irbesartan</td>
<td>Tab 75, 150, 300 mg.</td>
<td>150–300</td>
<td>3–6</td>
<td>24+</td>
<td>11–15</td>
<td>Hepatic, Renal.</td>
</tr>
<tr>
<td>Avapro</td>
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</tr>
<tr>
<td>Losartan</td>
<td>Tab 25, 50, 100 mg.</td>
<td>25–100&lt;sup&gt;b&lt;/sup&gt;</td>
<td>6</td>
<td>24+</td>
<td>2</td>
<td>Hepatic.</td>
</tr>
<tr>
<td>Cozaar</td>
<td></td>
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<td></td>
<td></td>
<td>Renal, Hepatic.</td>
</tr>
<tr>
<td>Telmisartan</td>
<td>Tab 40, 80 mg.</td>
<td>40–80</td>
<td>&gt;3</td>
<td>24+</td>
<td>6–9&lt;sup&gt;b&lt;/sup&gt;</td>
<td>Hepatic.</td>
</tr>
<tr>
<td>Micardis</td>
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<tr>
<td>Valsartan</td>
<td>Cap 80, 160 mg.</td>
<td>80–320</td>
<td>6</td>
<td>24+</td>
<td>6</td>
<td>Hepatic.</td>
</tr>
<tr>
<td>Diovan</td>
<td></td>
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</tr>
</tbody>
</table>

<sup>a</sup>Occasionally, the daily dosage can be given in 2 divided doses.

<sup>b</sup>For active metabolite, which is responsible for most or all pharmacologic effects.

*From references 197, 216, and 217 and product information.*
<table>
<thead>
<tr>
<th>DRUG</th>
<th>DOSAGE FORMS</th>
<th>DAILY ADULT DOSAGE (MG)</th>
<th>PEAK EFFECT (HR)</th>
<th>DURATION (HR)</th>
<th>HALF-LIFE (HR)</th>
<th>ELIMINATION ROUTES</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Doxazosin</strong>&lt;br&gt;Cardura</td>
<td>Tab 1, 2, 4, 8 mg.</td>
<td>1–16</td>
<td>2–3</td>
<td>24</td>
<td>10–22</td>
<td>Hepatic.</td>
</tr>
<tr>
<td><strong>Prazosin</strong>&lt;br&gt;Minipress Various</td>
<td>Cap 1, 2, 5 mg.</td>
<td>2–20</td>
<td>1–3</td>
<td>6–12</td>
<td>2–3</td>
<td>Hepatic.</td>
</tr>
<tr>
<td><strong>Terazosin</strong>&lt;br&gt;Hytrin Various</td>
<td>Tab 1, 2, 5, 10 mg.</td>
<td>1–20</td>
<td>1–2</td>
<td>24</td>
<td>9–16</td>
<td>Hepatic, Renal.</td>
</tr>
<tr>
<td><strong>Tamsulosin</strong>&lt;br&gt;Flomax</td>
<td>Cap 0.4 mg.</td>
<td>0.4–0.8&lt;sup&gt;b&lt;/sup&gt;</td>
<td>—</td>
<td>—</td>
<td>14–15</td>
<td>Hepatic.</td>
</tr>
</tbody>
</table>

<sup>a</sup>Usual maintenance dosage range for hypertension; higher dosages are sometimes effective. Dosage is the same in the elderly.

<sup>b</sup>Not for hypertension; for symptoms of benign prostatic hypertrophy only.

From references 147, and 218 and product information.
<table>
<thead>
<tr>
<th>DRUG</th>
<th>DOSAGE FORMS</th>
<th>ADULT DOSAGE</th>
<th>DURATION</th>
<th>ADVERSE EFFECTS</th>
<th>MECHANISM</th>
</tr>
</thead>
<tbody>
<tr>
<td>Guanabenz</td>
<td>Tab 4, 8 mg.</td>
<td>PO 4 mg bid, increasing q 1–2 weeks to a maximum of 32 mg bid.</td>
<td>12 hr</td>
<td>See clonidine monograph</td>
<td>See clonidine monograph.</td>
</tr>
<tr>
<td>Acetate</td>
<td>Wytensin</td>
<td></td>
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</tr>
<tr>
<td></td>
<td>Various</td>
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</tr>
<tr>
<td>Guanadrel</td>
<td>Tab 10, 25 mg.</td>
<td>PO 5 mg bid, increasing q 1–4 weeks to 20–75 mg/day. Usual maximum is 150 mg/day in 2 divided doses.</td>
<td>4–14 hr</td>
<td>Orthostatic hypotension, diarrhea, drowsiness, sexual dysfunction, peripheral edema, nasal stuffiness, palpitations, shortness of breath, leg cramps, aching limbs.</td>
<td>Postganglionic adrenergic blockade.</td>
</tr>
<tr>
<td>Sulfate</td>
<td>Hylorel</td>
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</tr>
<tr>
<td>Guanethidine</td>
<td>Tab 10, 25 mg.</td>
<td>PO 10 mg/day, increasing q 5–7 days to 25–50 mg once daily.</td>
<td>1–3 weeks</td>
<td>Same as guanadrel, but more frequent.</td>
<td>Postganglionic adrenergic blockade.</td>
</tr>
<tr>
<td>Sulfate</td>
<td>Ismelin</td>
<td></td>
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</tr>
<tr>
<td>Guanfacine</td>
<td>Tab 1, 2 mg.</td>
<td>PO 1 mg/day, increasing q 3–4 weeks to maximum of 3 mg/day.</td>
<td>2–4 days</td>
<td>See clonidine monograph.</td>
<td>See clonidine monograph.</td>
</tr>
<tr>
<td>Hydrochloride</td>
<td>Tenex</td>
<td></td>
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</tr>
<tr>
<td></td>
<td>Various</td>
<td></td>
<td></td>
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<td></td>
</tr>
<tr>
<td>Reserpine</td>
<td>Tab 0.1, 0.25 mg.</td>
<td>PO 0.5 mg/day for 1–2 weeks, then 0.1–0.25 mg/day.</td>
<td>24 hr</td>
<td>Drowsiness, weakness, GI disturbances, nasal congestion, sexual dysfunction, bradycardia. Dose-related mental depression occurs.</td>
<td>Depletes norepinephrine from post-ganglionic adrenergic neurons.</td>
</tr>
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</tbody>
</table>
### DRUGS FOR HYPERTENSIVE URGENCIES AND EMERGENCIES COMPARISON CHART

<table>
<thead>
<tr>
<th>DRUG</th>
<th>DOSAGE RANGE</th>
<th>ONSET (MIN)</th>
<th>DURATION</th>
<th>COMMENTS</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Captopril</strong></td>
<td>PO, SL 12.5–25 mg.</td>
<td>10–30</td>
<td>2–6 hr</td>
<td>Hypotensive effect is particularly large in patients on a diuretic or in hypertensive crisis. Subsequent doses may be less effective unless given with a diuretic. Acute renal failure can occur.</td>
</tr>
<tr>
<td><strong>Clonidine</strong></td>
<td>PO 0.1–0.2 mg initially, then 0.1 mg/hr, to a maximum total dosage of 0.8 mg.</td>
<td>30–120</td>
<td>6–8 hr</td>
<td>Rate of onset is slower after a meal; drowsiness or dry mouth can occur. Rebound hypertension is possible.</td>
</tr>
<tr>
<td><strong>Labetalol</strong></td>
<td>PO 200–400 mg, may repeat q 2–3 hr.</td>
<td>30–120</td>
<td>6–12 hr</td>
<td>Orthostatic hypotension, bronchoconstriction, and heart block can occur. Avoid in COPD and asthma.</td>
</tr>
<tr>
<td><strong>Prazosin</strong></td>
<td>PO 1–2 mg, may repeat q 1 hr.</td>
<td>30–90</td>
<td>1–10 hr</td>
<td>Useful in presence of increased circulating catecholamines. First-dose syncope, palpitations, tachycardia, and headache reported.</td>
</tr>
</tbody>
</table>

(continued)
<table>
<thead>
<tr>
<th>DRUG</th>
<th>ONSET (MIN)</th>
<th>DURATION</th>
<th>COMMENTS</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>INTRAVENTOUS DRUGS FOR HYPERTENSIVE EMERGENCIES</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Diazoxide</td>
<td>2–4</td>
<td>3–12 hr</td>
<td>Now obsolete, but can be useful in hypertensive encephalopathy, malignant hypertension, and eclampsia. Increases cardiac output; requires blood pressure monitoring at hourly intervals. Avoid with ischemic heart disease or intracranial hemorrhage.</td>
</tr>
<tr>
<td>Hyperstat I.V.</td>
<td>30 sec, may repeat q 5–15 min. Alternatively, IV infusion 10–30 mg/min. After 300 mg given, give furosemide IV 40 mg before subsequent doses.</td>
<td>15–30</td>
<td>4–6 hr</td>
</tr>
<tr>
<td>Enalaprilat</td>
<td></td>
<td></td>
<td>Use in CHF and those at risk for cerebral hypotension. Avoid in acute MI or severe renal impairment. Blacks may respond poorly. Hypotension may occur.</td>
</tr>
<tr>
<td>Enalapril I.V.</td>
<td></td>
<td></td>
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</tr>
<tr>
<td>Esmolol</td>
<td>1–2</td>
<td>10–20 min</td>
<td>Use in perioperative patients with aortic dissection. Does not cause tachycardia but does decrease heart rate.</td>
</tr>
<tr>
<td>Brevibloc</td>
<td></td>
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</tr>
<tr>
<td>Fenoldopam</td>
<td>&lt;5</td>
<td>30 min</td>
<td>Use in patients with renal insufficiency who risk cyanide toxicity with nitroprusside. Use with caution in glaucoma.</td>
</tr>
<tr>
<td>Corlopam</td>
<td></td>
<td></td>
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<tr>
<td>Hydralazine</td>
<td>10–20 (IV)</td>
<td>3–8 hr</td>
<td>Limited to treatment of severe pre-eclampsia and eclampsia. Increases cardiac output; many patients sensitive to parenteral doses, resulting in excessive hypotension.</td>
</tr>
<tr>
<td>Apresoline</td>
<td>20–30 (IM)</td>
<td></td>
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</tr>
<tr>
<td>Labetalol</td>
<td>&lt;10</td>
<td>3–6 hr</td>
<td>Hypotensive effect is predictable; contraindicated in CHF, head trauma, and intracranial hemorrhage; often causes marked postural hypotension. Avoid use in patients with COPD, CHF, or bradycardia.</td>
</tr>
<tr>
<td>Normodyne</td>
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<tr>
<td>Trandate</td>
<td></td>
<td></td>
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</tbody>
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(continued)
<table>
<thead>
<tr>
<th>DRUG</th>
<th>DOSAGE RANGE</th>
<th>ONSET (MIN)</th>
<th>DURATION</th>
<th>COMMENTS</th>
</tr>
</thead>
</table>
| Nicardipine
Cardene | IV infusion 5–15 mg/hr.       | <5–15       | 1–4 hr   | Predictable effect. Useful in coronary, cerebral, or peripheral artery disease and in surgical patients. Tachycardia can occur. Use with caution in patients with coronary ischemia. |
| Nitroglycerin
Various | IV 0.3–6 mg/hr by continuous infusion. | 1–5         | 3–5 min  | Useful in myocardial ischemia and hypertension associated with MI. Hypotension, headache, tachycardia, and tachyphylaxis occur. Avoid in constrictive pericarditis, pericardial tamponade, or intracranial hypertension. |
| Nitroprusside
Sodium
Nipride
Various | IV 0.3–10 µg/kg/min by continuous infusion. Infuse at maximal dosage for no more than 10 min. Average dosage is 3 µg/kg/min. | 0.5–1       | 1–2 min  | Especially useful in ischemic heart disease. Continuous monitoring required; arterial pressure response adjusted by changing infusion rate; hypotensive effect enhanced by elevating head of patient’s bed. Decreases cardiac output; cyanide toxicity with prolonged, high infusion rates. |

Adapted from references 173, 195, 219 and 220.
ESMOLOL HYDROCHLORIDE

**Pharmacology.** Esmolol is an ultrashort-acting, cardioselective, \( \beta_1 \)-adrenergic blocking agent. It is effective in controlling ventricular response in patients with atrial fibrillation and other supraventricular tachycardias and in slowing heart rate in patients with sinus tachycardia associated with acute MI or cardiac surgery. Esmolol is useful for treating hypertensive emergencies, particularly in patients with tachycardia, because it has a rapid onset, short duration of action, and reduces heart rate. It also can be effective in perioperative hypertension.\(^1\)\(\text{73,195,221,222}\)

**Adult Dosage.** Dilute injection to a final concentration of 10 mg/mL. \( \text{IV} \) loading dose is 500 \( \mu \text{g/kg/min} \) for 1 min and then 50 \( \mu \text{g/kg/min} \). The \( \text{IV} \) loading dose can be repeated as often as every 5 min, with a concomitant increase of infusion rate in 50 \( \mu \text{g/kg/min} \) increments, titrated to ventricular response, heart rate, and/or blood pressure. Most patients respond to infusions of 100–200 \( \mu \text{g/kg/min} \) once the desired endpoint is obtained, the infusion rate can be decreased in 25–50 \( \mu \text{g/kg/min} \) increments at 5- to 10-min intervals. Infusions up to 48 hr are well tolerated.

**Pediatric Dosage.** \( \text{IV} \) 500 \( \mu \text{g/kg/min} \) for 1 min and then 25–200 (average 120) \( \mu \text{g/kg/min} \).\(^2\) Weight-adjusted dosages can be higher than in adults because of its more rapid elimination in children; infusion rates as high as 1 mg/kg/min have been required to achieve complete \( \beta \) blockade.\(^2\)

**Dosage Forms.** \( \text{Inj} \) 10, 250 mg/mL.

**Pharmacokinetics.** Effective plasma levels are about 1–1.5 mg/L (3.4–5.1 \( \mu \text{mol/L} \)). The \( \alpha \) half-life is about 2 min; \( V_d \) averages 3.5 L/kg (range 2–5). Esmolol is rapidly hydrolyzed by plasma and blood esterases to a metabolite with weak, clinically unimportant \( \beta \)-blocking activity and small amounts of methanol. No unchanged esmolol appears in the urine. The elimination half-life is about 9 min in adults and 3 min in children.\(^2\)\(^,2\)\(^2\)

**Adverse Reactions.** The side effect profile is similar to that of other \( \beta_1 \)-selective \( \beta \)-blockers. Dose-related hypotension is frequent; \( \text{IV} \) site phlebitis occurs occasionally. Concurrent \( \text{IV} \) morphine can increase serum levels by 46%.

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PROPRANOLOL HYDROCHLORIDE

**Pharmacology.** Propranolol is a nonselective \( \beta \)-adrenergic blocker used in arrhythmias, hypertension, angina pectoris, and CHF. It is also effective in decreasing post-MI mortality. The antiarrhythmic mechanism is caused by decreased AV nodal conduction in supraventricular tachycardias and blockade of catecholamine-induced dysrhythmias. Propranolol and other \( \beta \)-blockers are effective in preventing postoperative atrial fibrillation. The antihypertensive mechanism is unknown, but contributing factors are a CNS mechanism, renin blockade, and decreases in myocardial contractility and cardiac output. Propranolol also lowers myocardial oxygen demand by decreasing contractility and heart rate, which symptomatically alleviates anginal pain and increases exercise tolerance in coronary artery disease. Metoprolol and carvedilol (and perhaps other \( \beta \)-blockers) are effective in reduc-
ing mortality and improving quality of life in patients with CHF by blocking dele-
terious neurohumoral compensatory factors. β-Blockers and diuretics are recom-
mended as first-line drugs for hypertension because of demonstrated reductions in
morbidity and mortality.173 (See β-Adrenergic Blocking Drugs Comparison
Chart.)

**Administration and Adult Dosage.** PO 10–20 mg q 6 hr initially, increasing gradu-
ally to desired effects. In hypertension, more than 1 g/day has been used; how-
ever, consider adding another drug if 480 mg/day is ineffective. In angina pector-
itis, the dosage is titrated to pain relief and exercise evidence of β-blockade
(bradycardia). The endpoint for dosage escalation in acute arrhythmias is the re-
turn to sinus rhythm or, in atrial fibrillation or flutter, to a ventricular rate below
100 beats/min with hemodynamic stability. Twice-daily administration is effect-
ive in angina pectoris and hypertension. Administer SR Cap in the same daily
dosage once or twice daily (not indicated post-MI). PO for post-MI prophy-
axis (non-SR) 180–240 mg/day in 2–3 divided doses. IV slow push 1 mg q 5 min, to
a maximum of 0.15 mg/kg; some investigators have recommended that the first
dose be given over 2–10 min.

**Special Populations. Pediatric Dosage.** PO for hypertension 0.5–1 mg/kg/day in
2–4 divided doses, increasing to a maximum of 8 mg/kg/day. IV slow push
0.01–0.1 mg/kg/dose over 10 min up to 1 mg (infants) or 3 mg (children); may re-
peat in 6–8 hr.4

**Geriatric Dosage.** Bioavailability is increased in the elderly, necessitating lower
initial doses.

**Other Conditions.** Therapeutic endpoints can be achieved with lower dosages in
hypothyroidism or liver disease. Begin with lower dosages and titrate to clinical
response. Patients with thyrotoxicosis require higher dosages to achieve the de-
sired effect.224

**Dosage Forms.** Soln 4, 8, 80 mg/mL; Tab 10, 20, 40, 60, 80, 90 mg; SR Cap 60,
80, 120, 160 mg; Inj 1 mg/mL.

**Patient Instructions.** Report any symptoms such as shortness of breath, swelling,
wheezing, fatigue, depression, nightmares, or inability to concentrate. Do not stop
therapy abruptly. Do not crush or chew SR capsule. A sustained-release capsule
core in the stool does not indicate lack of absorption.

**Missed Doses.** Take this drug at regular intervals. If you miss a dose take it as
soon as you remember. If it is about time for the next dose, take that dose only.
Leave at least 4 hours between regular tablet doses and 6–8 hours between
extended-release capsule doses. Do not double the dose or take extra.

**Pharmacokinetics. Onset and Duration.** PO onset is variable; the duration varies
from 6 to longer than 12 hr.224

**Serum Levels.** No definite relation has been established between serum concen-
trations and therapeutic effect in the treatment of arrhythmias, angina pectoris, or
hypertension. β-Blockade is associated with serum concentrations >100 μg/L
(340 nmol/L).225
Fate. Propranolol is rapidly and completely absorbed after oral administration; however, a large hepatic first-pass effect occurs, limiting systemic availability to 26 ± 10%. First-pass elimination is saturable with an oral dose greater than about 30 mg.225 The drug is 87 ± 6% bound to α1-acid glycoprotein and other plasma proteins.10,224 Vd is 4.3 ± 0.6 L/kg; Cl is 0.96 ± 0.3 L/hr/kg. Unlike most other drugs, displacement from plasma proteins increases elimination half-life and Vd because of high tissue affinity (nonrestrictive elimination). An active metabolite, 4-hydroxypropranolol, is formed after oral, but not IV, administration. Less than 0.5% of a dose is excreted unchanged in urine.10

\[ t_{1/2}\ alpha \] phase is about 10 min;224 \[ t_{1/2}\ beta \] phase after a single PO dose is 3.9 ± 0.4 hr.10 With long-term oral therapy, \[ t_{1/2}\ beta \] phase is 4–6 hr but can be as long as 10–20 hr in patients with liver disease.226

Adverse Reactions. Adverse effects often are not related to dose. Depression, nightmares, insomnia, fatigue, and lethargy occur frequently; less often, psychotic changes have been reported. CNS side effects probably occur more often with the lipophilic β-blockers (eg, propranolol). The drug can cause occasional life-threatening reactions when therapy (especially IV) is initiated, and acute CHF with pulmonary edema and hypotension or symptomatic bradycardia and heart block can occur. Acute drug cessation in patients with coronary artery disease can precipitate unstable angina pectoris or MI. The drug can precipitate hypoglycemia, but probably more important in diabetics is its ability to mask hypoglycemic symptoms (except for sweating). It can exacerbate symptoms of peripheral vascular disease or Raynaud’s disease. β-Blockers can exacerbate previously stable asthma or chronic airway obstruction by causing bronchospasm or renal dysfunction by further depressing GFR.

Contraindications. Severe obstructive pulmonary disease, asthma or active allergic rhinitis; cardiogenic shock or severe CHF; second- or third-degree heart block; severe sinus node disease.

Precautions. In coronary artery disease, discontinue drug by tapering the dosage over 4–7 days. Use cautiously in patients with Prinzmetal’s vasospastic angina to prevent worsening of chest pain. Use caution in peripheral vascular disease or CHF and in patients with brittle diabetes or history of hypoglycemic episodes. Can worsen atrial fibrillation associated with accessory AV pathway.

Drug Interactions. Concurrent digoxin therapy can lessen the β-blocker exacerbation of CHF. When taken with oral hypoglycemics, nonselective β-blockers such as propranolol prolong hypoglycemic episodes and inhibit tachycardia and tremors, which are signs of hypoglycemia (sweating is not inhibited); hypertension can occur during hypoglycemia. Epinephrine can produce hypertensive reactions in patients on propranolol (and probably other nonselective β-blockers); this can occur with other sympathomimetics such as phenylephrine and phenylpropanolamine. Barbiturates and rifampin can increase the metabolism of hepatically eliminated β-blockers such as propranolol. Cimetidine can increase propranolol effects. Combined use of clonidine and propranolol can result in hypertensive reactions, especially if clonidine is abruptly discontinued. β-Block-
ers can increase the first-dose hypotensive effect of prazosin and similar drugs. NSAIDs can blunt the hypotensive response of β-blockers.

**Parameters to Monitor.** During IV administration, obtain blood pressure and pulse q 5 min with constant ECG monitoring for signs of AV nodal block (lengthened PR interval) or bradycardia. Evaluate vital signs routinely for hemodynamic endpoints (eg, blood pressure in hypertension and heart rate or pressure rate product in angina pectoris). Question the patient about subjective complaints such as nightmares or fatigue. When a patient at risk for adverse reactions is first given propranolol, evaluate signs and symptoms of toxicity (eg, CHF, shortness of breath or edema; bronchospasm, wheezing or shortness of breath; diabetes, blood glucose; peripheral vascular disease, painful or cold extremities).

**Notes.** Propranolol can be beneficial for treatment of symptomatic hypertrophic obstructive cardiomyopathy by increasing end-diastolic volume, producing ventricular relaxation, and relieving ventricular outflow obstruction. Other uses include migraine prophylaxis, prevention of GI bleeding in patients with esophageal varices, prevention of sudden death in congenital long-QT syndromes, and as a cardiac protectant in patients with heart disease undergoing noncardiac surgery. If a β-blocker must be used in lung disease, β₁-selective drugs (eg, acebutolol, atenolol, or metoprolol) cause alterations in pulmonary function that are more easily reversed by bronchodilators; these drugs are probably a better choice than propranolol or other nonselective β-blockers. (See β-Adrenergic Blocking Drugs Comparison Chart.)
<table>
<thead>
<tr>
<th>DRUG</th>
<th>DOSAGE FORMS</th>
<th>CARDIO-SELECTIVITY</th>
<th>β HALF-LIFE (HR)</th>
<th>EXCRETED IN URINE</th>
<th>PROTEIN BINDING</th>
<th>Labeled USES</th>
<th>STARTING DOSAGE</th>
<th>MAXIMUM DOSAGE</th>
</tr>
</thead>
<tbody>
<tr>
<td>Acebutolol²</td>
<td>Cap 200, 400 mg.</td>
<td>+</td>
<td>3–4</td>
<td>30–40%</td>
<td>25%</td>
<td>Hypertension, arrhythmias.</td>
<td>PO 400 mg/day.</td>
<td>PO 1.2 g/day.</td>
</tr>
<tr>
<td>Sectral</td>
<td></td>
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<td></td>
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<tr>
<td>Various</td>
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<td></td>
</tr>
<tr>
<td>Atenolol</td>
<td>Tab 25, 50, 100 mg</td>
<td>+</td>
<td>6–7</td>
<td>85%</td>
<td>10%</td>
<td>Hypertension. Post-MI prophylaxis.</td>
<td>PO 50 mg/day.</td>
<td>PO 200 mg/day.</td>
</tr>
<tr>
<td>Tenormin</td>
<td>Tab 25, 50, 100 mg</td>
<td>+</td>
<td>6–7</td>
<td>85%</td>
<td>10%</td>
<td>Hypertension. Post-MI prophylaxis.</td>
<td>PO 50 mg/day.</td>
<td>PO 200 mg/day.</td>
</tr>
<tr>
<td>Various</td>
<td>Tab 25, 50, 100 mg</td>
<td>+</td>
<td>6–7</td>
<td>85%</td>
<td>10%</td>
<td>Hypertension. Post-MI prophylaxis.</td>
<td>PO 50 mg/day.</td>
<td>PO 200 mg/day.</td>
</tr>
<tr>
<td>Bevantolol</td>
<td>Tab 10, 20 mg.</td>
<td>+</td>
<td>14–20</td>
<td>15%</td>
<td>50%</td>
<td>Hypertension.</td>
<td>PO 10 mg/day.</td>
<td>PO 40 mg/day.</td>
</tr>
<tr>
<td>Kerlone</td>
<td></td>
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</tr>
<tr>
<td>Betaxolol</td>
<td>Tab 10, 20 mg.</td>
<td>+</td>
<td>14–20</td>
<td>15%</td>
<td>50%</td>
<td>Hypertension.</td>
<td>PO 10 mg/day.</td>
<td>PO 40 mg/day.</td>
</tr>
<tr>
<td>Bevantolol</td>
<td>Tab 10, 20 mg.</td>
<td>+</td>
<td>14–20</td>
<td>15%</td>
<td>50%</td>
<td>Hypertension.</td>
<td>PO 10 mg/day.</td>
<td>PO 40 mg/day.</td>
</tr>
<tr>
<td>Bisoprolol</td>
<td>Tab 5, 10 mg.</td>
<td>+</td>
<td>9–12</td>
<td>50%</td>
<td>30%</td>
<td>Hypertension.</td>
<td>PO 2–5 mg/day.</td>
<td>PO 20 mg/day.</td>
</tr>
<tr>
<td>Zebeta</td>
<td></td>
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</tr>
</tbody>
</table>

(continued)
<table>
<thead>
<tr>
<th>DRUG</th>
<th>DOSAGE FORMS</th>
<th>CARDIO-SELECTIVITY</th>
<th>β HALF-LIFE (HR)</th>
<th>EXCRETED UNCHANGED IN URINE</th>
<th>PROTEIN BINDING</th>
<th>LABELED USES</th>
<th>STARTING DOSAGE</th>
<th>MAXIMUM DOSAGE</th>
</tr>
</thead>
<tbody>
<tr>
<td>Carteolol</td>
<td>Tab 2.5, 5 mg</td>
<td>0</td>
<td>6–11</td>
<td>60%</td>
<td>15%</td>
<td>Hypertension.</td>
<td>PO 2.5 mg/day</td>
<td>PO 10 mg/day</td>
</tr>
<tr>
<td>Cartrol</td>
<td></td>
<td></td>
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</tr>
<tr>
<td>Carvedilol</td>
<td>Tab 3.125, 6.25, 12.5, 25 mg.</td>
<td>0</td>
<td>6–8</td>
<td>1%</td>
<td>95%</td>
<td>Hypertension, CHF.</td>
<td>PO 3.125 mg bid, increasing q 2 weeks.</td>
<td>PO (&lt;85 kg) 50 mg/day; PO (&gt;85 kg) 100 mg/day.</td>
</tr>
<tr>
<td>Coreg</td>
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</tr>
<tr>
<td>Esmolol</td>
<td>Inj 10, 250 mg/mL</td>
<td>+</td>
<td>9 min</td>
<td>0%</td>
<td>55%</td>
<td>Supraventricular tachycardia.</td>
<td>IV 50 µg/kg/min.</td>
<td>IV 200 µg/kg/min.</td>
</tr>
<tr>
<td>Brevibloc</td>
<td></td>
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</tr>
<tr>
<td>Labetalol</td>
<td>Tab 100, 200, 300 mg</td>
<td>0</td>
<td>4–9</td>
<td>5%</td>
<td>50%</td>
<td>Hypertension.</td>
<td>PO 100 mg/day; IV 20 mg, then 40–80 mg q 10 min.</td>
<td>PO 2.4 g/day; IV 300 mg.</td>
</tr>
<tr>
<td>Trandate</td>
<td>Inj 5 mg/mL</td>
<td></td>
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<tr>
<td>Normodyne</td>
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<tr>
<td>Various</td>
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</tr>
<tr>
<td>Metoprolol</td>
<td>Tab 50, 100 mg SR Tab 50, 100, 200 mg</td>
<td>+ (up to 100 mg)</td>
<td>3–7</td>
<td>39%</td>
<td>10%</td>
<td>Hypertension.</td>
<td>PO 100 mg/day; PO SR 50–100 mg/day.</td>
<td>PO 450 mg/day; PO SR 400 mg/day.</td>
</tr>
<tr>
<td>Lopressor</td>
<td>Inj 1 mg/mL</td>
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<td></td>
<td>Hypertension, angina pectoris. Acute MI.</td>
<td>IV 5 mg × 3, then PO 50 mg q 6 hr × 48 hr. PO SR 12.5–25 mg/day.</td>
<td>PO SR 200 mg/day.</td>
</tr>
<tr>
<td>Toprol-XL</td>
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(continued)
### β-Adrenergic Blocking Drugs Comparison Chart (continued)

<table>
<thead>
<tr>
<th>DRUG</th>
<th>DOSAGE FORMS</th>
<th>CARDIO-SELECTIVITY</th>
<th>β HALF-LIFE (HR)</th>
<th>EXCRETED UNCHANGED IN URINE</th>
<th>PROTEIN BINDING</th>
<th>LABELED USES</th>
<th>STARTING DOSAGE</th>
<th>MAXIMUM DOSAGE</th>
</tr>
</thead>
<tbody>
<tr>
<td>Nadolol</td>
<td>Tab 20, 40, 80, 120, 160 mg.</td>
<td>0</td>
<td>17–24</td>
<td>70%</td>
<td>25%</td>
<td>Hypertension, angina pectoris.</td>
<td>PO 40 mg/day.</td>
<td>PO 320 mg/day.</td>
</tr>
<tr>
<td>Corgard Various</td>
<td></td>
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</tr>
<tr>
<td>Penbutolol²</td>
<td>Tab 20 mg.</td>
<td>0</td>
<td>4–8</td>
<td>5%</td>
<td>80–90%</td>
<td>Hypertension.</td>
<td>PO 20 mg/day.</td>
<td>PO 80 mg/day.</td>
</tr>
<tr>
<td>Levatol</td>
<td></td>
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</tr>
<tr>
<td>Penbutolol²</td>
<td>Tab 5, 10 mg.</td>
<td>0</td>
<td>3–4</td>
<td>40%</td>
<td>57%</td>
<td>Hypertension.</td>
<td>PO 10 mg/day.</td>
<td>PO 60 mg/day.</td>
</tr>
<tr>
<td>Pindolol²</td>
<td></td>
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<tr>
<td>Levatol</td>
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<td></td>
<td></td>
</tr>
<tr>
<td>Propranolol</td>
<td>(See monograph.)</td>
<td>0</td>
<td>4–6</td>
<td>&lt;0.5%</td>
<td>87%</td>
<td>Hypertension, angina pectoris, arrhythmias.</td>
<td>PO 40–80 mg/day.</td>
<td>PO 480 mg/day.</td>
</tr>
<tr>
<td>Inderal Various</td>
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<tr>
<td>Sotalol³</td>
<td>Tab 80, 120, 160, 240 mg.</td>
<td>0</td>
<td>7–15</td>
<td>80–90%</td>
<td>0%</td>
<td>Life-threatening ventricular arrhythmias.</td>
<td>PO 180 mg/day.</td>
<td>PO 240 mg/day.</td>
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<tr>
<td>Betapace</td>
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</tr>
<tr>
<td>Timolol³</td>
<td>Tab 5, 10, 20 mg.</td>
<td>0</td>
<td>4–5</td>
<td>20%</td>
<td>&lt;10%</td>
<td>Hypertension, Post-MI prophylaxis.</td>
<td>PO 20 mg/day.</td>
<td>PO 60 mg/day.</td>
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<tr>
<td>Blockadren</td>
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</table>

²Acebutolol, carteolol, penbutolol, and pindolol have intrinsic agonist (sympathomimetic) activity (ISA).
³Carvedilol has α₁-blocking actions. Labetalol has potent α₁-blocking actions (ratio of α₁- to β-blockade 1:3 and 1:7 with PO and IV, respectively).
⁴Sotalol also has type III antiarrhythmic properties.

From references 123, 221, 227–232 and product information.
Calcium-Channel Blocking Drugs

Diltiazem Hydrochloride

**Pharmacology.** Diltiazem is a calcium-channel blocking drug that decreases heart rate, prolongs AV nodal conduction, and decreases arteriolar and coronary vascular tone. It also has negative inotropic properties. Diltiazem is effective in symptomatic angina pectoris, essential hypertension, and supraventricular tachycardias. It also can reduce early reinfarction rates in patients with non–Q-wave MI and normal left ventricular functions. (See Calcium-Channel Blocking Drugs Comparison Chart.)

**Administration and Adult Dosage.** IV loading dose 0.25 mg/kg (about 20 mg) over 2 min; can repeat in 15 min with 0.35 mg/kg (about 25 mg). IV infusion 5–15 mg/hr, titrated to ventricular response. PO for angina 30–60 mg q 6–8 hr initially; dosages up to 480 mg/day may be required for symptomatic relief of angina.\(^{233,234}\) 180–300 mg once daily with Cardizem CD. PO for hypertension 120–240 mg/day initially in 2 divided doses using Cardizem SR, or 180–300 mg once daily using Cardizem CD or Dilacor XR, titrated to clinical response; maintenance dosages of 180–480 mg/day are usually necessary.

**Special Populations.** Pediatric Dosage. Safety and efficacy are not established. PO 1.5–2 mg/kg/day in 3–4 divided daily doses up to a maximum of 3.5 mg/kg/day.\(^4\)

Geriatric Dosage. Same as adult dosage but titrate dosage slowly.

Other Conditions. Patients with liver disease may require lower dosages; titrate to clinical response.

**Dosage Forms.** Tab 30, 60, 90, 120 mg; SR Cap (12 hr; Cardizem SR, various) 60, 90, 120 mg; SR Cap (24 hr; Cardizem CD) 120, 180, 240, 300 mg; (24 hr; Dilacor XR) 120, 180, 240 mg; (24 hr; Tiazac) 120, 180, 240, 300, 360 mg; SR Tab 120, 180, 240 mg (Tiamate); Inj 5 mg/mL; SR Tab 180 mg with enalapril 5 mg (Teczem).

**Patient Instructions.** Report dizziness, leg swelling, or shortness of breath. (For angina) Maintain a diary to document the numbers of episodes of chest pain and sublingual nitroglycerin tablets used.

**Missed Doses.** Take this drug at regular intervals. If you miss a dose, take it as soon as you remember. If it is about time for the next dose, take that dose only. Do not double the dose or take extra.

**Pharmacokinetics.** Onset and Duration. PO onset 0.5–3 hr, duration 6–10 hr;\(^{235}\) 12–24 hr with SR cap, depending on the product.

Serum Levels. Levels $>95$ μg/L (230 nmol/L) are necessary to cause hemodynamic changes, but their clinical usefulness is questionable.\(^{236}\) Levels of dезaceteyldiltizem are similar to those of diltiazem.\(^{237}\)

Fate. Oral bioavailability is 38 ± 11% with the first dose and 90 ± 21% with long-term therapy.\(^{237}\) The drug is 78 ± 3% bound to plasma proteins; $V_d$ is 5.3 ± 1.7 L/kg;\(^{237}\) Cl is 0.72 ± 0.3 L/hr/kg.\(^{10}\) Enterohepatic recycling occurs. The drug is
almost entirely metabolized by the liver, with only 1–3% excreted unchanged in urine. One metabolite, desacetyldiltiazem, has 40–50% the activity of diltiazem. Metabolites are excreted primarily in the feces.

\( t_1/2 \) \( \alpha \) phase 2–5 min; \( \beta \) phase 4.9 ± 0.4 hr,\(^{235,237}\) longer in the elderly.\(^{234}\) \( \beta \) phase (desacetyldiltiazem) 6.1 ± 1.2 hr.\(^{237}\)

**Adverse Reactions.** Frequency of side effects is dose related. Headache, flushing, dizziness, and edema occur frequently. Sinus bradycardia and AV block occur frequently, often in association with concomitant \( \beta \)-blockers.\(^{234}\) CHF can worsen in patients with underlying left ventricular dysfunction. A variety of skin reactions have been occasionally reported.\(^{234}\) Hepatitis occurs rarely.

**Contraindications.** Second- or third-degree block or sick sinus syndrome without a ventricular pacemaker; symptomatic hypotension or severe CHF, acute MI, or pulmonary congestion; atrial fibrillation with accessory AV pathway.

**Precautions.** Use caution with concomitant use of \( \beta \)-blockers in patients with underlying CHF, especially those with poor left ventricular function.\(^{234}\)

**Drug Interactions.** Cimetidine and propranolol increase diltiazem serum levels.\(^{234}\) Diltiazem inhibits CYP3A4 and the metabolism of many drugs, including carbamazepine, cyclosporine, and theophylline.\(^{234}\) It also inhibits P-glycoprotein.\(^{238}\)

**Parameters to Monitor.** Monitor blood pressure, heart rate, and ECG, especially when initiating therapy. Watch for symptoms of hypotension and CHF. Serial treadmill exercise tests can assess efficacy in angina. Monitor the number of episodes of chest pain and SL nitroglycerin used.

**NIFEDIPINE** Adalat, Procardia

**Pharmacology.** Nifedipine is a dihydropyridine calcium-channel blocking drug with potent arterial and coronary vasodilating properties. A reflex increase in sympathetic tone (in response to vasodilation) counteracts the direct depressant effects on SA and AV nodal conduction. This renders nifedipine ineffective in the treatment of supraventricular tachycardias. It is used for vasospastic and chronic stable angina and in the treatment of hypertension. (See Calcium-Channel Blocking Drugs Comparison Chart.)

**Administration and Adult Dosage.** PO for angina (Cap) 10 mg tid initially, increasing to a usual maximum of 20–30 mg tid or qid; dosages above 180 mg/day are not recommended. PO for hypertension (SR Tab only) 30–60 mg/day initially, increasing up to 120 mg/day pm. PO for severe hypertension (non-SR) 10 mg, may repeat pm in 20 min. The capsule can be punctured or bitten and swallowed, usually resulting in a more rapid onset than SL administration.\(^{239}\)

**Special Populations.** Pediatric Dosage. Safety and efficacy not established. PO for hypertensive crisis 0.25–0.5 mg/kg q 4–6 hr.\(^{4}\)

**Geriatric Dosage.** Same as adult dosage.

**Other Conditions.** Patients with liver disease might require lower dosages;\(^{240}\) titrate to clinical response.

**Dosage Forms.** Cap 10, 20 mg; SR Tab 30, 60, 90 mg.
**Patient Instructions.** Report flushing, edema, dizziness, or increased frequency of chest discomfort. Do not split, chew, or crush sustained-release tablets. A sustained-release tablet core in the stool does not indicate lack of absorption. Maintain a diary to document the number of episodes of chest pain and sublingual nitroglycerin tablets used.

**Missed Doses.** Take this drug at regular intervals. If you miss a dose, take it as soon as you remember. If it is about time for the next dose, take that dose only. Do not double the dose or take extra.

**Pharmacokinetics. Onset and Duration.** PO onset 0.5–2 hr; duration (Cap) 4–8 hr; (SR Tab) 12–24 hr. PO (punctured capsule) onset 10–20 min; duration 3–4 hr.

**Serum Levels.** (Therapeutic) >90 μg/L (260 nmol/L), although clinical utility is questionable.²⁴¹

**Fate.** Bioavailability is 52 ± 37% in normals and 91 ± 26% in cirrhosis because of extensive and variable first-pass hepatic elimination.²⁴⁰ It is 96 ± 1% bound to plasma proteins; $V_d$ is 0.8 ± 0.2 L/kg.²⁴¹,²⁴² CI is 0.42 ± 0.12 L/hr/kg.¹⁰ Nifedipine is almost entirely eliminated by hepatic metabolism via the CYP3A4 isozyme, which is present in variable amounts (but is not a true polymorphism).²⁴³ Only traces of drug are excreted unchanged in urine.²⁴¹

$\tau_{1/2}$. α phase 4–7 min; β phase 2 ± 0.4 hr.²⁴¹,²⁴²

**Adverse Reactions.** Most side effects relate to vasodilatory actions and occur frequently; symptoms include dizziness (with or without hypotension), flushing, and headache. These types of side effects seem less frequent with SR dosage forms.²⁴⁴ Avoid long-term treatment of hypertension with immediate-release products because they can increase mortality.¹⁷³,²⁴⁵ Edema occurs frequently and is related to venous pooling and usually not exacerbation of CHF. Nifedipine paradoxically can worsen anginal chest pain, possibly because of a reflex increase in sympathetic tone or redistribution of coronary blood flow away from ischemic areas. Acute, reversible renal failure can occur in patients with chronic renal insufficiency;²⁴⁶ rare reactions include hepatitis and hyperglycemia.

**Contraindications.** Symptomatic hypotension.

**Precautions.** Use with caution in unstable angina pectoris when used alone (ie, without a β-blocker) and in patients with CHF caused by systolic dysfunction because mortality can be increased.²⁴⁵,²⁴⁷ Do not use immediate-release products to treat hypertension. Nifedipine has an antiplatelet action and can increase bleeding time.²⁴⁵ Nifedipine can worsen symptoms of obstructive cardiomyopathy.

**Drug Interactions.** Barbiturates increase nifedipine metabolism. Cimetidine can increase nifedipine serum levels. Nifedipine occasionally increases PT in patients on oral anticoagulants. Nifedipine and IV magnesium sulfate can cause neuromuscular blockade and hypotension.

**Parameters to Monitor.** Monitor blood pressure and heart rate, especially when initiating therapy. Observe for symptoms of hypotension and edema. Serial treadmill exercise tests can assess efficacy.

**Notes.** Other potential uses for nifedipine are migraine prophylaxis, achalasia, and Raynaud’s phenomenon.
Pharmacology. Verapamil is a calcium-channel blocking drug that prolongs AV nodal conduction. It is used to convert re-entrant supraventricular tachycardias and slow ventricular rate in atrial fibrillation or flutter. Because it decreases contractility and arteriolar resistance, it is used in angina caused by coronary obstruction or vasospasm. Verapamil also is effective in the treatments of hypertension, hypertrophic obstructive cardiomyopathy, and migraine prophylaxis. (See Calcium-Channel Blocking Drugs Comparison Chart.)

Administration and Adult Dosage. PO for angina 80–120 mg tid initially, increasing at daily (for unstable angina) or weekly intervals to a maximum of 480 mg/day. PO for hypertension usually 240 mg/day using SR tablet; SR dosages of 120 mg/day to 240 mg bid have been used. Covera HS is designed to be taken hs. PO for migraine prophylaxis 160–320 mg/day.

IV for supraventricular arrhythmias 5–10 mg (0.075–0.15 mg/kg) over at least 2 min (3 min in elderly); can repeat with 10 mg (0.15 mg/kg) in 30 min if arrhythmia is not terminated or desired endpoint is not achieved. IV constant infusion 5–10 mg/hr.

Special Populations. Pediatric Dosage. PO 4–8 mg/kg/day in 3 divided doses. IV (<1 yr) 0.1–0.2 mg/kg; (1–15 yr) 0.1–0.3 mg/kg, to a maximum of 5 mg over 2–3 min.

Geriatric Dosage. Same as adult dosage but administer over 3 min.

Other Conditions. Dosage might need to be decreased in patients with liver disease; titrate to clinical response.

Dosage Forms. Tab 40, 80, 120 mg; SR Tab 120, 180, 240 mg; SR Cap 100, 120, 180, 200, 240, 300 mg; Inj 2.5 mg/mL; SR Tab 180 mg with trandolapril 2 mg, 240 mg with trandolapril 1, 4 mg (Tarka).

Patient Instructions. Report any dizziness, shortness of breath, or edema. Constipation occurs often. Maintain a diary to document the number of episodes of chest pain and sublingual nitroglycerin tablets used.

Missed Doses. Take this drug at regular intervals. If you miss a dose, take it as soon as you remember. If it is about time for the next dose, take that dose only. Do not double the dose or take extra.

Pharmacokinetics. Onset and Duration. IV onset immediate; duration 2–6 hr, up to 12 hr with long-term use.

Serum Levels. 50–400 μg/L (100–800 nmol/L), although therapeutic range is not well established.

Fate. Although the drug is well absorbed orally, only 22 ± 8% is bioavailable because of extensive first-pass elimination; bioavailability increases in liver disease. Covera-HS provides a 4- to 5-hr delay before releasing the drug. Verapamil has stereospecific pharmacology and pharmacokinetics; L-verapamil is a more potent AV nodal blocking drug, but it undergoes greater first-pass metabolism. Norverapamil is an active metabolite. Verapamil is about 90 ± 2% bound to plasma proteins, with the more active L-isomer having a greater unbound fraction. Vd is 5 ± 2 L/kg and increases in liver disease; Cl is 0.9 ± 0.36 L/hr/kg. About 1% is excreted unchanged in urine.
(Verapamil) α phase 5–30 min; β phase 4 ± 1.5 hr; can increase during long-term use; 13.6 ± 3.9 hr in severe liver disease; (norverapamil) 8 ± 1.9 hr.\(^{10,251,253}\)

**Adverse Reactions.** Constipation occurs frequently (5–40%), particularly in elderly patients. CHF can occur in patients with left ventricular dysfunction. Serious hemodynamic side effects (eg, severe hypotension) and conduction abnormalities (eg, symptomatic bradycardia or asystole) have been reported; these reactions usually occur when the patient is concurrently receiving a β-blocker or has underlying conduction disease.\(^{254}\) Infants appear to be particularly susceptible to arrhythmias. IV calcium (gluconate or chloride salts, 10–20 mL of a 10% solution) and/or isoproterenol can, in part, reverse these adverse effects.\(^{254}\) The administration of IV calcium before verapamil can prevent hypotension without abolishing the antiarrhythmic actions.\(^{255}\)

**Contraindications.** Shock or severely hypotensive states; second- or third-degree AV nodal block; sick sinus syndrome, unless functioning ventricular pacemaker is in place; hypotension or CHF unless caused by supraventricular tachyarrhythmias amenable to verapamil therapy; atrial fibrillation and an accessory AV pathway.

**Precautions.** Use caution with any wide-QRS tachycardia; severe hypotension and shock can ensue if the tachycardia is ventricular in origin. Use with caution in combination with oral β-blockers and poor left ventricular function.

**Drug Interactions.** Verapamil can increase serum levels of several drugs, including carbamazepine, cyclosporine, digoxin (probably by inhibiting P-glycoprotein\(^{238}\)), and theophylline. Barbiturates and rifampin can increase verapamil metabolism.

**Parameters to Monitor.** Monitor blood pressure and ECG continuously during IV administration. Pay particular attention to signs and symptoms of CHF and hypotension. Also, monitor the ECG for PR prolongation and bradycardia.
<table>
<thead>
<tr>
<th>DRUG</th>
<th>DOSAGE FORMS</th>
<th>ADULT DOSAGE</th>
<th>CONTRACTILITY</th>
<th>HEART RATE</th>
<th>AV NODAL CONDUCTION</th>
<th>VASCULAR RESISTANCE</th>
</tr>
</thead>
<tbody>
<tr>
<td>Amlodipinea</td>
<td>Tab 2.5, 5, 10 mg.</td>
<td>PO for hypertension or angina 5–10 mg/day.</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>↓↓</td>
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<tr>
<td>Norvasc</td>
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<tr>
<td>Bepridilb</td>
<td>Tab 200, 300, 400 mg.</td>
<td>PO for refractory angina 200–400 mg/day.</td>
<td>↓↓</td>
<td>↓↓</td>
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<td>Vasocor</td>
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<tr>
<td>Diltiazemc</td>
<td>(See monograph.)</td>
<td>(See monograph.)</td>
<td>↓</td>
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<tr>
<td>Cardizem</td>
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<td>Dilacem</td>
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<td>Cardene</td>
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<td>Various</td>
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<tr>
<td>Felodipinef</td>
<td>SR Tab 2.5, 5, 10 mg.</td>
<td>PO for hypertension 2.5–20 mg once daily.</td>
<td>↑±</td>
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<tr>
<td>Plendil</td>
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<tr>
<td>Isradipinef</td>
<td>Cap 2.5, 5 mg</td>
<td>PO for hypertension 2.5–10 mg bid or SR 5–10 mg once daily.</td>
<td>↑±</td>
<td>↑±</td>
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<tr>
<td>DynaCirc</td>
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<tr>
<td>Nicardipinef</td>
<td>Cap 20, 30 mg</td>
<td>PO for angina or hypertension 20–40 mg tid or SR 30–60 mg q 12 hr. or IV for hypertension 5–15 mg/hr.</td>
<td>↑±</td>
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<tr>
<td>Cardene</td>
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<td>Various</td>
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<tr>
<td>Nifedipinef</td>
<td>Cap 10, 20 mg</td>
<td>PO for hypertension (See monograph.)</td>
<td>↑±</td>
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<tr>
<td>Adalat</td>
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<tr>
<td>Procardia</td>
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(continued)
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<tr>
<th>DRUG</th>
<th>DOSAGE FORMS</th>
<th>ADULT DOSAGE</th>
<th>CONTRACTILITY</th>
<th>HEART RATE</th>
<th>AV NODAL CONDUCTION</th>
<th>VASCULAR RESISTANCE</th>
</tr>
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<tbody>
<tr>
<td>Nimodipine&lt;sup&gt;a&lt;/sup&gt;</td>
<td>Cap 30 mg.</td>
<td>PO postsubarachnoid hemorrhage 60 mg q 4 hr for 21 days.</td>
<td>0/↑</td>
<td>0/↑</td>
<td>0/↑</td>
<td>↓↓ ↓ ↓</td>
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<td>Nimotop</td>
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<tr>
<td>Nisoldipine&lt;sup&gt;b&lt;/sup&gt;</td>
<td>SR Tab 10, 20, 30, 40 mg.</td>
<td>PO for hypertension</td>
<td>0/↑</td>
<td>0/↑</td>
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<tr>
<td>Sular</td>
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<td>SR 20–40 mg once daily.</td>
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<tr>
<td>Verapamil&lt;sup&gt;c&lt;/sup&gt;</td>
<td>(See monograph.)</td>
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<td>Calan</td>
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<td>Isoptin</td>
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<td>Verelan</td>
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↑ = increase; ↓↓ = marked decrease, ↓ = decrease, 0 = no change.

<sup>a</sup>Selective vascular actions.

<sup>b</sup>Complex pharmacology with probable sodium- and potassium-channel blockade (quinidine-like).

<sup>c</sup>Vascular and electrophysiologic actions.

<sup>d</sup>Predominantly vascular actions.

From reference 256 and product information.
**Hypolipidemic Drugs**

**Class Instructions. Hypolipidemics.** There is a strong relationship between elevated serum cholesterol and death caused by coronary heart disease (CHD). Lowering cholesterol decreased events related to CHD and can slow or even reverse atherosclerosis. These effects are associated with a decrease in CHD mortality. In general, each 1% decrease in serum cholesterol results in a 2% decrease in the risk of coronary events. Hypolipidemic drugs must be taken daily to achieve these results. Drug therapy does not eliminate the need for appropriate diet and other measures such as weight reduction (if appropriate), smoking cessation, and physical activity. Use of estrogen and progestin in postmenopausal women also has beneficial effects on lipoprotein levels, but the overall risk/benefit assessment of hormone-replacement therapy remains controversial. Depending on their overall state of health, elderly patients can benefit from secondary prevention with hypolipidemic therapy. Hypolipidemic drug therapy (with the exception of niacin) has been associated with an increase in cancer in animals, but it is not known if they have this effect in humans.

**Missed Doses.** Take this drug at regular intervals. If you miss a dose, take it as soon as you remember. If it is about time for the next dose, take that dose only. Do not double the dose or take extra.

**CHOLESTYRAMINE RESIN Questran, Various**

**Pharmacology.** Cholestyramine is a bile acid sequestrant that acts as an anion exchange resin; it releases chloride ions and adsorbs bile acids in the intestine to form a nonabsorbable complex that is excreted in feces. The resulting increase in activity of hepatic low-density lipoprotein cholesterol (LDL-c) receptors leads to the oxidation of cholesterol to form new bile acids. Despite a compensatory increase in hepatic cholesterol synthesis, total serum cholesterol and LDL-c levels are reduced by 15–30%. The increase in cholesterol synthesis sometimes results in an increase in VLDL cholesterol levels, which can increase triglyceride levels by 10–50%. Cardioprotective HDL-c levels can increase by 3–8%.

**Administration and Adult Dosage.** PO for hyperlipidemia 4 g daily–bid initially, increasing slowly to a maintenance dosage of 8–16 g/day in 1–6 (usually 2) divided doses, to a maximum of 24 g/day. Compliance appears to be best in the range of 8–10 g/day in 1–2 divided doses. PO for treatment of cholestatic pruritus 4–8 g/day is usual. PO for treatment of relapsing enterocolitis caused by *Clostridium difficile* 4 g tid or qid (with or without vancomycin) has been used.

**Special Populations. Pediatric Dosage.** Limited data are available, especially concerning long-term use. Drug therapy is generally reserved for children at least 10 yr old, initiated at the lowest possible dosage, and gradually increased until the desired response is achieved. Base initial dosage on serum LDL-c level rather than body weight, and adjust dosage based on response: PO for hyperlipidemia (LDL-c <195 mg/dL) 4 g/day; (LDL-c 195–235 mg/dL) 8 g/day; (LDL-c 236–280 mg/dL) 12 g/day; (LDL-c >280 mg/dL) 16 g/day. (See Precautions.)
**Geriatric Dosage.** Initiate therapy at lowest possible dosage and slowly titrate to desired effect. Maximum dosage might not be required or tolerated.

**Other Conditions.** In patients with histories of constipation, start at the low end of the dosage range. In patients with GI intolerance, reduce dosage and increase gradually. (See Adverse Reactions.)

**Dosage Forms.** Pwdr 4 g resin/9 g powder (Questran); 4 g resin/5, 5.5, 5.7 g powder (Questran Light, various); Tab 1 g.

**Patient Instructions.** (See Hypolipidemics Class Instructions.) It is preferable to take this drug before meals, but you can adjust the time of the dosages around the scheduling of other oral medications. Take other oral medications at least 1 hour before or 4–6 hours after taking cholestyramine. Do not take dry; mix each packet or level scoopful with at least 60–180 mL (2–6 fluid ounces) of water or noncarbonated beverage, highly fluid soup, or pulpy fruit such as applesauce or crushed pineapple. You can experiment with different products and vehicles to determine your preference based on taste, cost, and caloric restrictions. Mixtures can be refrigerated to improve palatability but do not cook because the drug can be inactivated. This drug frequently causes constipation. If this becomes a problem, contact your physician or pharmacist to discuss measures to minimize constipation. It can cause other gastrointestinal symptoms that usually decrease over time.

**Pharmacokinetics.**

**Onset and Duration.** Reduction in cholesterol begins the first month.

**Fate.** It is not absorbed from the GI tract. Resin and complex are excreted in the feces.

**Adverse Reactions.** Almost 70% of patients experience at least one GI side effect.\(^{261}\) Constipation frequently occurs, especially with higher dosages, in the elderly and patients with previous constipation; fecal impaction is rare. Nausea, heartburn, abdominal pain, bloating, steatorrhea, and belching also occur frequently but tend to decrease over time.\(^{257}\) GI side effects tend to be milder in children than in adults. Hemorrhoids can be aggravated or develop. Rash can occur. Chloride absorption in place of bicarbonate can lead to hyperchloremic acidosis, especially in children, and calcium excretion can increase. Absorption of vitamins D and K can be impaired, leading to osteomalacia and bleeding, respectively. Absorption of folic acid also can be impaired, especially in children. Alimentary cancers in rats are somewhat more prevalent with cholestyramine treatment because of enhancement of other carcinogens, but the importance of this in humans is unknown.\(^{262}\)

**Contraindications.** Complete biliary obstruction.

**Precautions.** Pregnancy and lactation because of possible malabsorption of fat-soluble vitamins. Avoid constipation in patients with symptomatic coronary artery disease. Constipation can be controlled by reducing dosage, slowly titrating dosage, increasing dietary fiber, or using stool softeners. Avoid use in the presence of diverticular disease and local intestinal tract lesions because constipation can be a problem.\(^{263}\) Discontinue if a clinically important elevation in serum triglycerides occurs. Vitamin supplementation might be needed with high dosage or long-term therapy. Children in particular might need multivitamins with folate.
and iron.\textsuperscript{260} Patients with osteoporosis might need to restrict dietary chloride to limit calcium excretion.\textsuperscript{264} Phenylketonurics should avoid Questran Light because it contains aspartame.

**Drug Interactions.** Absorption of many drugs can be delayed or reduced, including acetaminophen, coumarin anticoagulants, digoxin, furosemide, gemfibrozil, hydrocortisone, oral hypoglycemic drugs,\textsuperscript{265} iron, loperamide, methotrexate, naproxen, penicillin G, phenobarbital, oral phosphate supplements, pravastatin, propranolol, tetracyclines, thyroid hormones, thiazides, and vancomycin. Monitor for concurrent drug therapy effects when initiating and altering sequestrant therapy, particularly for drugs with a narrow therapeu tic index.

**Parameters to Monitor.** Monitor LDL-c and triglycerides 4 weeks and 3 months after initiation of therapy. If therapy goals are achieved, monitor q 4 months unless adverse effects are suspected. Periodically monitor hemoglobin and serum folic acid during long-term therapy. Monitor efficacy of and appropriate tests for concurrent drug therapy that might be affected by cholestyramine. In children, monitor serum concentrations of vitamins A, D, and E and erythrocyte folate, liver function tests, and CBC annually.\textsuperscript{260}

**Notes.** Bile acid sequestering resins are indicated as an adjunct to diet for primary hypercholesterolemia (types IIA and IIB) in patients for whom hypertriglyceridemia is not a primary concern (triglyceride levels <300 mg/dL).\textsuperscript{264} Bile acid sequestrants moderately lower LDL-c compared with some of the other hypolipidemic drugs but are considered safer because they are not absorbed. These drugs can be particularly useful with moderately elevated LDL-c and when the risk of CHD is low and long-term safety is of concern (eg, primary prevention and in young men and premenopausal women).\textsuperscript{257,266} They can be used in combination with other hypolipidemic drugs for additive effects when a larger decrease in LDL-c is required. Long-term use reduces cardiovascular morbidity and mortality, including the incidence of first heart attacks. Because over one-third of patients discontinue bile acid sequestrants in the first year, primarily because of adverse effects, conservative dosage titration, education, and support are needed to manage and avoid adverse effects.\textsuperscript{261,267} Maximum dosage is rarely needed.\textsuperscript{258} Low-dose therapy (8–10 g/day) appears to be best tolerated\textsuperscript{257} and the most cost effective, alone or in combination therapy.\textsuperscript{268} Increased dosage can increase adverse effects without meaningful decreases in cholesterol. Resins are not effective in patients with homozygous familial hypercholesterolemia.\textsuperscript{258} (See Recommendations for Initiation of Drug Therapy in Hypercholesterolemia Chart.)

The resins also are used to reduce pruritus caused by dermal deposition of bile acids in patients with partial biliary obstruction, and cholestyramine has been used to treat relapsing *Clostridium difficile* colitis.\textsuperscript{269,270} Interference with digoxin absorption suggests a possible role in the management of the mild intoxication caused by these drugs; however, do not rely on cholestyramine alone in cases of severe digoxin toxicity.\textsuperscript{265} Questran contains 14 kcal/9 g packet or scoop; Questran Light is flavored with aspartame and contains 1.6 kcal and 16.8 mg of phenylalanine/5 g packet or scoop.
Pharmacology. Colesevelam is a nonabsorbed, polymeric, lipid-lowering agent that binds intestinal bile acids, resulting in the increased clearance of LDL-c and a reduction of total cholesterol. Unlike cholestyramine and colestipol, colesevelam is not an anion exchange resin but binds bile acids and impedes their reabsorption. Clinical trials have demonstrated a mean LDL-c reduction of 15–18% after 24 weeks of therapy. HDL-c was increased by approximately 3% and triglyceride levels were elevated 4–5% compared with placebo.

Administration and Adult Dosage. PO for hyperlipidemia 3 tablets bid with meals or 6 tablets once daily with a meal.²⁷¹ PO combination therapy for hyperlipidemia 4–6 tablets/day is safe and effective when coadministered with an HMG-CoA reductase inhibitor. The drugs can be administered together or separately.²⁷¹

Special Populations. Pediatric Dosage. Safety and efficacy not established.

Geriatric Dosage. Same as adult dosage.

Dosage Forms. Tab 625 mg.

Patient Instructions. Take this drug with meals for maximum benefit.

Pharmacokinetics. Onset and Duration. Maximum effect occurs after 2 weeks.²⁷¹

Fate. Colesevelam is not absorbed orally. It is excreted unchanged in the feces.

Adverse Reactions. Unlike cholestyramine and colestipol, colesevelam is generally well tolerated. GI effects, including flatulence, constipation, diarrhea, nausea, and dyspepsia, are the most common side effects, but the frequency is similar to that of placebo.²⁷¹,²⁷²

Contraindications. Bowel obstruction.

Precautions. Caution in patients with elevated triglyceride levels or GI disorders (ie, GI motility disorders, dysphagia, swallowing disorders, or recent GI surgery).

Drug Interactions. Colesevelam does not affect the bioavailability of digoxin, lovastatin, metoprolol, quinidine, valproic acid, or warfarin. The bioavailability of SR verapamil can be reduced by colesevelam. Colesevelam does not interfere with the lipid-lowering activity of the HMG-CoA reductase inhibitors. Colesevelam did not appear to affect the bioavailability of vitamin A, D, E, or K during clinical trials of up to 1 yr.²⁷¹ The manufacturer states that caution should be exercised when treating patients with a susceptibility to vitamin K or fat-soluble vitamin deficiencies.

Parameters to Monitor. Monitor serum total cholesterol, LDL-c, and triglyceride levels initially and periodically during therapy.

Notes. The tolerability and the apparent lack of GI side effects can make colesevelam a good alternative to other bile acid binding agents and potentially the drug of choice in this class.²⁷¹ It is a good choice for use with an HMG-CoA reductase inhibitor in patients with inadequate responses to the maximal HMG-CoA dose.
Pharmacology. Colestipol is a bile acid sequestrant similar to cholestyramine, with equivalent lipid-lowering effects in most patients. Selection of a bile acid sequestrant is generally based on patient preference and cost. The palatability of cholestyramine–vehicle combinations is often preferred over colestipol granules, although colestipol tablets are well tolerated. A 5 g dose of colestipol lowers cholesterol in an amount equivalent to 4 g of cholestyramine; a 4 g dose of colestipol tablets is about equivalent to 5 g of the granules.273–279 (See Hypolipidemic Drugs Comparison Chart and Recommendations for Initiation of Drug Therapy in Hypercholesterolemia Chart.)

Adult Dosage. PO for hypercholesterolemia (Granules) 5 g bid initially, increasing in 5 g/day increments at 1- to 2-month intervals to a maximum of 30 g/day in 1–4 doses; (tablets) 2 g bid initially, increasing in 2 g/day increments at 1- to 2-month intervals to a maximum of 16 g/day. (See Adverse Effects.) PO for relapsing enterocolitis caused by Clostridium difficile 5 g q 12 hr has been used with oral vancomycin but avoid coadministration with vancomycin to prevent vancomycin binding.

Dosage Forms. Granules 5 g resin/7.5 g packets and bulk containers; Tab 1 g.

Adverse Effects. Adverse effects, precautions, monitoring instructions, and drug interactions are similar to those of cholestyramine. Patients with moderate hypercholesterolemia who cannot tolerate colestipol granules due to GI side effects can benefit from one-half the colestipol dose mixed with 2.5 g of psyllium.

Fenofibrate is a fibric acid derivative indicated for the treatment of type IV and V hyperlipidemias. It reduces serum LDL-c by 17–35% and triglycerides by 15–43% and increases HDL. It appears to act by enhancing lipoprotein lipase activity, inhibiting VLDL synthesis, and reducing cholesterol synthesis, possibly by inhibiting acyltransferase activity. It also can reduce platelet aggregation and decrease serum uric acid.280–282

Adult Dosage. PO for type IV or V hyperlipidemia in those at risk of pancreatitis 67 mg/day initially, increasing q 4–8 weeks to a maximum of 201 mg/day. Take doses with a meal.

Dosage Forms. Cap 67, 134 mg.

Pharmacokinetics. After absorption, fenofibrate is hydrolyzed to the active drug, fenofibric acid, which is more than 99% bound to plasma proteins. It is excreted predominantly unchanged in urine with a half-life of 20 hr, which is prolonged in renal dysfunction.

Adverse Reactions. Side effects include GI disturbances, skin rash, muscle pain, and headache. Elevations in serum transaminases and CPK have occurred. Cholelithiasis has been reported, but it is not clear if the frequency is as great as with clofibrate. Avoid fenofibrate in those with liver, gallbladder, or kidney disease.
Pharmacology. Gemfibrozil is a fibric acid derivative that decreases triglyceride and VLDL-c concentrations and increases HDL-c concentrations. Effects on LDL-c are variable. LDL-c can increase in some patients, especially those with type IV hyperlipoproteinemia. Its exact mechanism is unclear, but it appears to act through many mechanisms. There is increased secretion of cholesterol into bile, increased affinity of LDL receptors for LDL particles, activation of lipoprotein lipase, inhibition of triglyceride synthesis, suppression of free fatty acid release from adipose tissue, and a change in LDL-c toward a potentially less atherosclerotic form.

Administration and Adult Dosage. PO as a hypolipidemic 600 mg bid.

Special Populations. Pediatric Dosage. Safety and efficacy not established.

Geriatric Dosage. Initiate therapy at lowest possible dosage and slowly titrate to desired effect. Maximum dosage might not be required or tolerated.

Other Conditions. Some investigators advise decreasing dose by one-half with Clcr of 20–50 mL/min.

Dosage Forms. Tab 600 mg.

Patient Instructions. (See Hypolipidemics Class Instructions.) Take doses 30 minutes before morning and evening meals. Gemfibrozil can slightly increase the risk of cancer and is similar to another medication that increases the risk of cancer, gallstones, and pancreatitis. You and your physician might decide that the benefit of reducing the risk of coronary heart disease is worth these other risks. Promptly report any muscle pain, tenderness, or weakness, especially if you also are taking lovastatin or a similar drug.

Pharmacokinetics. Onset and Duration. The maximum decrease in serum triglyceride and total cholesterol occurs within 4–12 weeks; lipids return to pretreatment levels after drug discontinuation.

Fate. The drug is rapidly and completely absorbed after oral administration. Mean peak serum concentrations of 15–25 mg/L (60–100 μmol/L) occur 1–2 hr after administration of 600 mg bid. Serum concentrations are directly proportional to dose. The drug is 97–98.6% bound to albumin. Cl appears to be independent of renal function. Gemfibrozil is metabolized in the liver to a number of compounds. Approximately 70% of a dose is excreted in the urine, primarily as glucuronide conjugates of the drug and metabolites; less than 2% is excreted renally as unchanged drug.

Adverse Reactions. Dyspepsia (20%), abdominal pain (10%), diarrhea (7%), fatigue (3%), and nausea and vomiting (3%) are frequent; acute appendicitis, dizziness, eczema, rash, vertigo, constipation, headache, paresthesia (all 1–2%) also occur. Occasional side effects include atrial fibrillation and elevations in liver function tests (AST, ALT, LDH, bilirubin, and alkaline phosphatase) that return to normal with drug discontinuation. Occasional mild decreases in WBC count, hematocrit, and hemoglobin occur but usually stabilize. However, there have been rare reports of severe blood dyscrasias. Serum glucose can be slightly elevated, as
can LDL-c in some patients with high triglyceride levels. Gemfibrozil can increase biliary lipogenicity and possibly increase long-term risk of cholelithiasis.\textsuperscript{258} Cholelithiasis requiring gallbladder surgery developed in 0.9\% of gemfibrozil-treated patients, compared with 0.5\% of patients in a placebo group. This excess was similar to that which occurred with clofibrate. (See Notes.) An acute infection-like syndrome characterized by arthralgia, myalgia, and myositis has occurred during therapy. Rhabdomyolysis with elevated CPK levels can precipitate acute renal failure, especially with the combination of gemfibrozil and lovastatin.\textsuperscript{263} This has occurred as early as several weeks to months after initiating therapy. Routine monitoring of CPK might not detect rhabdomyolysis in a timely manner. Many have recommended avoiding the combination of gemfibrozil and any HMG-CoA reductase inhibitor.\textsuperscript{265} Worsening of renal insufficiency has been reported with an initial Cr $>2$ mg/dL. Carcinogenesis, impairment of fertility, and development of cataracts occur in rats. (See Precautions and Notes.)

**Contraindications.** Hepatic or severe renal dysfunction; primary biliary cirrhosis; pre-existing gallbladder disease.

**Precautions.** Pregnancy, lactation. Evaluate any reports of muscle pain, tenderness, or weakness for myositis, including a determination of serum CPK. Discontinue if an adequate effect does not occur after 3 months, if cholelithiasis is suspected, or if liver function tests remain elevated.

**Drug Interactions.** Gemfibrozil can potentiate the effect of oral anticoagulants. Cholesterol-binding resins can decrease absorption of gemfibrozil. Insulin or an oral hypoglycemic might be required.\textsuperscript{283,285} Displacement of glyburide from plasma protein binding sites, an action that produces hypoglycemia, has been reported.\textsuperscript{286} Gemfibrozil and lovastatin (and possibly other HMG-CoA reductase inhibitors) together might increase the risk of myotoxicity. (See Adverse Reactions.)

**Parameters to Monitor.** Serum lipids, initially every few weeks, and then about q 3 months.\textsuperscript{266} Liver function tests and CBC q 3–6 months. Monitor serum glucose if the patient is receiving insulin or an oral hypoglycemic and prothrombin time if patient is taking an oral anticoagulant.

**Notes.** Gemfibrozil is not considered a major treatment for hypercholesterolemia because of its effects on LDL-c, but it does increase HDL-c and decrease triglycerides, so it is useful in some patients.\textsuperscript{257} Gemfibrozil is indicated for the treatment of type IV and V hyperlipidemias with very high serum triglycerides (usually $>2000$ mg/dL) in patients at risk for pancreatitis and not responding to diet. It is also indicated in type IIB patients (only those without history or symptoms of CHD) with low HDL-c and an inadequate response to weight loss, diet, exercise, and other drugs that raise HDL-c (eg, bile acid sequestrants, niacin), but is not indicated in type I or IIa hyperlipidemias or in those patients who have low HDL-c only. The Helsinki Heart Study showed a 34\% reduction in the incidence of CHD in middle-aged men (initially without CHD symptoms) treated with gemfibrozil in a 5-yr study, although total death rate was no different between treated and placebo groups.\textsuperscript{287,288} A substudy of the Helsinki Heart Study showed an increase in gallstone and gallbladder surgery in gemfibrozil-treated patients. After a 3.5-yr extension of this study, all-cause mortality was slightly higher in the original gem-
fibrozil group, primarily because of cancer deaths. An ancillary study of the Helsinki Heart Study investigated the use of gemfibrozil in patients with signs or symptoms of CHD. The rate of serious adverse cardiac advents and total mortality with gemfibrozil treatment did not significantly differ from that of placebo; however, information on key prognostic indicators and their distribution was not known. Gemfibrozil is chemically and pharmacologically similar to clofibrate. A 44% relative increase in age-adjusted, all-cause mortality occurred in a study of long-term clofibrate use related to a 33% increase in noncardiovascular disease such as malignancy, gallbladder disease, and pancreatitis. Because of the smaller size of the gemfibrozil studies, the increase in mortality in the gemfibrozil group relative to placebo might not be statistically significantly different from the excess mortality associated with clofibrate use.

**Pharmacology.** Hydroxymethylglutaryl-CoA (HMG-CoA) reductase inhibitors competitively inhibit conversion of HMG-CoA to mevalonate, an early rate-limiting step in cholesterol synthesis. A compensatory increase in LDL receptors, which bind and remove circulating LDL-C, results. Production of LDL-C also can decrease because of decreased production of VLDL or increased VLDL removal by LDL receptors. These drugs produce dose-dependent, maximum reductions in LDL of 30–40% (up to 60% with atorvastatin) and triglycerides of 10–30% and increases in HDL levels of 2–15%. Drug effects are dose dependent until the following doses are reached: 80 mg for atorvastatin, 0.3 mg for cerivastatin, 20 mg for fluvastatin and simvastatin, and 40 mg for lovastatin and pravastatin. Fluvastatin is about 30% less effective in lowering lipids than the other drugs. These drugs stabilize arterial plaques, which might be an important factor in their reduction of MI risk. They also appear to reduce the risk of bone fracture by increasing bone density, the risk of DVT and the risk of becoming diabetic.

**HMG-COA REDUCTASE INHIBITORS:**

| ATORVASTATIN | Lipitor |
| CERIVASTATIN | Baycol |
| FLUVASTATIN | Lescol, Lescol XL |
| LOVASTATIN | Mevacor |
| PRAVASTATIN | Pravachol |
| SIMVASTATIN | Zocor |

**Pharmacology.** Hydroxymethylglutaryl-CoA (HMG-CoA) reductase inhibitors competitively inhibit conversion of HMG-CoA to mevalonate, an early rate-limiting step in cholesterol synthesis. A compensatory increase in LDL receptors, which bind and remove circulating LDL-C, results. Production of LDL-C also can decrease because of decreased production of VLDL or increased VLDL removal by LDL receptors. These drugs produce dose-dependent, maximum reductions in LDL of 30–40% (up to 60% with atorvastatin) and triglycerides of 10–30% and increases in HDL levels of 2–15%. Drug effects are dose dependent until the following doses are reached: 80 mg for atorvastatin, 0.3 mg for cerivastatin, 20 mg for fluvastatin and simvastatin, and 40 mg for lovastatin and pravastatin. Fluvastatin is about 30% less effective in lowering lipids than the other drugs. These drugs stabilize arterial plaques, which might be an important factor in their reduction of MI risk. They also appear to reduce the risk of bone fracture by increasing bone density, the risk of DVT and the risk of becoming diabetic.

**Administration and Adult Dosage.** Adjust dosage at no less than 4-week intervals. PO for hyperlipidemia (Atorvastatin) 10 mg/day initially, increasing to a maximum of 80 mg/day. (Cerivastatin) 0.2–0.8 mg/day in the evening. (Fluvas- statin) 20 mg hs initially, increasing up to 80 mg/day; a slight increase in fluvas- statin LDL-C lowering occurs with a bid schedule. (Lovastatin) 20 mg (40 mg with serum cholesterol >300 mg/dL) with the evening meal initially. Start with 10 mg/day in patients requiring <20% decrease in LDL-C or when concurrent use with cyclosporine is unavoidable. Increase to a maintenance dosage of 20–80 mg/day. Do not exceed a lovastatin dosage of 20 mg/day when used with
cyclosporine. (Pravastatin) 10–20 mg hs initially, increasing to 10–40 mg/day hs.
(Simvastatin) 5–10 mg/day in the evening initially, increasing to 5–40 mg/day in
the evening.

**Special Populations.** *Pediatric Dosage.* Safety and efficacy not established.

**Geriatric Dosage.** PO (Pravastatin) 10 mg/day initially; (Simvastatin) 5 mg/day
initially; the elderly can achieve maximal reductions in LDL-c at doses ≤20 mg/day. Maximum dosage might not be required or tolerated.

**Other Conditions.** (Atorvastatin) no dosage adjustment necessary in renal impair-
ment. (Cerivastatin) start with 0.2 mg/day in moderate to severe renal impairment.
(Lovastatin) start with 10 mg/day when concurrent use with cyclosporine is un-
avoidable; maximum dosage is 20 mg/day with concurrent immunosuppressant
therapy. (Pravastatin) start with 10 mg/day with renal or hepatic impairment and
in patients concurrently taking cyclosporine; do not exceed a dose of 20 mg/day in
the latter case. (Simvastatin) start with 5 mg/day in patients requiring <20% re-
ductions in LDL-c and with severe renal insufficiency. Use dosages >20 mg/day
only with extreme caution in patients with Clcr <30 mL/min.

**Dosage Forms.** (See Hypolipidemic Drugs Comparison Chart.)

**Patient Instructions.** (See Hypolipidemics Class Instructions.) Atorvastatin can
be taken any time of day without regard to meals. Take lovastatin with the
evening meal to increase its absorption; other drugs can be taken without regard to
meals. Take pravastatin 1 hour before or 4 hours after a dose of a cholesterol-
binding resin. Promptly report any unexplained muscle pain or tenderness, espe-
cially if accompanied by malaise or fever. Avoid excessive concurrent use of alco-
hol, but abstinence is not required. Do not take these drugs during pregnancy
because of possible harm to the fetus. Inform your physician if you become or in-
tend to become pregnant.

**Pharmacokinetics.** *Onset and Duration.* Onset is within 2 weeks; peak effect is
within 4–6 weeks; cholesterol levels return to baseline after drug discontinuation.

**Fate.** Absorption is rapid. Lovastatin and simvastatin are prodrugs that undergo
extensive first-pass metabolism to active metabolites. Serum concentrations of the
active lovastatin metabolite when taken under fasting conditions are two-thirds of
that when taken with food. Absorption of other drugs is unaffected by food. Sys-
temic bioavailability of all drugs is low because of extensive (>60%) first-pass ex-
traction. Peak serum concentrations of active inhibitors are achieved in 1–4 hr for
all drugs. Protein binding is 98% for atorvastatin, 99% for cerivastatin, 55–60% for
pravastatin, 95% for simvastatin, >95% for lovastatin, and 98% for fluvastatin.
Atorvastatin, lovastatin, and simvastatin are lipophilic and cross the blood–brain
barrier; cerivastatin, fluvastatin, and pravastatin are hydrophilic and do not. All
the drugs are primarily (>70%) hepatically metabolized to active and inactive
metabolites, which then undergo extensive fecal elimination. Renal elimination
accounts for <2% for atorvastatin, 5% for fluvastatin, 10% for lovastatin, 13% for
simvastatin, 20% for pravastatin, and 26% for cerivastatin. Severe renal insuffi-
ciency (Clcr 10–30 mL/min) results in a 2-fold increase in serum concentrations of
renally excreted drugs (ie, cerivastatin, pravastatin). 257,291–297
The drugs are generally well tolerated, with discontinuation rates less than other hypolipidemic drugs. The frequency of side effects appears to be similar for all drugs. GI complaints such as diarrhea, constipation, flatulence, abdominal pain, and nausea occur in about 5% of patients. Headache (4–9%), rash (3–5%), dizziness (3–5%), and blurred vision (1–2%) are other frequent side effects. Myopathy and myositis occur rarely with single therapy and can be associated with mild elevations of CPK. Rhabdomyolysis leading to acute renal failure is a rare complication but occurs more frequently in combination with gemfibrozil, cyclosporine, or ≥1 g/day of niacin. Pravastatin can cause less myopathy than lovastatin with cyclosporine. Increases in liver function tests greater than 3 times normal occur in up to 2% of patients, but most are asymptomatic and reverse with discontinuation. There have been 62 cases of serious liver disease directly associated with statin use reported to the Food and Drug Administration. Anomalies have been reported with intrauterine exposure. Carcinogenicity occurs in mice and rats at higher than human doses, but the clinical implications are unclear.

Contraindications. Pregnancy; lactation; active liver disease; unexplained persistent elevations of serum transaminases.

Precautions. Administer to women of childbearing age only when possibility of becoming pregnant is unlikely. Use with caution in patients who consume substantial quantities of alcohol and/or have histories of liver disease. Discontinue therapy if liver function tests are >3 times normal. Consider withholding the drug in any patient with risk factors for renal failure secondary to rhabdomyolysis, such as severe acute infection; hypotension, major surgery, or trauma; severe electrolyte, endocrine, or metabolic abnormalities; or uncontrolled seizures.

Drug Interactions. Myositis and rhabdomyolysis can be more common in combination with cyclosporine (lovastatin levels are quadrupled), erythromycin, gemfibrozil, itraconazole, ketoconazole (and possibly other inhibitors of CYP3A4), or lipid-lowering dosages of niacin (>1 g/day). HMG-CoA reductase inhibitors can increase the effect of warfarin. Bile acid sequestrants can markedly decrease pravastatin oral bioavailability when taken together; take pravastatin 1 hr before or 4 hr after resin doses.

Parameters to Monitor. Obtain serum lipid and liver function tests on initiation, 6 and 12 weeks after initiation, after dosage increases, and at least semiannually thereafter. Others have recommended more frequent monitoring. Increase the frequency of monitoring if adverse effects are suspected. Routine monitoring of muscle enzymes might not adequately identify patients at risk for rhabdomyolysis but might be warranted in patients with skeletal muscle complaints and risk factors.

Notes. HMG-CoA reductase inhibitors are indicated for the treatment of type IIa and IIb hyperlipoproteinemias. They are the most effective in decreasing LDL-c, and their effect on decreasing coronary morbidity and mortality is proven. The choice of drug depends on cost, amount of cholesterol lowering desired, and risk
factors for CHD. Lovastatin, pravastatin, and simvastatin are more effective than fluvastatin and are therefore preferable when a >25% reduction in LDL-c is required.\textsuperscript{292,301} They can have additive effects with the bile acid sequestrants and have been used in combination with niacin and gemfibrozil. (See Adverse Reactions and Drug Interactions.) Cholesterol reduction with these agents can reduce the risk of stroke and total mortality.\textsuperscript{302} One large study found a reduced CHD risk with lovastatin in men and women with “average” serum cholesterol and no evidence of pre-existing heart disease.\textsuperscript{303} Pravastatin appears to reduce inflammation in vessel walls based on its effective lowering of C-reactive protein levels in patients with coronary artery disease.\textsuperscript{304} C-reactive protein concentrations can be a predictive and sensitive measure of vessel wall inflammation resulting from elevated cholesterol levels.\textsuperscript{305}

**Pharmacology.** Niacin (nicotinic acid), in dosages of 1 g/day or more, decreases serum total cholesterol, LDL-c and VLDL-c, and triglycerides, and increases HDL-c. Mean serum cholesterol and triglyceride levels are reduced by 10% and 26%, respectively. The mechanism for these effects is not entirely known but might involve inhibition of lipolysis, reduced LDL and VLDL synthesis, and increased lipoprotein lipase activity.\textsuperscript{306,307} Niacin consistently decreases lipoprotein (a).\textsuperscript{257} Niacinamide (nicotinamide) does not have hypolipidemic effects and cannot be substituted for niacin.

**Administration and Adult Dosage.** PO as a hypolipidemic 250 mg/day with evening meal initially, increasing at 4- to 7-day intervals to 1.5–2 g/day in 3 divided doses. Continue this dosage for 2 months. If necessary, the dosage can then be increased at 2- to 4-week intervals to 1 g tid; the dose can be increased again, if necessary, to reach the desired clinical effect or the maximum tolerated dosage to a usual maximum of 2 g tid. Use of >3–4 g/day does not appear to increase efficacy appreciably and is associated with increased side effects.\textsuperscript{306} (Niaspan) 375 mg hs initially, increasing as tolerated to a maximum of 2 g hs. (See Notes.) If an SR product is substituted for an immediate-release form, reduce the dosage by one-half. Risk of hepatotoxicity increases at SR doses >1.5–3 g/day, and these are not recommended.\textsuperscript{258}

**Special Populations.** **Pediatric Dosage.** Safety and efficacy not established. Although niacin is effective in reducing triglycerides and cholesterol in children and adolescents, adverse effects are common and can be severe. Some investigators have recommended niacin use be avoided in children or used only under close supervision by a lipid specialist if diet and bile acid sequestrants have failed.\textsuperscript{259,308} In these cases, niacin should generally be used in combination with diet and a bile acid sequestrant, if tolerated.\textsuperscript{259} Close monitoring is necessary. (See Adverse Reactions and Precautions.)

**Geriatric Dosage.** Initiate therapy at lowest possible dosage and slowly titrate to desired effect. Maximum dosage might not be required or tolerated.

**Patient Instructions.** Use only under medical supervision. Effects can differ with different preparations. Almost everyone experiences some flushing. Tolerance to
flushing generally occurs with time. Taking 325 mg of aspirin or 200 mg of ibuprofen (or an equivalent dose of another NSAID) 30–60 minutes before each niacin dose might reduce flushing. Taking niacin with meals also might minimize flushing and reduce GI upset. Avoid hot liquids or alcohol after taking a dose. Avoid interruptions in therapy; tolerance can be lost if therapy is interrupted, so slow resumption of the usual dosage is recommended. Avoid sudden changes in posture if you are also taking medicine for high blood pressure. Report any persistent nausea.

**Dosage Forms.** Tab 50, 100, 250, 500 mg; Elxr 10 mg/mL; SR Cap 125, 250, 400, 500 mg; SR Tab 250, 500, 750, 1000 mg. (See Notes.)

**Pharmacokinetics.**

**Onset and Duration.** The onset of triglyceride and cholesterol reductions usually occurs within several days, although some studies have demonstrated a response after a single dose. Pretreatment lipid levels return 2–6 weeks after drug discontinuation.

**Fate.** The drug is almost completely absorbed from standard formulations; peak serum levels 30–60 min after 1 g standard formulations are 15–30 μg/L (120–240 nmol/L). Niacin is largely metabolized in the liver to niacinamide (nicotinamide) and its derivatives (eg, nicotinuric acid), which can contribute to the lipid-lowering activity, especially after long-term use. The majority is excreted as unchanged drug or metabolites in urine. $t_{1/2}$ 20–48 min.

**Adverse Reactions.** Dose-related flushing of the neck and face (usually in “blush” areas) occurs in almost all patients and is related to the rate of rise of serum levels rather than the absolute serum concentrations. Tolerance can develop but disappear if therapy is interrupted. Administration with food, gradual upward dosage titration, use of a SR formulation, or premedication with 325 mg of aspirin or 200 mg of ibuprofen (or equivalent dose of another NSAID) 30–60 min before each dose can reduce flushing. Postural hypotension can occur, especially when niacin is used with antihypertensive drugs or when it is taken with alcohol or hot liquids. Vasodilatory effects can precipitate or aggravate angina. Rash, pruritus, and stomach discomfort also occur frequently; the latter can occur more commonly with SR preparations. Increases in AST, ALT, bilirubin, and LDH concentrations occur frequently and might be related to increasing the daily dosage by more than 2.5 g/month. Severe hepatotoxicity (hepatic necrosis) is rare and tends to occur with abrupt dosage increases, SR forms substituted for immediate-release products without dosage reduction, or brand interchange. Hepatotoxicity can occur at low dosages (≤3 g/day) of SR products and as soon as 2 days after initiation. Discontinue therapy if liver function tests remain over 3 times pretreatment values.

**Contraindications.** Arterial hemorrhage; severe hypotension; hepatic dysfunction; unexplained elevations of transaminases, active peptic ulcer disease.

**Precautions.** Pregnancy; lactation. Use with caution in patients with gallbladder disease or histories of liver disease, unstable angina, gout, gouty arthritis, glaucoma, or diabetes.

Nausea can be a presenting sign of hepatotoxicity. If the SR form is substituted for the immediate-release form, reduce dosage by about
one-half. If use with an HMG-CoA reductase inhibitor is unavoidable, use with extreme caution. Some formulations contain tartrazine dye, which can cause allergic reactions in sensitive patients.

**Drug Interactions.** Concurrent use with HMG-CoA reductase inhibitors can increase the risk of rhabdomyolysis. Concurrent therapy with α-adrenergic blocking antihypertensives can result in hypotension. Diet and/or dosage of oral hypoglycemic drugs or insulin might require adjustment with concurrent niacin use. Hepatotoxic drugs can have additive effects.\(^{257}\)

**Parameters to Monitor.** Monitor serum lipids q 2 weeks initially and then q 1–3 months. Periodic liver function tests, blood glucose, and serum uric acid levels are recommended, especially at dosages >1.5 g/day.\(^{257}\) Obtain liver function tests at baseline and q 6–12 weeks for the first year and then semiannually unless hepatotoxicity is suspected.

**Notes.** Indicated for type IIa, IIb, III, IV, and V hyperlipoproteinemias. Reductions in sudden cardiac deaths and fatal and nonfatal MI and an 11% decrease in mortality (compared with placebo) occur in patients treated with niacin.\(^{298}\) The cost of standard niacin formulations can be much lower than alternative drugs; the cost of SR products can be higher. SR products appear to have equal efficacy at lowering LDL-c at one-half the dosage of standard products, but the SR form might be less effective in lowering triglycerides or raising HDL at dosages equal to standard forms.\(^{257,308,310}\) Some have advocated avoidance of SR preparations altogether for treatment of hyperlipidemia because of their potentially greater hepatotoxicity.\(^{308,310}\)

Niaspan is an extended-release product that may be safer than other SR products because of its release rate. It is available as 500, 750 and 1000 mg tablets.\(^{369}\)
<table>
<thead>
<tr>
<th>DRUG</th>
<th>DOSAGE FORMS</th>
<th>ADULT ORAL DOSAGE&lt;sup&gt;a&lt;/sup&gt; RELATIVE EFFECT ON LIPIDS&lt;sup&gt;b&lt;/sup&gt; INDICATIONS BY WHO CATEGORY</th>
<th>COMMENTS</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>BILE ACID SEQUESTRANTS</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Cholestyramine</td>
<td>Pwdr 4 g resin/9 g, Questran</td>
<td>(I) 4 g daily–bid. ↓↓ ↑ ↑ IIa Constipation frequent. Can be used in children. Numerous interactions. Decrease in mortality demonstrated. Take before meals.</td>
<td></td>
</tr>
<tr>
<td>Questran Light</td>
<td>Pwdr 4 g resin/5, 5.5, various</td>
<td>1–6 doses, to a maximum of 24 g/day.↑↓ ↑↑ ↑↓↓↓↓ IV, V Avoid in liver, gallbladder, or kidney disease. Take with food.</td>
<td></td>
</tr>
<tr>
<td>Various</td>
<td>5.7 g packet or scoop, Tab 1 g.</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Colesevelam</td>
<td>Tab 625 mg. Welchol</td>
<td>6 tablets/day in 1–2 doses. ↓↓ ↑ ↑ IIa Better tolerated than cholestyramine or colestipol.</td>
<td></td>
</tr>
<tr>
<td>Colestipol HCl</td>
<td>Gran 5 g resin/7.5 g packet or scoop, Colestid</td>
<td>(I) (gran) 5 g daily–bid. (M) (gran) 15–30 g/day in 1–4 doses.↑↓ ↑↑ ↑↓↓↓↓ IIb, IV Avoid overall mortality rate with long-term therapy is increased. Take 30 min before AM and PM meals.</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Tab 1 g.</td>
<td></td>
<td></td>
</tr>
<tr>
<td>FIBRIC ACID DERIVATIVES</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Fenofibrate</td>
<td>Cap 67, 134 mg. Tricor</td>
<td>(I) 67 mg/day, (M) up to 201 mg/day. ↑↓↓ ↑↑ ↑↓↓↓↓ IV, V Avoid in liver, gallbladder, or kidney disease. Take with food.</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Gemfibrozil</td>
<td>Tab 600 mg. Lopid</td>
<td>(M) 600 mg bid. ↑↓ ↑↑ ↓↓↓↓ Ilb, IV V Overall mortality rate with long-term therapy is increased. Take 30 min before AM and PM meals.</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Various</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Clofibrate</td>
<td>Not recommended because of a large increase in overall mortality. (continued)</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
## Hypolipidemic Drugs Comparison Chart (continued)

<table>
<thead>
<tr>
<th>Drug</th>
<th>Dosage Forms</th>
<th>Oral Dosage*</th>
<th>Relative Effect on Lipids*</th>
<th>Indications by WHO Category</th>
<th>Comments</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Nicotinic Acid</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Niacin</td>
<td>Tab 50, 100, 250, 500 mg</td>
<td>(I) Titrate slowly. (See monograph.)</td>
<td>↓↓ ↑↑ ↓↓↓</td>
<td>IIa, IIb, III, IV, V</td>
<td>Frequent flushing and GI side effects. Must titrate dosage slowly. Decrease in mortality demonstrated. Avoid SR products. Low cost. Take with food or milk.</td>
</tr>
<tr>
<td>Various</td>
<td>Elixir 10 mg/mL</td>
<td>(M) 1.5–3 g/day, to a maximum of 6 g/day.</td>
<td>↓↓ ↑↑ ↓↓↓</td>
<td>IIa, IIb, III, IV, V</td>
<td></td>
</tr>
<tr>
<td></td>
<td>SR Cap 125, 250, 400, 500 mg</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>SR Tab 250, 500, 750, 1000 mg</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Inj 100 mg/mL</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>HMG-CoA Reductase Inhibitors</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Atorvastatin</td>
<td>Tab 10, 20, 40, 80 mg</td>
<td>(I) 10 mg/day. (M) 10–80 mg/day.</td>
<td>↓↓↓↓↑ ↓↓↓</td>
<td>IIa, IIb</td>
<td>Take at any time without regard to meals.</td>
</tr>
<tr>
<td>Lipitor</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Cerivastatin</td>
<td>Tab 0.2, 0.3, 0.4, 0.8 mg</td>
<td>0.2–0.8 mg/day.</td>
<td>↓↓ ↑↑ ↓↓</td>
<td>IIa, IIb</td>
<td>Take in evening without regard to meals.</td>
</tr>
<tr>
<td>Baycol</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Fluvastatin</td>
<td>Cap 20, 40 mg</td>
<td>(I) 20 mg q hs. (M) 20–80 mg q hs.</td>
<td>↓↓ ↑↑ ↓↓</td>
<td>IIa, IIb</td>
<td>Take without regard to meals. Monitor for myositis and hepatic dysfunction. Important drug interactions.</td>
</tr>
<tr>
<td>Lescol</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Lescol XL</td>
<td>SR Tab 80 mg</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

*(continued)*
**Hypolipidemic Drugs Comparison Chart (continued)**

<table>
<thead>
<tr>
<th>Drug</th>
<th>Dosage Forms</th>
<th>Initial (I) Maintenance (M)</th>
<th>Relative Effect on Lipids</th>
<th>Indications by WHO Category</th>
<th>Comments</th>
</tr>
</thead>
<tbody>
<tr>
<td>Lovastatin</td>
<td>Tab 10, 20, 40 mg.</td>
<td>(I) 20 mg/day. (M) 20–80 mg/day in 1–2 divided doses.</td>
<td>↓↓↓ ↑ ↓↓</td>
<td>IIa, IIb</td>
<td>Take with evening meal. (See Mevacor (M) 20–80 mg/day Fluvastatin.)</td>
</tr>
<tr>
<td>Pravastatin</td>
<td>Tab 10, 20, 40 mg.</td>
<td>(I) 10–20 mg q hs. (M) 10–40 mg q hs.</td>
<td>↓↓↓ ↑ ↓↓</td>
<td>IIa, IIb</td>
<td>Take without regard to meals. (See Fluvastatin.)</td>
</tr>
<tr>
<td>Rosuvastatin</td>
<td>—</td>
<td>(I) 5–10 mg/day. (M) 10–80 mg/day.</td>
<td>↓↓↓↓↓↓ ↑↑ ↓↓</td>
<td>IIa, IIb</td>
<td>Take at anytime of day.</td>
</tr>
<tr>
<td>Simvastatin</td>
<td>Tab 5, 10, 20, 40, 80 mg.</td>
<td>(I) 5–10 mg q hs. (M) 5–80 mg/day.</td>
<td>↓↓↓ ↑ ↓↓</td>
<td>IIa, IIb</td>
<td>Take without regard to meals. (See Fluvastatin.)</td>
</tr>
</tbody>
</table>

LDL = serum low-density lipoprotein cholesterol, shown to have a graded, positive relationship with coronary heart disease (CHD); HDL = serum high-density lipoprotein, shown to have a cardioprotective effect against CHD; TG = serum triglycerides, which have a positive relationship to CHD.

- Clinical trials in the elderly are limited, but the drugs are considered effective. Base decision on drug treatment on life expectancy, concomitant disease, potential adverse effects, patient desire for treatment, quality of life, assessment of coronary heart disease risk, and cost. A conservative approach is recommended; generally initiating therapy at the lowest possible dosage and slowly titrating to desired effect while closely monitoring possible side effects. Maximum dosage might not be required or tolerated.
- Arrows represent approximate relative effects based on results achieved with usual dosage. Study results differ because of differences in patient groups, administration schedules, dosages, and degrees of hypercholesteremia.
- World Health Organization (WHO) classification by characteristics of dyslipidemia.
- All dosages are expressed in terms of anhydrous resin.
- Do not substitute SR niacin products for an equal dosage of immediate-release products; use one-half the dosage of immediate-release products.

From references 266, 285, 311, and 312 and product information.
### RECOMMENDATIONS FROM THE THIRD REPORT OF THE NATIONAL CHOLESTEROL EDUCATION PROGRAM (NCEP) EXPERT PANEL ON DETECTION, EVALUATION, AND TREATMENT OF HIGH BLOOD CHOLESTEROL IN ADULTS (ADULT TREATMENT PANEL [ATP] III)

**STEP 1. Classification of lipoprotein levels**

#### ATP III CLASSIFICATION OF LDL, TOTAL AND HDL CHOLESTEROL (MG/DL)

<table>
<thead>
<tr>
<th>LDL CHOLESTEROL-PRIMARY TARGET OF THERAPY</th>
<th>TOTAL CHOLESTEROL</th>
<th>HDL CHOLESTEROL</th>
</tr>
</thead>
<tbody>
<tr>
<td>&lt;100 mg/dL</td>
<td>&lt;200 mg/dL</td>
<td>&lt;40 mg/dL</td>
</tr>
<tr>
<td>100–129 mg/dL</td>
<td>200–239 mg/dL</td>
<td>≥60 mg/dL</td>
</tr>
<tr>
<td>130–159 mg/dL</td>
<td>Borderline High</td>
<td>High</td>
</tr>
<tr>
<td>160–189 mg/dL</td>
<td>High</td>
<td>≥60 mg/dL</td>
</tr>
<tr>
<td>≥190 mg/dL</td>
<td>Very High</td>
<td></td>
</tr>
</tbody>
</table>

**STEP 2.** Identify the presence or absence of clinical atherosclerotic disease that confers a high risk for coronary heart disease (CHD) events. These include clinical CHD, symptomatic carotid artery disease, peripheral arterial disease or abdominal aortic aneurysm.

**STEP 3.** Identify major risk factors other than LDL: cigarette smoking, hypertension (BP ≥140/90 mmHg or on antihypertensive medication, HDL cholesterol <40 mg/dL, family history of premature CHD, age (men ≥45 years; women ≥55 years). Diabetes is considered a risk equivalent.

**STEP 4.** If 2+ risk factors (other than LDL) are present without CHD or CHD risk equivalent, assess 10-year (short-term) CHD risk (refer to Framingham tables).

**STEP 5.** Determine Risk Category

<table>
<thead>
<tr>
<th>RISK CATEGORY</th>
<th>LDL GOAL</th>
<th>LDL LEVEL AT WHICH TO INITIATE THERAPEUTIC LIFESTYLE CHANGES (TLC)</th>
<th>LDL LEVEL AT WHICH TO CONSIDER DRUG THERAPY</th>
</tr>
</thead>
<tbody>
<tr>
<td>CHD or CHD risk equivalents (10-yr risk &gt;20%)</td>
<td>&lt;100 mg/dL</td>
<td>≥100 mg/dL</td>
<td>≥130 mg/dL (100–129 mg/dL: drug therapy optional)</td>
</tr>
<tr>
<td>2+ risk factors (10-yr risk ≤20%)</td>
<td>&lt;130 mg/dL</td>
<td>≥130 mg/dL</td>
<td>10-yr risk 10–20%: ≥130 mg/dL; 10-yr risk &lt; 10%: ≥160 mg/dL</td>
</tr>
<tr>
<td>0–1 risk factor</td>
<td>&lt;160 mg/dL</td>
<td>≥160 mg/dL</td>
<td>≥190 mg/dL (160–189 mg/dL: LDL-lowering drug is optional)</td>
</tr>
</tbody>
</table>

**STEP 6.** Initiate therapeutic lifestyle changes (TLC): diet (refer to guidelines for precise recommendations), weight management, increased physical activity.

**STEP 7.** Consider adding drug therapy if LDL exceeds levels shown in Step 5 of table: consider drug simultaneously with TLC for CHD and CHD equivalents; consider adding drug to TLC after 3 months for other risk categories.
**RECOMMENDATIONS FROM THE THIRD REPORT OF THE NATIONAL CHOLESTEROL EDUCATION PROGRAM (NCEP) EXPERT PANEL ON DETECTION, EVALUATION, AND TREATMENT OF HIGH BLOOD CHOLESTEROL IN ADULTS (ADULT TREATMENT PANEL [ATP] III) (continued)**

**STEP 8.** Identify metabolic syndrome and treat, if present, after 3 months of TLC.

**CLINICAL IDENTIFICATION OF METABOLIC SYNDROME**

<table>
<thead>
<tr>
<th>Risk Factor</th>
<th>Defined Level</th>
</tr>
</thead>
<tbody>
<tr>
<td>Abdominal obesity</td>
<td>For men: waist circumference &gt;102 cm (40 in); For women: waist circumference &gt;88 cm (35 in)</td>
</tr>
<tr>
<td>Triglycerides</td>
<td>≥150 mg/dL</td>
</tr>
<tr>
<td>HDL cholesterol</td>
<td>For men: &lt;40 mg/dL; For women: &lt;50 mg/dL</td>
</tr>
<tr>
<td>Blood pressure</td>
<td>≥130/≥85 mmHg</td>
</tr>
<tr>
<td>Fasting glucose</td>
<td>≥110 mg/dL</td>
</tr>
</tbody>
</table>

**STEP 9.** Treat elevated triglycerides

**ATPIII—CLASSIFICATION OF SERUM TRIGLYCERIDES (TG) (MG/DL)**

<table>
<thead>
<tr>
<th>TG Level</th>
<th>Classification</th>
<th>Recommendations</th>
</tr>
</thead>
<tbody>
<tr>
<td>&lt;150 mg/dL</td>
<td>Normal</td>
<td>- Primary aim of therapy is to reach LDL goal</td>
</tr>
<tr>
<td>150–199 mg/dL</td>
<td>Borderline High</td>
<td>- Intensify weight management</td>
</tr>
<tr>
<td>200–499 mg/dL</td>
<td>High</td>
<td>- Increase physical activity</td>
</tr>
<tr>
<td>≥500 mg/dL</td>
<td>Very High</td>
<td>If triglycerides are ≥200 mg/dL after LDL goal is reached, set secondary goal for non-HDL cholesterol (total HDL) 30 mg/dL higher than LDL goal.</td>
</tr>
</tbody>
</table>

**COMPARISON OF LDL CHOLESTEROL AND NON-HDL CHOLESTEROL GOALS FOR THREE RISK CATEGORIES**

<table>
<thead>
<tr>
<th>Risk Category</th>
<th>LDL Goal</th>
<th>Non-HDL Goal</th>
<th>Recommendations</th>
</tr>
</thead>
<tbody>
<tr>
<td>CHD and CHD RE (10-yr risk for CHD &gt;20%)</td>
<td>&lt;100 mg/dL</td>
<td>&lt;130 mg/dL</td>
<td>If TG ≥500 mg/dL: reduce to prevent pancreatitis (low fat diet, weight management, a fibrate or niacin).</td>
</tr>
<tr>
<td>Multiple (2+) risk factors and 10-yr risk ≤20%)</td>
<td>&lt;130 mg/dL</td>
<td>&lt;160 mg/dL</td>
<td>Treatment of low HDL: weight management; increased physical activity; if TG = 200–499 mg/dL, then achieve non-HDL goal; if TG &lt;200 mg/dL in CHD or CHD risk equivalent, consider niacin or a fibrate.</td>
</tr>
<tr>
<td>0–1 risk factor</td>
<td>&lt;160 mg/dL</td>
<td>&lt;190 mg/dL</td>
<td></td>
</tr>
</tbody>
</table>

*From reference 266 and The National Heart Lung and Blood Institute [http://www.nhlbi.nih.gov/].*
Inotropic Drugs

DOBUTAMINE HYDROCHLORIDE

Pharmacology. Dobutamine is a synthetic sympathomimetic amine that exists as the racemic mixture of an L-isomer with predominantly \( \beta_1 \)- and \( \beta_2 \)-adrenergic agonist actions and a D-isomer that has \( \beta_1 \)- and \( \beta_2 \)-adrenergic agonist actions. The net clinical effect is typically that of a potent \( \beta_1 \)-agonist with mild vasodilatory properties. At low dosages, it increases myocardial contractility without markedly increasing heart rate; this specificity is dose dependent and is lost at high dosages. Unlike dopamine, dobutamine does not release stored catecholamines and has no effect on dopaminergic receptors.\(^{313,314}\)

Administration and Adult Dosage. IV for inotropic support, by infusion only (in any nonalkaline IV fluid) 2.5 \( \mu \)g/kg/min initially, increasing gradually in 2.5 \( \mu \)g/kg/min increments to 20 \( \mu \)g/kg/min, and adjusting dosage to desired response. Maintenance dosages are typically 2–10 \( \mu \)g/kg/min.\(^{315}\) Although dosages up to 40 \( \mu \)g/kg/min have been used, use dosages above 20 \( \mu \)g/kg/min with caution because of increased risks of tachycardia, arrhythmias, and myocardial ischemia.\(^{316}\)

Special Populations. Pediatric Dosage. Safety and efficacy not established. However, IV infusion 2 \( \mu \)g/kg/min initially, followed by adjustment to desired hemodynamic response, up to 20 \( \mu \)g/kg/min, has been used.\(^{317}\)

Geriatric Dosage. Same as adult dosage.

Dosage Forms. Inj 12.5 mg/mL.

Pharmacokinetics. Onset and Duration. Onset <2 min; peak within 10 min; duration <10 min.\(^{318}\)

Fate. Wide interpatient variability exists, especially between adult and pediatric patients. \( V_d \) in CHF is 0.2 ± 0.08 L/kg; Cl in CHF is 3.5 ± 1.3 L/hr/kg. The drug is eliminated primarily in the liver to inactive glucuronide conjugates and 3-O-methyldobutamine.\(^{10,313}\) \( t_{1/2} \). 2.4 ± 0.7 min.\(^{10}\)

Adverse Reactions. Precipitation or exacerbation of ventricular ectopy occurs frequently; ventricular arrhythmias can occur (although these are less likely than with other sympathomimetics).\(^{313}\) Modest increases in heart rate or systolic blood pressure occur frequently; dosage reduction usually reverses these effects rapidly. Occasionally, nausea, headache, angina, nonspecific chest pain, palpitations, and shortness of breath are noted. Patients with atrial fibrillation might be at risk of developing rapid ventricular responses because dobutamine facilitates AV conduction.

Contraindications. Idiopathic hypertrophic subaortic stenosis.

Precautions. Correct hypovolemia before using in patients who are hypotensive. Although most cases of extravasation cause no signs of tissue damage, at least one case of dermal necrosis after extravasation of a 2.5 \( \mu \)g/kg/min infusion has been reported. Dobutamine contains a sulfite preservative that can be problematic in sensitive individuals, especially asthmatics.
**Drug Interactions.** Bretylium, guanethidine, and heterocyclic antidepressants can potentiate the pressor response to direct-acting vasopressors. Oxytocics used in obstetrics can cause severe, persistent hypertension when used with vasopressors. Halogenated hydrocarbon anesthetics can predispose patients to serious arrhythmias.

**Parameters to Monitor.** Monitor heart rate, arterial blood pressure, urine output, pulmonary capillary wedge pressure, cardiac index, ECG for ectopic activity, and infusion rate of solution continuously in the acute care setting and during periods of dosage titration or adjustment.

**Notes.** The drug is physically incompatible with sodium bicarbonate, furosemide, and other alkaline solutions. Use the reconstituted solution within 24 hr. (See Sympathomimetic Drugs for Hemodynamic Support Comparison Chart.)

---

**DOBUTAMINE DILUTION GUIDE**

<table>
<thead>
<tr>
<th>AMOUNT ADDED</th>
<th>VOLUME OF DILUENT</th>
<th>FINAL CONCENTRATIONa</th>
</tr>
</thead>
<tbody>
<tr>
<td>mg</td>
<td>Volume</td>
<td></td>
</tr>
<tr>
<td>250</td>
<td>1 vial (20 mL)</td>
<td>1000 mL 250 mg/L</td>
</tr>
<tr>
<td>250</td>
<td>1 vial (20 mL)</td>
<td>500 mL 500 mg/L</td>
</tr>
<tr>
<td>250</td>
<td>1 vial (20 mL)</td>
<td>250 mL 1 g/L</td>
</tr>
</tbody>
</table>

*aRecommended concentrations, but concentrations up to 5 g/L have been used.

---

**DOPAMINE HYDROCHLORIDE**

**Pharmacology.** Dopamine is a catecholamine that acts directly, in a dose-dependent fashion, on postsynaptic dopaminergic (DA1) receptors to produce renal and mesenteric vasodilation and on postsynaptic α1-, α2-, and β1-adrenergic receptors. It also acts indirectly by releasing norepinephrine from sympathetic nerve storage sites. Clinical response depends on the patient’s clinical condition and baseline sympathetic nervous system activity.319 Approximate ranges: dopaminergic 0.5–2 μg/kg/min; β1 5–10 μg/kg/min; mixed α and β 10–20 μg/kg/min; predominantly α >20 μg/kg/min.313

**Administration and Adult Dosage.** **IV for shock,** by infusion only (in any nonalkaline IV fluid) 2.5 μg/kg/min initially, increasing gradually in 5–10 μg/kg/min increments up to 20–50 μg/kg/min, and adjusting dosage to desired response. If a dosage over 20 μg/kg/min is required, consider other pressors.316 Use dosages over 50 μg/kg/min only with careful monitoring of hemodynamic parameters and urine output. **IV for chronic refractory CHF** 2–3 μg/kg/min initially and then increasing gradually until desired increases in urine flow, diastolic blood pressure, or heart rate are observed.320 Dosages over 20 μg/kg/min are rarely used in CHF.315,319,320 (See Notes.)

**Special Populations.** **Pediatric Dosage.** Safety and efficacy not established. However, **IV for shock** (recommendations for pediatric advance life support by the
American College of Cardiology and the American Heart Association)
2–5 μg/kg/min initially, increasing to 10–20 μg/kg/min to improve blood pressure, perfusion, and urine output.317

Geriatric Dosage. Same as adult dosage.

Dosage Forms. Inj 40, 80, 160 mg/mL. Also available prediluted, 0.8, 1.6, 3.2 mg/mL.

Pharmacokinetics. Onset and Duration. Onset within 5 min, duration <10 min.

Fate. There is large interpatient variability.313 One evaluation in adult surgery patients derived a $V_d$ of 0.89 ± 0.25 L/kg.321 $Cl$ is usually 3–4.2 L/hr/kg and 4.48 ± 0.94 L/hr/kg in adult surgery patients.313,316 The drug is metabolized primarily to homovanillic acid (HVA) and related metabolites; the remainder is metabolized to norepinephrine and excreted in urine as HVA and metabolites of both HVA and norepinephrine; very little is excreted as unchanged dopamine.

$t_{\frac{1}{2}}$. α phase 1–2 min, β phase 6–9 min.313,318

Adverse Reactions. Increases in ventricular ectopy and ventricular arrhythmias can occur, particularly at high dosages (although this is less likely than with other sympathomimetics); reduce dosage if the number of ventricular ectopic beats increases. Hypertension can occur at high infusion rates. Nausea, vomiting, headache, anxiety, and angina pectoris also have been observed. Gangrene of the extremities has occurred in patients given large dosages of dopamine for long periods and in patients with occlusive vascular disease given low dosages.313,322

Contraindications. Pheochromocytoma; presence of uncorrected tachyarrhythmias or ventricular fibrillation.

Precautions. Correct hypovolemia before using in patients with shock. If increased diastolic pressure, decreased pulse pressure, or decreased urine flow occurs, decrease infusion rate and monitor patient for signs of excessive vasoconstriction. Use with caution in patients with occlusive vascular disease and extreme caution in patients receiving halogenated hydrocarbon anesthesia. Avoid extravasation of solution; however, if it occurs, the area can be infiltrated with 5–10 mg of phentolamine diluted in 10–15 mL of NS. Dopamine contains a sulfite preservative that can be problematic in sensitive individuals, especially asthmatics.

Drug Interactions. MAOIs (including furazolidone and linezolid) can increase the pressor response to dopamine by up to 20-fold; avoid these combinations. Bretylium, guanethidine, and heterocyclic antidepressants can potentiate the pressor response to direct-acting vasopressors. Oxytocics used in obstetrics can cause severe, persistent hypertension when used with vasopressors. IV phenytoin can produce hypotension in severely ill patients receiving IV dopamine. Halogenated hydrocarbon anesthetics can predispose patients to serious arrhythmias.

Parameters to Monitor. In shock, closely monitor heart rate, ECG, pulmonary capillary wedge pressure, cardiac index, arterial blood pressure, arterial blood gases, acid–base balance, toe temperature, urine output, and infusion rate of solution and watch for signs of vasoconstriction or extravasation (eg, blanching). With low dosages for dopaminergic effects, monitor urine output and ECG.
Notes. The drug is physically incompatible with sodium bicarbonate, furosemide, and other alkaline solutions. (See Sympathomimetic Drugs for Hemodynamic Support Comparison Chart.) Low-dose dopamine is ineffective in preventing renal failure in critically ill patients with signs of early renal dysfunction.323

### INOTROPIC DRUGS

#### DOPAMINE DILUTION GUIDE

<table>
<thead>
<tr>
<th>AMOUNT ADDED</th>
<th>Volume</th>
<th>VOLUME OF DILUENT</th>
<th>FINAL CONCENTRATIONa</th>
</tr>
</thead>
<tbody>
<tr>
<td>200 mg</td>
<td>5 mL (1 amp, 40 mg/mL)</td>
<td>250 mL</td>
<td>800 mg/L</td>
</tr>
<tr>
<td>200 mg</td>
<td>5 mL (1 amp, 40 mg/mL)</td>
<td>500 mL</td>
<td>400 mg/L</td>
</tr>
<tr>
<td>400 mg</td>
<td>5 mL (1 amp, 80 mg/mL)</td>
<td>500 mL</td>
<td>800 mg/L</td>
</tr>
<tr>
<td>800 mg</td>
<td>5 mL (1 amp, 160 mg/mL)</td>
<td>500 mL</td>
<td>1.6 g/L</td>
</tr>
</tbody>
</table>

aRecommended concentrations, but concentrations up to 3.2 g/L have been used.

#### EPINEPHRINE AND SALTS

**Adrenalin, Sus-Phrine, Various**

**Pharmacology.** Epinephrine stimulates \( \alpha_1 \), \( \alpha_2 \) (vasoconstriction, pressor effects), \( \beta_1 \) (increased myocardial contractility and conduction), and \( \beta_2 \)-adrenergic (bronchodilation and vasodilation) receptors. It is used for reversible bronchospasm, anaphylactic reactions, laryngeal edema (croup), open-angle glaucoma, and cardiac arrest. (See Medical Emergencies.)

**Administration and Adult Dosage.** SC for anaphylaxis 0.2–0.5 mg (0.2–0.5 mL of 1:1000 aqueous soln), may repeat q 10–15 min prn; if SC is ineffective, then IV 0.1–0.25 mg (1–2.5 mL of 1:10,000) q 5–15 min may be given and followed by an IV infusion, if necessary. SC for asthma same dosage as SC for anaphylaxis; may repeat q 20 min to 4 hr as needed.144 SC aqueous suspension for asthma 0.5–1.5 mg (0.1–0.3 mL of 1:200), may repeat with 0.5–1.5 mg no sooner than q 6 hr. IV infusion for hemodynamic support 1 \( \mu \)g/min (1 mg in 500 mL NS or D5W) initially, adjust to hemodynamic response (usually 2–10 \( \mu \)g/min).316 Inhal (metered dose) not recommended because of low efficacy and ultrashort duration of action.

**Special Populations.** Pediatric Dosage. SC for anaphylaxis or asthma 0.01 mL/kg/dose of 1:1000 aqueous solution, to a maximum of 0.5 mL.; may repeat q 15–20 min for 2 doses, then q 4 hr prn. SC aqueous suspension for asthma (1 month–12 yr) 0.005 mL/kg/dose of 1:200, to a maximum of 0.15 mL/dose for children \( \leq 30 \) kg; repeat no sooner than q 6 hr.144 IV for croup 0.25–0.5 mL of 2.25% racemic aqueous solution diluted in 1.5–4.5 mL of NS q 1–2 hr prn by nebulizer.324,325 Alternatively, 5 mL of prediluted 1-epinephrine in NS (1:1000) has been used.325 IV for hemodynamic support 0.05–0.3 \( \mu \)g/kg/min initially, adjusted to desired hemodynamic response; avoid dosages >0.3 \( \mu \)g/kg/min, if possible, because they are associated with marked vasoconstrictor effects.317,326
Geriatric Dosage. Same as adult dosage. (See Precautions.)

Dosage Forms. Inhal Pwdr 200 µg/spray; Inhal Pwdr (bitartrate) 160 µg/spray; Inhal Soln (HCl) 1% (1:100); Inhal Soln (racpinephrine) 2.25%; Inj (aqueous solution as HCl) 0.01 mg/mL (1:100,000), 0.1 mg/mL (1:10,000), 0.5 mg/mL (1:2000), 1 mg/mL (1:1000); Inj (aqueous suspension as free base) 5 mg/mL (1:200).

Patient Instructions. (Autoinjectors) Periodically familiarize yourself with instructions for use so you maintain an adequate comfort level. Obtain new kit by expiration date or sooner if precipitate or color change is noted in solution.

Pharmacokinetics. Onset and Duration. Onset SC (aqueous soln or susp) 3–10 min; inhal peak 3–5 min. Duration SC (aqueous soln) 0.5–2 hr, (aqueous susp) up to 6–10 hr; inhal 15–60 min.323,327

Fate. Parenteral action is terminated by uptake into adrenergic neurons. Metabolism is by MAO and COMT.313,323 Cl is 2.1–5.3 L/hr/kg.313

Adverse Reactions. Dose-related restlessness, anxiety, tremor, cardiac arrhythmias, palpitations, hypertension, weakness, dizziness, and headache occur. Cerebral hemorrhage can be caused by a sharp rise in blood pressure from overdose. Angina can be precipitated when coronary insufficiency is present, and elevation of blood glucose has been reported. Local necrosis from repeated injections and tolerance with prolonged use also can occur.313,323

Contraindications. Intra-arterial administration is not recommended because of marked vasoconstriction. Do not use with local anesthetics in fingers or toes or during general anesthesia with halogenated hydrocarbons. Other contraindications include α-adrenergic blocker-induced (including phenothiazines) hypotension; cerebral arteriosclerosis; organic heart disease; narrow-angle glaucoma; shock; labor.

Precautions. Use with caution in patients with cardiovascular disease, hypertension, diabetes, or hyperthyroidism and in psychoneurotic patients. Caution is usually recommended in the elderly because of a higher frequency of cardiovascular intolerance. However, one evaluation of patients with acute asthma attacks found no difference in hemodynamic alterations or arrhythmias between those older and younger than 40 yr in response to SC epinephrine.328 IM injection can produce local tissue necrosis. Epinephrine infusions are administered preferably through a central venous line. Extravasation can cause necrosis; if extravasation occurs, infiltrate the area with phentolamine 5–10 mg diluted in 10–15 mL of NS.

Drug Interactions. Bretylium, guanethidine, and heterocyclic antidepressants can potentiate the pressor response to epinephrine. Oxytocics used in obstetrics can cause severe, persistent hypertension when used with vasopressors. Halogenated hydrocarbon anesthetics can predispose patients to serious arrhythmias. A hypertensive reaction can occur when epinephrine is given with nonselective β-adrenergic blockers (eg, propranolol, nadolol).
Parameters to Monitor. (IV infusion) ECG, infusion rate and site; (in the elderly) ECG; (asthma or allergy) blood pressure, heart rate, relief of symptoms.

Notes. Do not use solution if it is brown or contains a precipitate. Protect solution from light. The solution is incompatible with sodium bicarbonate, furosemide, and other alkaline solutions. Suspension provides a sustained effect; shake suspension well before use. Nonprescription inhalers have only a transient effect because of their low dosage and should be used only by patients who have infrequent symptoms (less than once a week) and obtain total relief of symptoms from administration of two inhalations. Parenteral administration offers no advantage over inhalation for the treatment of acute bronchospasm. (See Sympathomimetic Drugs for Hemodynamic Support Comparison Chart.)

INAMRINONE LACTATE

Pharmacology. Inamrinone (formerly amrinone) increases cyclic AMP and calcium availability through the inhibition of phosphodiesterase III, which improves cardiac output through vasodilatory and positive inotropic actions.

Administration and Adult Dosage. IV loading dose 0.5–1 mg/kg (usually 0.75 mg/kg) over 2–3 min; can repeat in 30 min based on response. IV maintenance dosage by continuous infusion 5–10 μg/kg/min.

Special Populations. Pediatric Dosage. Safety and efficacy not established, but the following dosages have been suggested: IV loading dosage (neonates and infants) 3–4.5 mg/kg, then IV maintenance dosage by continuous infusion (neonates) 3–5 μg/kg/min; (infants) 10 μg/kg/min.

Geriatric Dosage. Same as adult dosage.

Dosage Forms. Inj 5 mg/mL.

Pharmacokinetics. Onset and Duration. IV onset 2–5 min after bolus, peak 10 min, duration 60–90 min.

Serum Levels. A level of 1.5–4 mg/L (8–21.3 μmol/L) is associated with therapeutic response. Although a correlation exists between inamrinone levels and increased cardiac output, the clinical usefulness of serum level monitoring is not established.

Fate. Bioavailability is 93 ± 12%. The drug is 20–50% bound to plasma proteins; Vd is 1.8 ± 0.9 L/kg; Cl is 0.28 ± 0.1 L/hr/kg. The drug is eliminated primarily by hepatic metabolism, with 30 ± 20% excreted unchanged in urine. t½α phase 1–5 min; β phase 4.3 ± 1.3 hr in normals, 7.3 ± 4.6 hr in CHF.

Adverse Reactions. Dose-dependent, asymptomatic thrombocytopenia occurs frequently. Platelet counts return to normal within 2–4 days of discontinuing therapy; in some cases, this side effect is reversible when dosage is maintained or reduced. Thrombocytopenia might be caused primarily by the metabolite N-acetylinamrinone. Nausea and vomiting are unusual with IV use. Occasional side effects include nephrogenic diabetes insipidus, liver enzyme elevation, fever, taste disturbances, flu-like syndrome, rash, and aggravation of underlying arrhythmias.
Precautions. Caution in hypertrophic obstructive cardiomyopathy. One study found decreased survival rates in patients on long-term inamrinone therapy. Use concomitant antiplatelet drugs with caution.

Drug Interactions. None known.

Parameters to Monitor. Continuous ECG and frequent vital signs. Invasive hemodynamic monitoring is necessary in seriously ill patients for adequate dosage titration.

Notes. Do not dilute with dextrose-containing solutions but can infuse through dextrose-containing IV lines. Do not administer furosemide through IV lines containing inamrinone.

MILRINONE LACTATE

Pharmacology. Milrinone is a phosphodiesterase inhibitor, positive inotropic agent, and a vasodilator similar to inamrinone, but 10–15 times more potent on a weight basis. Milrinone is labeled for temporary use in patients with severe left ventricular dysfunction and CHF. It is also used for postoperative hemodynamic support and those awaiting cardiac transplantation.

Adult Dosage. IV loading dose 50 μg/kg over 10 min and then IV continuous infusion 0.375–0.5 μg/kg/min; adjust dosage depending on the patient’s response and hemodynamic variables to a maximum of 0.75 μg/kg/min. In renal impairment, reduce dosage as follows: Cl, 50 mL/min, 0.43 μg/kg/min; 40 mL/min, 0.38 μg/kg/min; 30 mL/min, 0.33 μg/kg/min; 20 mL/min, 0.28 μg/kg/min; 10 mL/min, 0.23 μg/kg/min; 5 mL/min, 0.2 μg/kg/min.

Pediatric Dosage. Safety and efficacy not established, but weight-based dosages similar to adult dosages have been used.

Dosage Forms. Inj 200 μg/mL (100 mL), 1 mg/mL.

Pharmacokinetics. Although well absorbed orally, milrinone is available only for IV use; 70% bound to plasma proteins; Vd is 0.47 ± 0.3 L/kg. It is 88–90% eliminated as unchanged drug by renal excretion. Elimination half-life in patients with CHF is 2.3 ± 0.1 hr, longer than in normal volunteers.

Adverse Reactions. Thrombocytopenia is less frequent (0.4%) with milrinone than with inamrinone. Milrinone can cause or worsen existing ventricular and supraventricular arrhythmias. Hypotension and headaches occur frequently.

Precautions. Long-term milrinone therapy can lead to increased mortality in patients with CHF. Use with caution in patients with severe obstructive aortic or pulmonary disease (eg, hypertrophic subaortic stenosis).

Parameters to Monitor. Continuous ECG and frequent vital signs. Invasive hemodynamic monitoring is necessary in seriously ill patients for adequate dosage titration.

Notes. Physically incompatible with IV furosemide.
Pharmacology. Norepinephrine is a catecholamine that directly stimulates $\beta_1$-, $\alpha_1$-, and $\alpha_2$-adrenergic receptors. It has little action on $\beta_2$-receptors.

Administration and Adult Dosage. IV for shock, by infusion only (in any nonalkaline IV fluid) 8–12 $\mu$g of base/min initially; adjust rate to maintain a systolic blood pressure of about 80–100 mm Hg or to a specific hemodynamic response; average maintenance dosage range is 2–4 $\mu$g of base/min. Very large dosages (up to 1.5 $\mu$g/kg/min) have been used in patients with septic shock.

Special Populations. Pediatric Dosage. Safety and efficacy not established. IV for shock, by infusion only 0.05–0.1 $\mu$g/kg/min of base initially, adjust dosage to blood pressure response, to a maximum of 1.5 $\mu$g/kg/min.

Geriatric Dosage. Same as adult dosage. (See Precautions.)

Dosage Forms. Inj 1 mg (of base)/mL.

Pharmacokinetics. Onset and Duration. Onset 1–2 min; duration 1–2 min after discontinuing infusion.

Fate. Action is terminated primarily by uptake into adrenergic neurons. Free drug is metabolized primarily by COMT and, to a lesser extent, MAO to inactive metabolites and their conjugates. $t_{1/2}$: 2–2.5 min.

Adverse Reactions. Dose-related hypertension (sometimes indicated by headache), reflex bradycardia, increased peripheral vascular resistance, and decreased cardiac output occur. Volume depletion can occur if fluid is not replaced. Arrhythmias can occur in extreme hypoxia or hypercarbia.

Contraindications. Hypotension secondary to uncorrected blood volume deficit; severe visceral or peripheral vasoconstriction; mesenteric or peripheral vascular thrombosis, unless drug is life-saving; halogenated hydrocarbon anesthesia.

Precautions. Use with caution in patients receiving MAOIs or heterocyclic antidepressants. Administer into a large vein (antecubital preferred) to avoid necrosis secondary to vasoconstriction; avoid the leg veins whenever possible, especially in the elderly or in those with occlusive vascular diseases. Avoid extravasation of solution; however, if it occurs, the area can be infiltrated with 5–10 mg of phentolamine diluted in 10–15 mL of NS.

Drug Interactions. Bretylium, guanethidine, MAOIs, methyldopa, and heterocyclic antidepressants can potentiate the pressor response to direct-acting vasoressors. Oxytocics used in obstetrics can cause severe, persistent hypertension when used with vasopressors. Halogenated hydrocarbon anesthetics can predispose patients to serious arrhythmias.

Parameters to Monitor. In shock, closely monitor heart rate, pulmonary capillary wedge pressure, cardiac index, arterial blood pressure, arterial blood gases, acid–base balance, urine output, and infusion rate of solution and watch for signs of vasoconstriction or extravasation (eg, blanching).
**Notes.** Do not use solution if it is brown or contains precipitate; 2 mg of norepinephrine bitartrate = 1 mg norepinephrine base. (See Sympathomimetic Drugs for Hemodynamic Support Comparison Chart.)

<table>
<thead>
<tr>
<th>AMOUNT ADDED</th>
<th>VOLUME OF 5% DEXTROSE</th>
<th>FINAL CONCENTRATION</th>
</tr>
</thead>
<tbody>
<tr>
<td>2 mg</td>
<td>2 mL</td>
<td>500 mL</td>
</tr>
<tr>
<td>4 mg</td>
<td>4 mL</td>
<td>500 mL</td>
</tr>
<tr>
<td>8 mg</td>
<td>8 mL</td>
<td>500 mL</td>
</tr>
</tbody>
</table>

<sup>a</sup>Recommended pediatric concentration<sup>340</sup>
# Sympathomimetic Drugs for Hemodynamic Support Comparison Chart

## Adrenergic Receptor Selectivity

<table>
<thead>
<tr>
<th>DRUG</th>
<th>Inotropic Activity ((\beta_1))</th>
<th>Chronotropic Activity ((\beta_2))</th>
<th>Vasodilation ((\alpha_1))</th>
<th>Vasoconstriction ((\alpha_2))</th>
<th>Renal/Mesenteric Vasodilation ((DA_1))</th>
<th>Total Peripheral Resistance</th>
<th>Cardiac Output</th>
</tr>
</thead>
<tbody>
<tr>
<td>Dobutamine</td>
<td>++</td>
<td>0/+/b</td>
<td>+</td>
<td>0/+/b</td>
<td>0</td>
<td>↓</td>
<td>↑</td>
</tr>
<tr>
<td>Dobutrex</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Dopamine</td>
<td>++</td>
<td>+/++c</td>
<td>++</td>
<td>+/++</td>
<td>+++</td>
<td>↓/↑h</td>
<td>↑</td>
</tr>
<tr>
<td>Inotropin</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Epinephrine</td>
<td>+++</td>
<td>+++</td>
<td>++</td>
<td>++++</td>
<td>0</td>
<td>↓</td>
<td>↑</td>
</tr>
<tr>
<td>Adrenalin</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Norepinephrine&lt;sup&gt;a&lt;/sup&gt;</td>
<td>++</td>
<td>++&lt;sup&gt;e&lt;/sup&gt;</td>
<td>0</td>
<td>++++</td>
<td>0</td>
<td>↑</td>
<td>0/↓/↑</td>
</tr>
<tr>
<td>Norepinephrine&lt;sup&gt;d&lt;/sup&gt;</td>
<td>++</td>
<td>++&lt;sup&gt;e&lt;/sup&gt;</td>
<td>0</td>
<td>++++</td>
<td>0</td>
<td>↑</td>
<td>0/↓/↑</td>
</tr>
<tr>
<td>Levophed</td>
<td>0</td>
<td>g</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>↓</td>
<td>↑</td>
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(continued)
### ADRENERGIC RECEPTOR SELECTIVITY

<table>
<thead>
<tr>
<th>DRUG</th>
<th>Inotropic Activity $(\beta_1)$</th>
<th>Chronotropic Activity $(\beta_2)$</th>
<th>Vasodilation $(\beta_2)$</th>
<th>Vasoconstriction $(\alpha_1)$</th>
<th>Renal/Total Resistance</th>
<th>Cardiac Output</th>
</tr>
</thead>
<tbody>
<tr>
<td>Phenylephrine</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>+++++</td>
<td>0</td>
<td>↑</td>
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<tr>
<td>Neo-Synephrine</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>↓</td>
</tr>
</tbody>
</table>

+++ +++ = Pronounced effect; + = Minimal effect; 0 = No effect; ↓ = Decreased; ↑ = Increased.

*This table compares only a few of the many factors important in the treatment of shock. Consult references 313, 315 and 341–343 for clinical use. Cross-table comparisons of the adrenergic selectivity properties between this table and the Sympathomimetic Bronchodilators Comparison Chart cannot be made because: (1) the rating scale of this table reflects a finer degree of differentiation of effects (hence 0–5+ vs 0–4+); (2) the routes of administration are different; and (3) vascular $\beta_2$-receptors appear to respond slightly differently from bronchiolar $\beta_2$-receptors.

bDose dependent.

cReleases stored norepinephrine via tyramine-like mechanism.
dUsed primarily to increase peripheral vascular resistance in volume-repleted hypotensive patients.
a$\alpha$-Adrenergic blocking drug; useful in severe vasoconstriction (e.g., extravasation of norepinephrine or dopamine).
iIncrease in heart rate can result from reflex and direct mechanisms.
hPrimary use is to increase blood pressure to reflexly increase vagal tone in paroxysmal supraventricular tachycardias. Other pressors are preferred in most shock states because they also have positive inotropic activity. Phenylephrine has no inotropic activity and with its strong $\alpha_1$-agonist properties functions as a pure vasopressor (afterload increaser).
Nitrates

**Class Instructions.** Nitrates. This drug can cause headache, dizziness, and/or flushing; alcohol can worsen these side effects. Tolerance to side effects of long-acting nitrates such as headache can occur with continued therapy. If necessary, a mild analgesic can be used until tolerance to side effects occurs. During an acute angina attack, discontinue activity, assume a sitting position, and dissolve one sublingual tablet under the tongue. If chest discomfort does not improve after use of the tablets, seek medical attention. Keep tablets in the tightly closed original container. If you have been taking this medication for a long time, do not discontinue it abruptly.

**Pharmacology.** (See Nitroglycerin.)

**Administration and Adult Dosage.**
- **SL tab for acute anginal attack** 2.5–10 mg q 2–3 hr prn;344,345
- **Chew Tab for acute anginal attack** 5 mg initially and then 5–10 mg q 2–3 hr prn.345
- **PO for prophylaxis of angina and for CHF** 10–60 mg q 4–6 hr; individual doses up to 120 mg have been used.344,346 (See Vasodilators in Heart Failure Comparison Chart.)
- **SR products for prophylaxis of angina** 40–80 mg q 8–12 hr (once daily–bid at 8 AM and 2 PM preferred). Start the dosage low and adjust upward slowly over several days to weeks to patient tolerance or to the desired therapeutic effect. A daily nitrate-free period of at least 12 hr is desirable to minimize tolerance.346

**Special Populations.**
- **Pediatric Dosage.** Safety and efficacy not established.
- **Geriatric Dosage.** Same as adult dosage. However, some clinicians have recommended lower doses and that initial SL doses be given under medical observation because of increased likelihood of postural hypotension.347

**Dosage Forms.**
- **Chew Tab** 5, 10 mg;
- **SL Tab** 2.5, 5, 10 mg;
- **SR Cap** 40 mg;
- **SR Tab** 40 mg;
- **Tab** 5, 10, 20, 30, 40 mg.

**Patient Instructions.** (See Nitrates Class Instructions.) Do not crush or chew sustained-release preparations.

**Missed Doses.** Take this drug at regular intervals. If you miss a dose, take it as soon as you remember. If it is about time for the next dose, take that dose only. Leave a minimum of 2 hours between regular tablet doses and 6 hours between sustained-release tablet or capsule doses. Do not double the dose or take extra.

**Pharmacokinetics.**
- **Onset and Duration.** Onset is 5–20 min after SL and Chew Tab administration, 15–45 min after PO tab administration, up to 4 hr in rare cases or with SR products; peak occurs 15–60 min after SL administration and 45–120 min after PO tab administration.344,345 Duration is 1–3 hr after SL or Chew Tab, 2–6 hr after PO tab, up to 8 hr after SR.344,348 Although PO administration can improve exercise tolerance for 4–8 hr after the first dose, with long-term, around-the-clock administration, the duration of action declines to 2–3 hr, probably because of nitrate tolerance.348

**Fate.** Oral bioavailability is 22 ± 14%. There is extensive first-pass metabolism by the liver after oral administration to less active isosorbide mononitrate metabolites (2-ISMN, 5-ISMN). Larger doses and long-term administration saturate metabolic
processes, with appreciable increases in serum concentrations of the parent compound and metabolites.\textsuperscript{349} The drug is $28 \pm 12\%$ bound to plasma proteins; $V_d$ is $1.5 \pm 0.8$ L/kg; $Cl$ is $2.7 \pm 1.2$ L/hr/kg.\textsuperscript{10} (See Isosorbide Mononitrate Fate.)

$\frac{t_\text{½}}{}$. (Isosorbide dinitrate) $50 \pm 20$ min; (2-ISMN) $1.9 \pm 0.5$ hr; (5-ISMN) $4.6 \pm 0.7$ hr.\textsuperscript{10,350}

**Adverse Reactions.** (See Nitroglycerin.)

**Contraindications.** (See Nitroglycerin.)

**Precautions.** (See Nitroglycerin.)

**Drug Interactions.** (See Nitroglycerin.)

**Parameters to Monitor.** Monitor for headache, orthostatic hypotension, and dizziness. In angina, monitor frequency of angina. In CHF, monitor hemodynamic and functional measurements.

**Notes.** Because of their slower onset of action, reserve SL and chewable isosorbide dinitrate (ISDN) for acute anginal attacks only in patients intolerant of or unresponsive to nitroglycerin. ISDN is a mainstay of antianginal and CHF therapy because of its long record of efficacy in these disorders. It is also less expensive than other long-term nitrate preparations. However, patients must take multiple doses daily on an eccentric schedule to achieve and maintain efficacy.

<table>
<thead>
<tr>
<th>ISOSORBIDE MONONITRATE</th>
<th>Imdur, ISMO, Monoket</th>
</tr>
</thead>
</table>

**Pharmacology.** Isosorbide mononitrate (ISMN) is the active 5-mononitrate metabolite of isosorbide dinitrate. (See Nitroglycerin.)

**Administration and Adult Dosage.** PO for prophylaxis of angina (ISMO, Monoket) 20 mg bid, doses 7 hr apart. SR for prophylaxis of angina (Imdur) 30–60 mg once daily in the morning, can increase to 120 mg once daily, to a maximum (rarely) of 240 mg once daily. There can be some attenuation of antianginal efficacy after 6 weeks of therapy with the 30 and 60 mg doses, but not with the 120 mg dose.\textsuperscript{348,351}

**Special Populations.** *Pediatric Dosage.* Safety and efficacy not established.

**Geriatric Dosage.** Same as adult dosage.

**Other Conditions.** For persons of particularly small stature, the manufacturer of Monoket recommends an alternative initial dosage of 5 mg bid, but to increase it to at least 10 mg bid by the second or third day of therapy.

**Dosage Forms.** SR Tab (Imdur) 30, 60, 120 mg; Tab (ISMO, Monoket) 10, 20 mg.

**Patient Instructions.** (See Nitrates Class Instructions.) Follow the prescribed administration schedule closely. Do not crush the sustained-release product.

**Missed Doses.** Take this drug at regular intervals. If you miss a dose, take it as soon as you remember. If it is about time for the next dose, take that dose only. Leave a minimum of 6 hours between regular tablet doses and 12 hours between sustained-release tablet doses. Do not double the dose or take extra.

**Pharmacokinetics.** *Onset and Duration.* Non-SR tablet onset 30–60 min,\textsuperscript{351} peak 1–2 hr, duration 12–14 hr with bid administration.\textsuperscript{348} SR Tab onset within 4 hr, peak 4 hr, duration about 12 hr.\textsuperscript{348,351}
**Fate.** The tablet is rapidly absorbed and essentially 100% bioavailable; SR Tab bioavailability is 78–86%. The drug is distributed into total body water with negligible plasma protein binding and a $V_d$ of $0.62 \pm 0.05$ L/kg. CI is $0.094 \pm 0.005$ L/hr/kg. It is primarily hepatically metabolized by denitration and glucuronidation to inactive products that are renally eliminated; less than 2% of a dose is excreted unchanged in urine.

$\frac{1}{2}$ $t_{\text{sp}}$ 4.6 ± 0.7 hr.

**Adverse Reactions.** (See Nitroglycerin.)

**Contraindications.** (See Nitroglycerin.)

**Precautions.** (See Nitroglycerin.)

**Drug Interactions.** (See Nitroglycerin.)

**Parameters to Monitor.** Monitor for headache, orthostatic hypotension, and dizziness; antianginal efficacy; and compliance with 7-hr dosage regimen for non-SR tablet.

**Notes.** ISMN is an effective nitrate with dosage schedules proven to avoid tolerance. Patient compliance is favored with the use of ISMN because the products are administered once (Indur) or twice (ISMO, Monoket) daily. Although the available preparations are comparable to each other in cost, at relatively low dosages (ie, 20 mg bid for the immediate-release products and 60 mg/day or less of the SR products) they are much more expensive than generic ISDN, which is also effective when taken properly. Compliance and cost factors must be assessed carefully in the individual patient when choosing an oral nitrate product.

**Nitroglycerin**

**Pharmacology.** Nitroglycerin and other organic nitrates are believed to be converted to nitric oxide (NO) by vascular endothelium. NO activates guanylate cyclase, increasing cyclic GMP that in turn decreases intracellular calcium, resulting in direct relaxation of vascular smooth muscle. The venous (capacitance) system is affected to a greater degree than the arterial (resistance) system. Venous pooling, decreased venous return to the heart (preload), and decreased arterial resistance (afterload) reduce intracardiac pressures and left ventricular size, thereby decreasing myocardial oxygen consumption and ischemia. In myocardial ischemia, nitrates dilate large epicardial vessels, enhance collateral size and flow, and reduce coronary vasoconstriction. The various organic nitrate preparations have the same pharmacologic effects and differ only in bioavailability and pharmacokinetics.

**Administration and Adult Dosage.** SL Tab for acute anginal attack 150–600 µg prn, up to 3 doses in 15 min; SL aerosol for acute anginal attack 400–800 µg prn, up to 1200 µg/15 min; Buccal for acute anginal attack and/or prophylaxis and treatment of angina pectoris or CHF 1–3 mg q 4–6 hr; PO SR for prophylaxis of angina or CHF 2.5–19.5 mg bid or tid; Top ointment for prophylaxis and treatment of angina pectoris or CHF 1.3–5 cm (0.5–2 inches) q 6–8 hr. Start the dosage low and adjust slowly upward over a several days to weeks to patient tolerance or to the desired therapeutic effect. SR Patch for prophylaxis and treatment of angina pectoris 0.2–0.8 mg/hr, patch applied once daily, dosage adjusted...
Greater antianginal efficacy has been noted with patches delivering at least 0.4 mg/hr. A daily nitrate-free period of 12 hr is desirable to minimize nitrate tolerance. IV for CHF, post-MI, angina pectoris, perioperative blood pressure control, or hypotensive anesthesia 5 μg/min initially by constant infusion using an infusion pump. Dosage must be adjusted to the individual patient’s response. Increase dosage initially in 5 μg/min increments q 3–5 min until response is noted. If no response occurs at 20 μg/min, increments of 10 μg/min and then perhaps 20 μg/min can be used. Once partial blood pressure response occurs, decrease incremental increases and increase intervals. (See Notes.)

Special Populations. Pediatric Dosage. Safety and efficacy not established. IV 0.5–20 μg/kg/min has been suggested.

Geriatric Dosage. Same as adult dosage but some clinicians have recommended lower doses and that initial SL doses be given under medical observation because of increased likelihood of postural hypotension.

Dosage Forms. Buccal SR Tab 1, 2, 3 mg; Oint 2%; SL Aerosol 400 μg/spray; SL Tab 300, 400, 600 μg; SR Cap 2.5, 6.5, 9, 13 mg; SR Tab 2.6, 6.5, 9 mg; SR Patch 0.1, 0.2, 0.3, 0.4, 0.6, 0.8 mg/hr; Inj 0.5, 5 mg/mL.

Patient Instructions. (See Nitrates Class Instructions.)

Pharmacokinetics. Onset and Duration. Onset immediate after IV, 2–5 min after SL or buccal, 20–45 min after SR Cap or Tab, 15–60 min after topical ointment, and 30–60 min after transdermal administration. Peak 4–8 min after SL, 4–10 min after buccal, 45–120 min after SR Cap or Tab, 30–120 min after topical ointment, and 1–3 hr after transdermal administration. Duration 10–30 min after IV and SL, 0.5–5 hr after buccal, 2–6 hr after oral, 4–8 hr after SR, and 3–8 hr after topical administration. Although the SR patch can act longer after the first application, long-term continuous therapy limits the duration of action to 4 hr or less, probably because of tolerance. With sustained intermittent therapy (removal of patch after 12 hr), duration is 8–12 hr.

Fate. Bioavailability of SL and Top are 38 ± 26% and 72 ± 20%, respectively. Extensive first-pass metabolism occurs after oral administration. The drug is 87 ± 1% bound to plasma proteins. Vd is 3.3 ± 1.2 L/kg; Cl is 13.8 ± 5.4 L/hr/kg. It is metabolized in the liver to less active dinitro and inactive mononitro metabolites. Larger doses and long-term administration can saturate metabolism and result in increased serum concentrations of drug and metabolites.

t½. β phase estimated to be 2.3 ± 0.6 min.

Adverse Reactions. Headache occurs very frequently; dizziness occurs frequently, especially with oral or topical administration. Occasionally, flushing, weakness, nausea, vomiting, palpitations, tachycardia, and postural hypotension occur. Many of these effects are dose related and can be minimized by slowly increasing the dosage. Tolerance and dependence can occur with prolonged use. Contact dermatitis occurs in up to 40% of patients using transdermal patches.

Contraindications. Severe anemia; severe hypotension or uncompensated hypovolemia; increased intracranial pressure; purported hypersensitivity or idiosyncrasy to nitroglycerin, nitrates, or nitrites; use of sildenafil. (See Drug Interac-
Constrictive pericarditis, pericardial tamponade, and inadequate cerebral circulation are also considered contraindications by some clinicians.348

**Precautions.** Some tolerance and cross-tolerance with other nitrates can occur with long-term or excessive use.348 Use with caution in patients with severe renal or hepatic disease, those with low or normal pulmonary capillary wedge pressure, and those receiving drugs that lower blood pressure. With intermittent therapy, anginal episodes can increase during the nitrate-free interval.359

**Drug Interactions.** Nitrates can produce additive vasodilation and severe postural hypotension when combined with alcohol or hypotensive drugs. Use in patients taking sildenafil can result in profound hypotension with serious consequences, including death.

**Parameters to Monitor.** Observe for headache, dizziness, and other side effects. Monitor for orthostatic hypotension, especially with first SL dose in elderly. (Angina) monitor frequency of angina. (CHF) obtain hemodynamic and functional measurements. (IV use) monitor blood pressure and heart rate constantly in all patients; monitoring pulmonary capillary wedge pressure in some patients also can be useful.

**Notes.** Large and unpredictable amounts of nitroglycerin are lost through polyvinylchloride (PVC) containers, most IV administration sets and tubing, and certain IV filters.360,361 The manufacturers recommend that IV nitroglycerin infusions be prepared and stored in glass bottles and infused through special non-PVC tubing to avoid the use of in-line filters. However, because nitroglycerin infusion rate is usually adjusted to response rather than by a microgram/kilogram dosage, the need for special tubing has been questioned.344 Some institutions have discontinued use of nitroglycerin tubing to reduce costs and achieved good results clinically when using PVC tubing to infuse nitroglycerin.362,363 However, because substantial amounts of nitroglycerin are adsorbed onto PVC tubing, special attention to patient response is advisable at the time of IV tubing changes. Stored in glass containers, the diluted injection is stable for 48 hr at room temperature and 7 days under refrigeration. When administration sets with large dead spaces are used, flush the line whenever the concentration of solution is changed. (See Vasodilators in Heart Failure Comparison Chart.)

### APPROXIMATE EQUIVALENT DOSAGES OF NITRATES

<table>
<thead>
<tr>
<th>PRODUCT</th>
<th>LOW DOSAGE</th>
<th>HIGH DOSAGE</th>
</tr>
</thead>
<tbody>
<tr>
<td>Nitroglycerin Ointment</td>
<td>≤1 inch q 6 hr</td>
<td>1–2 inches q 6 hr</td>
</tr>
<tr>
<td>Nitroglycerin Patch</td>
<td>0.4 mg/hr</td>
<td>0.4–0.6 mg/hr</td>
</tr>
<tr>
<td>Isosorbide Dinitrate</td>
<td>20 mg tid</td>
<td>20–40 mg tid</td>
</tr>
<tr>
<td>Isosorbide Mononitrate</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Immediate-Release</td>
<td>10–20 mg bid</td>
<td>20 mg bid</td>
</tr>
<tr>
<td>Sustained-Release</td>
<td>30–60 mg/day</td>
<td>60–120 mg/day</td>
</tr>
<tr>
<td>DRUG</td>
<td>DOSAGE(^a)</td>
<td>DURATION</td>
</tr>
<tr>
<td>------------</td>
<td>------------------------</td>
<td>----------</td>
</tr>
<tr>
<td><strong>ACE INHIBITORS</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td><em>Captopril</em></td>
<td>PO 25–100 mg tid.</td>
<td>hours</td>
</tr>
<tr>
<td>Capoten</td>
<td></td>
<td></td>
</tr>
<tr>
<td><em>Enalapril</em></td>
<td>PO 2.5–10 mg bid;</td>
<td></td>
</tr>
<tr>
<td>Vasotec</td>
<td>IV 0.625–5 mg q 6–12 hr.</td>
<td></td>
</tr>
<tr>
<td><em>Lisinopril</em></td>
<td>PO 5–20 mg/day.</td>
<td></td>
</tr>
<tr>
<td>Prinivil</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Zestril</td>
<td></td>
<td></td>
</tr>
<tr>
<td><em>Quinapril</em></td>
<td>PO 5–20 mg q 12 hr.</td>
<td></td>
</tr>
<tr>
<td>Accupril</td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>HYDRAZINE</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td><em>Hydralazine</em></td>
<td>PO 50–75 mg</td>
<td>hours</td>
</tr>
<tr>
<td>Apresoline</td>
<td>q 6–8 hr; usual</td>
<td></td>
</tr>
<tr>
<td>Various</td>
<td>maintenance 200–600 mg/day.</td>
<td></td>
</tr>
<tr>
<td><strong>NITRATES</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td><em>Isosorbide</em></td>
<td>PO 10–60 mg</td>
<td>hours</td>
</tr>
<tr>
<td>Dinitrates</td>
<td>q 4–6 hr.</td>
<td></td>
</tr>
<tr>
<td>Isordil</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Sorbitrate</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Various</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

(continued)
### VASODILATORS IN HEART FAILURE COMPARISON CHART (continued)

<table>
<thead>
<tr>
<th>DRUG</th>
<th>DOSAGE</th>
<th>DURATION</th>
<th>SITE OF ACTION</th>
<th>HR</th>
<th>MAP</th>
<th>PCWP</th>
<th>CI</th>
<th>SVR</th>
</tr>
</thead>
<tbody>
<tr>
<td>Nitroglycerin</td>
<td>(See monograph.)</td>
<td>minutes</td>
<td>V,(A)</td>
<td>sl↑/↓</td>
<td>↓</td>
<td>↓</td>
<td>↑/↓</td>
<td>sl↓</td>
</tr>
<tr>
<td>Various</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

**NITROPRUSSIDE**

<table>
<thead>
<tr>
<th>Nitroprusside Sodium</th>
<th>IV 0.1–3 µg/Kg/min.</th>
<th>minutes</th>
<th>A,V</th>
<th>0</th>
<th>sl↓</th>
<th>↓</th>
<th>↑</th>
<th>↓</th>
</tr>
</thead>
<tbody>
<tr>
<td>Various</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

A = arterial; V = venous; HR = heart rate; MAP = mean arterial pressure; PCWP = pulmonary capillary wedge pressure; CI = cardiac index; SVR = systemic vascular resistance; ↑ = increase; ↓ = decrease; sl = slight; 0 = no change

*Start with low dosages of these drugs and increase gradually with continuous hemodynamic monitoring. To avoid adverse rebound effects, carefully taper the dosages of these drugs if they are to be discontinued. (See Nitroglycerin Notes.)

*Predominant site of action. Parentheses denote lesser activity.

*From references 320, 364–368 and product information.*
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Anticonvulsants

Class Instructions. Anticonvulsants. It is important to take this medication as prescribed to control seizures; stopping it suddenly can increase seizures. This medication can cause drowsiness. Until the extent of this effect is known, use caution when driving, operating machinery, or performing other tasks requiring mental alertness. Avoid concurrent use of alcohol or other drugs that cause drowsiness. Report unusual or bothersome side effects. Always use an effective contraceptive; contact your physician if you plan to become or become pregnant.

Pharmacology. Carbamazepine is an iminostilbene compound related structurally to the tricyclic antidepressants. In animals, carbamazepine acts presynaptically to block firing of action potentials, which decreases the release of excitatory neurotransmitters, and postsynaptically by blocking high-frequency repetitive discharge initiated at cell bodies.

Administration and Adult Dosage. PO for epilepsy 100–200 mg bid with meals initially, increasing in increments of up to 200 mg/day at weekly intervals to effective dosage. Usual maintenance dosage is 10–30 mg/kg/day or 600–1600 mg/day in 2–4 divided doses; tid or qid administration is recommended when enzyme-inducing antiepileptic drugs are administered concurrently. SR product can be given bid. PO for trigeminal neuralgia 100 mg bid initially, increasing in 200 mg/day increments until relief of pain, to a maximum of 1.2 g/day. Usual maintenance dosage is 400–800 mg/day in 2–3 divided doses. PO loading dose 8 mg/kg in a single dose achieves therapeutic levels in 2 hr (with suspension) or 5 hr (with tablets) and is well tolerated. Rectal administration has been reported. (See Fate.)

Special Populations. Pediatric Dosage. PO for epilepsy (<6 yr) 10–20 mg/kg/day in 3–4 divided doses with the chewable tablets or in 4 divided doses with the suspension; (6–12 yr) 10–20 mg/kg/day with meals initially, increasing weekly in 100 mg/day increments as needed to achieve optimal clinical response. Usual maintenance dosage is 15–35 mg/kg/day or 400–800 mg/day in 3–4 divided doses. (>12 yr) same as adult dosage.

Geriatric Dosage. Clearance of carbamazepine is reduced in some elderly patients, so a lower maintenance dosage might be required.
Other Conditions. During pregnancy, increases in carbamazepine clearance can occur; dosage increases guided by serum levels and patient status might be necessary.2

Dosage Forms. Chew Tab 100 mg; Susp 20 mg/mL; Tab 200 mg; SR Cap 200, 300 mg (Carbatrol); SR Tab 100, 200, 400 mg (Tegretol XR).

Patient Instructions. (See Anticonvulsants Class Instructions.) Immediately report sore throat, fever, mouth ulcers, or easy bruising, which can be an early sign of a severe, but rare, blood disorder. The Tegretol XR shell might appear in the stool, but does not indicate a lack of absorption.

Missed Doses. Take this drug at regular intervals. If you miss a dose, take it as soon as you remember. If it is about time for the next dose, take that dose only. Do not double the dose or take extra. If you miss more than one dose in a day, call your physician.

Pharmacokinetics. Onset and Duration. Steady-state serum levels are attained within 2–4 days and subsequently can decline because of autoinduction of metabolism.2 (See Fate.)

Serum Levels. (Anticonvulsant) 4–12 mg/L (17–50 μmol/L). Variability exists in the relationship between serum levels and CNS side effects. (See Notes.)

Fate. Absorption from tablets is slow and erratic, with a bioavailability of 75–85%; peak serum levels occur 4–8 hr after a dose of immediate-release product and 19 ± 7 hr after a dose of SR product.2 Absorption is rapid with the suspension, with a peak serum level of 7.9 ± 1.9 mg/L at 1.6 ± 1.3 hr in the fasting state or 3.4 ± 3.4 hr with concomitant enteral tube feeding after a 500 mg oral dose.4 A peak serum level of 5.1 ± 1.6 mg/L occurs 6.3 ± 1.5 hr after rectal administration of 6 mg/kg oral suspension (100 mg/5 mL) diluted with an equal volume of water.5 The drug is 75–78% bound to plasma proteins.2,6 Vd is 0.88 ± 0.06 L/kg in adults4 and 1.2 ± 0.2 L/kg in children.7 Large differences in Cl occur because of autoinduction of liver enzymes; autoinduction is completed within 1–2 weeks of monotherapy; Cl is 0.052 ± 0.04 L/hr/kg at end of week 1, 0.04 ± 0.02 L/hr/kg at week 2, and 0.05 ± 0.04 L/hr/kg at week 4.8 Carbamazepine is metabolized to pharmacologically active carbamazepine-10,11-epoxide; the epoxide metabolite (CBZ-E) serum level ratios at steady state are 0.19 ± 0.06 at a carbamazepine level of 6.9 ± 1.5 mg/L and 0.28 ± 1.4 at a carbamazepine level of 10.5 ± 2.6 mg/L.9 (See Notes.) Only about 2% of drug is excreted unchanged in the urine.2,6

t1/2. There are large interindividual differences because of autoinduction of liver enzymes. (Adults) 31.3 ± 5.9 hr after a single dose,4 14.5 ± 5.3 hr after 2 months;9 (children) 29.4 ± 3.6 hr after a single dose, 15.2 ± 5.2 hr after 5 months.7

Adverse Reactions. Dizziness, drowsiness, headache, diplopia, nausea, and vomiting occur frequently with initiation of therapy and are minimized by slow titration of dosage. Mild, transient, morbilliform rash and thrombocytopenia also occur frequently. Occasionally, confusion, stomatitis, or rash occur. Hyponatremia and water intoxication occur, and risk factors include carbamazepine monotherapy, elevated serum levels, patient age >25 yr, diuretic use, vomiting, or diarrhea.2 Transient leukopenia has been observed in 10–20% of patients; persistent leukopenia occurs in 2% of patients.5 Discontinue drug if leukopenia (ANC <1500/µL) persists.
or any evidence of bone marrow depression develops. Rare effects include aplastic anemia, agranulocytosis, hepatitis, lenticular opacities, and arrhythmias.

**Contraindications.** History of bone marrow depression; hypersensitivity to tricyclic antidepressants.

**Precautions.** Pregnancy; history of liver disease. Abrupt withdrawal of the drug in patients with epilepsy can precipitate status epilepticus. Exacerbation of atypical absence seizures can occur in children receiving carbamazepine for mixed seizure disorders.² Use carbamazepine cautiously in patients with histories of severe hypersensitivity reactions to phenytoin or phenobarbital.

**Drug Interactions.** Because of structural similarities to tricyclic antidepressants, discontinue MAOIs for a minimum of 14 days before starting carbamazepine. Carbamazepine can stimulate the metabolism of many drugs metabolized by CYP3A4, including oral anticoagulants, oral contraceptives, corticosteroids, cyclosporine, doxycycline, haloperidol, heterocyclic antidepressants, protease inhibitors, and theophylline. Many drugs inhibit carbamazepine metabolism, including cimetidine, clarithromycin, danazol, erythromycin, fluoxetine, isoniazid, ketoconazole, propoxyphene, quinine, troleandomycin, verapamil, and diltiazem.

**Parameters to Monitor.** Baseline CBC and platelet counts; monitor more frequently if WBC or platelet counts decrease. Monitor liver function tests periodically during long-term therapy. Monitor serum levels at least weekly during the first month of therapy because of autoinduction. Periodic serum level monitoring is useful in evaluating therapeutic efficacy or potential for adverse effects.²

**Notes.** The contribution of CBZ-E to the therapeutic or adverse effects of carbamazepine is uncertain. One study found no significant correlation between toxicity score or seizure frequency and serum levels of carbamazepine, CBZ-E, or their sum in patients receiving carbamazepine monotherapy, or combination therapy with phenytoin or valproic acid.⁹ (See Anticonvulsants Comparison Chart.)

### CLONAZEPAM  Klonopin

**Pharmacology.** Clonazepam is a benzodiazepine anticonvulant that limits the spread of seizure activity, possibly by enhancing the postsynaptic effect of the inhibitory neurotransmitter, γ-aminobutyric acid (GABA).

**Administration and Adult Dosage.** PO for epilepsy no more than 0.5 mg tid initially; increase in 0.5–1 mg/day increments q 3 days to effective dosage or to a maximum of 20 mg/day. **Usual maintenance dosage** is 4–8 mg/day.¹ PO for panic disorder 0.25 mg bid initially, increasing to 1 mg/day after 3 days. **Usual maintenance dosage** is 1 mg/day; some patients require up to 4 mg/day. (See Notes.) Rectal administration has been reported. (See Fate.)

**Special Populations.** Pediatric Dosage. PO for epilepsy (≤10 yr or ≤30 kg) 0.01–0.03 mg/kg/day initially in 2–3 divided doses, increase in 0.25–0.5 mg/day increments q 3 days to effective dosage, to a maximum of 0.2 mg/kg/day in 3 divided doses. Rectal administration has been reported. (See Fate.)

**Geriatric Dosage.** PO for epilepsy and as an antipanic agent same as adult dosage initially. **Maintenance dosage** requirements might be lower in elderly pa-
patients because of reduced drug clearance and enhanced pharmacodynamic response.10

**Dosage Forms.** Tab 0.5, 1, 2 mg.

**Patient Instructions.** *(See Anticonvulsants Class Instructions.)*

**Missed Doses.** Take this drug at regular intervals. If you miss a dose, take it as soon as you remember. If it is about time for the next dose, take that dose only. Do not double the dose or take extra.

**Pharmacokinetics.**

**Onset and Duration.** Steady-state serum levels are attained in 4–8 days.2 *(See Notes.)*

**Serum Levels.** 13–72 μg/L (40–230 nmol/L); however, some patients controlled with clonazepam can have levels below this range. There is a poor correlation between serum levels and efficacy or adverse effects.2

**Fate.** Rapidly absorbed orally; peak serum levels occur 1–3 hr after a dose. Serum levels of 18–40 μg/L occur 20–120 min after rectal administration of a 0.1 mg/kg dose of clonazepam suspension.11 The drug is 86 ± 0.5% bound to plasma proteins; Vₐ is 3.1 ± 1.2 L/kg; Cl is 0.059 ± 0.011 L/hr/kg.12 The principal metabolite, 7-aminoclonazepam, is inactive. Less than 0.5% of clonazepam is excreted unchanged in urine.2

**t½.** 34.1 ± 7.5 hr.12

**Adverse Reactions.** Drowsiness, ataxia, behavior disturbances, and personality changes (hyperactivity, restlessness, and irritability, especially in children) occur frequently and require dosage reduction.2 Occasionally, hypersalivation and bronchial hypersecretion occur and can cause respiratory difficulties. Rarely, anemia, leukopenia, thrombocytopenia, and respiratory depression occur. Nearly 50% of patients receiving long-term clonazepam can experience transient exacerbations of seizures, dysphoria, restlessness, or autonomic signs during clonazepam withdrawal.13 An increased seizure frequency and status epilepticus occur rarely, possibly associated with supratherapeutic serum levels.2

**Contraindications.** Severe liver disease; acute narrow-angle glaucoma.

**Precautions.** Pregnancy; lactation; patients with chronic respiratory disease. Clonazepam can increase frequency of generalized tonic-clonic seizures in patients with mixed seizure types. Abrupt withdrawal of the drug in patients with epilepsy can precipitate status epilepticus. Absence status has been reported in patients receiving valproic acid concurrently.

**Drug Interactions.** Concurrent use with other CNS depressants can potentiate the sedation caused by clonazepam.

**Parameters to Monitor.** Periodic serum level monitoring is of limited value. Close attention to changes in patient’s seizure frequency is necessary to monitor for the development of tolerance to the therapeutic effect. *(See Notes.)*

**Notes.** Tolerance to the anticonvulsant effect of clonazepam occurs in approximately one-third of patients within 3–6 months of starting the drug. Taper and discontinue clonazepam if therapeutic benefit cannot be demonstrated. Because of
prominent CNS adverse effects and the development of tolerance, it is considered an alternative to valproic acid for myoclonic seizures and an alternative to ethosuximide or valproic acid for absence seizures. Clonazepam also has become an alternative to alprazolam for treatment of panic disorder. For patients who experience interdose symptom recurrence or morning rebound with alprazolam, clonazepam offers an equally effective alternative with the benefit of a longer duration of effect. When switching a patient from alprazolam to clonazepam, an equivalent dosage of clonazepam is one-half that of alprazolam.

**ETHOSUXIMIDE**

**Pharmacology.** Ethosuximide is a succinimide that produces an anticonvulsant effect by blockade of T-type calcium currents in the thalamus. In humans, it suppresses 3 cycle per second spike and wave activity. (See Notes.)

**Administration and Adult Dosage.** PO for epilepsy 250 mg bid initially; increase in 250 mg/day increments at 4- to 7-day intervals to an effective dosage, to a maximum of 1.5 g/day. Usual maintenance dosage 750–1250 mg/day in 1–2 doses.1,2

**Special Populations.** Pediatric Dosage. PO for epilepsy (3–6 yr) 250 mg/day initially; increase in 250 mg/day increments q 4–7 days to effective dosage, to a maximum of 1 g/day. Usual maintenance dosage 20–40 mg/kg/day in 1–2 doses.1,2 (>6 yr) same as adult dosage.

**Geriatric Doses.** Same as adult dosage.

**Dosage Forms.** Cap 250 mg; Syrup 50 mg/mL.

**Patient Instructions.** (See Anticonvulsants Class Instructions.) This drug can be taken with food or milk to minimize stomach upset.

**Missed Doses.** Take this drug at regular intervals. If you miss a dose, take it as soon as you remember. If it is about time for the next dose, take that dose only. Do not double the dose or take extra.

**Pharmacokinetics.** Onset and Duration. Steady-state serum levels are attained in 7–12 days.2

**Serum Levels.** 40–100 mg/L (280–710 μmol/L).2

**Fate.** The drug is well absorbed orally, with peak serum level in 3–7 hr in adults and children. Plasma protein binding is less than 10%. Vd is 0.69 L/kg;17 Cl is 0.01 ± 0.04 L/hr/kg, greater in children.2 Ethosuximide is metabolized to three inactive metabolites. About 20% of the drug is excreted unchanged in urine.2,6 t½. (Adults) 52.6 hr,18 (children) 31.6 ± 5.4 hr.17

**Adverse Reactions.** Nausea, vomiting, drowsiness, headache, hiccups, and dizziness occur frequently during initiation of therapy and usually are dose related. Occasionally, behavior changes or rashes occur. Rarely, SLE, leukopenia, aplastic anemia, or Stevens-Johnson syndrome occur.

**Precautions.** Pregnancy; patients with known liver or renal disease. Generalized tonic-clonic seizures can occur in patients with mixed seizure types who are
treated with ethosuximide alone. Abrupt withdrawal of the drug can precipitate absence status epilepticus.

**Contraindications.** None known.

**Drug Interactions.** Ethosuximide can increase phenytoin serum levels and decrease levels of primidone (and its phenobarbital metabolite).

**Parameters to Monitor.** Periodic serum level monitoring, after attaining steady state (7–12 days), is useful in evaluating therapeutic efficacy or potential adverse effects. Periodically monitor CBC, urinalysis, and liver function tests.

**Notes.** Ethosuximide is indicated only for treatment of absence seizures. Because of the drug’s low potential for serious or long-term toxicity and its proven efficacy, it is considered the drug of choice for absence seizures. (See Anticonvulsants Comparison Chart.)

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**FELBAMATE**  
Felbatol

**Pharmacology.** Felbamate is a dicarbamate that is structurally related to meprobamate; its mechanism of action is not known but might involve inhibition of \(\text{N}-\text{methyl-D-aspartate}\) responses and potentiation of \(\text{GABA}_A\) receptor chloride currents.\(^{19}\)

**Administration and Adult Dosage.** PO for epilepsy 1.2 g/day initially in 3–4 divided doses while reducing the dosage of concomitant antiepileptic drugs (phenytoin, carbamazepine, valproic acid, or phenobarbital) by 20–30%. Increase in 1200 mg/day increments at weekly intervals to a maximum of 3.6 g/day. Further reduction in the dosage of concomitant antiepileptic drugs might be required during felbamate titration.

**Special Populations.**  
**Pediatric Dosage.** PO for epilepsy 15 mg/kg/day initially in 3–4 divided doses while reducing the dosage of concomitant antiepileptic drugs (phenytoin, carbamazepine, valproic acid, or phenobarbital) by 20–30%. Increase in 15 mg/kg/day increments at weekly intervals to a maximum of 45 mg/kg/day. Further reduction in the dosage of concomitant antiepileptic drugs may be required during felbamate titration.

**Geriatric Dosage.** Dosage reduction might be required in patients with reduced hepatic or renal function and should be guided by clinical response.

**Dosage Forms.** Tab 400, 600 mg; Susp 120 mg/mL.

**Patient Instructions.** (See Anticonvulsants Class Instructions.) Felbamate has been associated with severe blood and liver disorders that can be fatal. Report signs of infection, bleeding, easy bruising, or signs of anemia (fatigue, weakness) immediately; also report abdominal pain or yellowing of the skin immediately. Give the patient the information/consent section of the Felbatol prescribing information and obtain informed consent at the time of initial prescribing.

**Missed Doses.** Take this drug at regular intervals. If you miss a dose, take it as soon as you remember. If it is about time for the next dose, take that dose only. Do not double the dose or take extra.

**Pharmacokinetics.** **Onset and Duration.** Steady-state serum levels are attained in 2–3 days.\(^2\)
**Serum Levels.** A therapeutic range has not been established. Serum concentrations reported in clinical studies are 20–137 mg/L.\textsuperscript{20,21}

**Fate.** Rapidly absorbed with over 90% bioavailability; peak serum levels occur 1–4 hr after an oral dose.\textsuperscript{22} Food and antacids have no appreciable effect on absorption.\textsuperscript{2} The drug is 22–25% bound to plasma proteins, primarily to albumin. V\textsubscript{d} is 0.76 ± 0.08 L/kg; Cl is 0.030 ± 0.008 L/hr/kg in adults. From 40% to 50% is excreted unchanged in urine; 5% is excreted unchanged in feces; the remainder is excreted as inactive metabolites.\textsuperscript{2}

\( t_{1/2} \) 20.2 hr in normal volunteers;\textsuperscript{22} 14.7 ± 2.8 hr in epileptic patients on concomitant enzyme-inducing antiepileptic drugs.\textsuperscript{23}

**Adverse Reactions.** Anorexia, vomiting, insomnia, nausea, headache, weight loss, dizziness, and somnolence occur frequently. Adverse reaction frequency is lower when felbamate is used as monotherapy, and reactions often resolve during long-term therapy. Adverse reactions during adjunctive therapy can be the result of drug interactions. Felbamate is occasionally associated with aplastic anemia and hepatic failure (see Precautions), with fatality rates of 20–30%. It is not known whether the risks of aplastic anemia and hepatic failure are related to the duration of felbamate exposure.\textsuperscript{24}

**Contraindications.** History of any blood dyscrasia or hepatic dysfunction.

**Precautions.** Do not use felbamate as a first-line antiepileptic drug. Because of the risk of aplastic anemia and hepatic failure, use felbamate only in patients whose seizures cannot be controlled with other antiepileptic drugs or whose epilepsy is so severe that the risks are deemed acceptable. Fully inform patients of the risks of felbamate therapy. (See Patient Instructions.)

**Drug Interactions.** Felbamate increases serum concentrations of phenytoin, CBZ-E (the active metabolite of carbamazepine), and valproate; therefore, reduce the dosage of these antiepileptics by 20–30% when felbamate is initiated.

**Parameters to Monitor.** Close monitoring of CBC, platelets, liver function tests, and clinical signs or symptoms of infection, bruising, bleeding, or hepatitis is essential. Monitor liver function tests (ie, AST, ALT, bilirubin) q 1–2 weeks while treatment continues. The acceptable frequency of hematologic monitoring is not established. Routine monitoring of serum levels is of limited value because of the lack of a well-defined therapeutic range.

**Notes.** Felbamate is effective for the treatment of partial and secondarily generalized seizures in adults and for partial and generalized seizures associated with the Lennox–Gastaut syndrome in children.

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**FOSPHENYTOIN**  
**Pharmacology.** Fosphenytoin is a phosphate ester prodrug that is rapidly and completely converted to phenytoin in vivo by phosphatases after parenteral administration. Fosphenytoin has no pharmacologic activity before its conversion to phenytoin. (See Phenytoin.)

**Administration and Adult Dosage.** IV loading dose 15–20 mg phenytoin equivalents (PE)/kg at a maximum rate of 150 mg PE/min. IM loading dose 10–20 mg
PE/kg in 1 or more injection sites. **IV or IM maintenance dosage** 4–6 mg PE/kg/day in 1 or 2 divided doses. Safety and effectiveness have not been established for therapy lasting more than 5 days. **IV or IM substitution for oral phenytoin therapy** use same total daily dosage in PEs.

**Special Populations. Pediatric Dosage.** Safety and efficacy not established.

**Geriatric Dosage.** Advanced age has no effect on fosphenytoin pharmacokinetics, but phenytoin clearance and protein binding might be reduced. (See Phenytoin.)

**Other Conditions.** In patients with renal or hepatic disease, fosphenytoin conversion to phenytoin might be increased because of protein binding changes. Because of an increased fraction of unbound phenytoin in patients with renal or hepatic diseases or hypoalbuminemia, dosage adjustments should be guided by patient status and measurement of unbound phenytoin concentrations.

**Dosage Forms. Inj** 50 mg PE/mL.

**Patient Instructions.** Itching or tingling can occur during intravenous infusion, particularly in the facial and groin areas. These sensations are usually mild and disappear within minutes of stopping the infusion. These symptoms do not indicate an allergic reaction to fosphenytoin or phenytoin.

**Pharmacokinetics. Onset and Duration.** Therapeutic concentrations of unbound phenytoin (≥1 μg/mL) are attained within 8–10 min after the start of an IV infusion of fosphenytoin (administered at a rate of 100–150 mg PE/min) and are similar to those attained after an equivalent dose of phenytoin administered at 50 mg/min. Therapeutic concentrations of unbound phenytoin are attained within 30 min after IM injection of fosphenytoin.

**Serum Levels.** Monitoring fosphenytoin serum concentration is not clinically useful. Phenytoin serum concentrations correlate with efficacy and toxicity. (See Phenytoin.)

**Fate.** Fosphenytoin is completely converted to phenytoin by phosphatases. Peak-free phenytoin concentrations occur 30 min after completion of fosphenytoin infusion administered at rates of 100–150 mg PE/min and occur 3 hr after an IM dose of fosphenytoin. (See Onset and Duration.) Fosphenytoin is 95–99% bound to plasma proteins, primarily albumin. Fosphenytoin binding is saturable and the Vₐ of fosphenytoin is 4.3–10.8 L depending on plasma concentration. Fosphenytoin displaces phenytoin from protein binding sites. The free (unbound) fraction of phenytoin ranges from 0.30 (in the presence of fosphenytoin) to 0.12 (after complete conversion of fosphenytoin to phenytoin).²⁶,²⁷ (See Phenytoin.)

\[ t_{1/2} \] (Conversion to phenytoin) 15 min.

**Adverse Reactions.** Fosphenytoin commonly causes burning, itching, or paresthesias during IV infusion, particularly in the groin and facial areas. These symptoms usually disappear within minutes and can be minimized by slowing or stopping the infusion. The frequency and severity of these symptoms increase with fosphenytoin dose and infusion rate and might be related to the phosphate load. Venous irritation including pain, erythema, swelling, tenderness, and cording (hardening of the vessel) occurs less often than with phenytoin.²⁸ IM fosphenytoin
injections are well tolerated and no significant differences in local symptoms were reported when compared with IM saline injections. No significant differences between fosphenytoin and phenytoin have been reported with regard to adverse cardiovascular effects with IV infusion. CNS adverse effects are common and likely represent reactions to phenytoin. (See Phenytoin.)

**Contraindications.** Sinus bradycardia; sinoatrial block; second- or third-degree AV block; Adams–Stokes syndrome.

**Precautions.** Consider the phosphate load of fosphenytoin (0.0037 mmol phosphate/mg PE fosphenytoin) in patients with renal insufficiency and those requiring phosphate restriction. (See Phenytoin.)

**Drug Interactions.** No drugs are known to affect the conversion of fosphenytoin to phenytoin. (See Phenytoin.)

**Parameters to Monitor.** Common immunoassays (eg, TDx, TDx/FLx) overestimate phenytoin concentrations when fosphenytoin is present. Determine serum phenytoin concentrations no earlier than 2 hr after an IV infusion or 4 hr after an IM dose of fosphenytoin. Obtain samples in tubes containing ethylenediaminetraacetic acid to minimize the ex vivo conversion of fosphenytoin to phenytoin. Monitor blood pressure and ECG during and 1–2 hr after IV fosphenytoin infusion.

**Notes.** Unlike parenteral phenytoin, fosphenytoin is not formulated with propylene glycol and is compatible with most common IV solutions (including those containing dextrose). Fosphenytoin, undiluted or admixed in NS or D5W, is stable for 30 days at room temperature.

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**GABAPENTIN** Neurontin

**Pharmacology.** Gabapentin is a cyclohexane compound that is structurally related to GABA; its mechanism of action is not known. Gabapentin does not interact with GABA receptors or alter the formation, release, degradation, or reuptake of GABA.

**Administration and Adult Dosage.** PO for epilepsy 300 mg hs on day 1; 300 mg bid on day 2; 300 mg tid on day 3. However, many patients tolerate initiation with 300 mg bid or tid. Dosage can be increased according to clinical response. **Usual maintenance dosage** 900–2400 mg/day in 3 divided doses. 1 Dosages of 3.6–4.8 g/day have been well tolerated in some patients. 31 Give dosages ≥3.6 g/day in 4 divided doses qid. (See Notes.)

**Special Populations.** **Pediatric Dosage.** (<12 yr) safety and efficacy not established.

**Geriatric Dosage.** Lower dosages might be required because of normal age-related decreases in renal function.

**Other Conditions.** Reduce dosage in patients with compromised renal function as indicated on the following page:
Dosage Forms. Cap 100, 300, 400 mg.

Patient Instructions. (See Anticonvulsants Class Instructions.) Do not take this drug with antacids.

Missed Doses. Take this drug at regular intervals. If you miss a dose, take it as soon as you remember. If it is about time for the next dose, take that dose only. Do not double the dose or take extra.

Pharmacokinetics. Onset and Duration. Steady-state serum levels are attained in 1–2 days in patients with normal renal function.32

Serum Levels. A therapeutic range has not been established. One study found that therapeutic effect correlated with serum concentrations greater than 2 mg/L.33

Fate. Rapidly absorbed and food has no effect on absorption. Absorption occurs via a saturable transport mechanism, so bioavailability decreases with dosages greater than 1.8 g/day; at this dosage it is 60%. At a dosage of 4.8 g/day, bioavailability is 35%.33 The drug is not bound to plasma proteins. Vd is 58 ± 6 L in adults; Cl is 0.17 ± 0.05 L/kg/hr in adults with normal renal function; Cl is linearly related to Clcr. Gabapentin is not appreciably metabolized but is eliminated unchanged in urine and normal renal function.34

\[ t_{1/2} = 4.8 \pm 1.4 \text{ hr in adults with epilepsy and normal renal function.} \]

Adverse Reactions. Somnolence, dizziness, ataxia, nystagmus, and headache occur frequently. Symptoms are of mild to moderate severity and resolve within 2 weeks with continued treatment.35 Weight gain (mean 4.9% of body weight) and peripheral edema also occur frequently.36 Occasionally, rash occurs. Rarely, behavioral changes in children occur.37

Contraindications. None known.

Precautions. Abrupt withdrawal of gabapentin in patients with epilepsy can precipitate status epilepticus. In vivo carcinogenicity studies have demonstrated a high incidence of pancreatic acinar cell tumors in male rats; the relevance of this observation to humans is not known.

Drug Interactions. Gabapentin does not induce or inhibit hepatic microsomal enzymes and does not affect the metabolism of other antiepileptic drugs or oral contraceptives. Antacids decrease the oral bioavailability of gabapentin by about 20%.

Parameters to Monitor. Serum level monitoring is of limited value because of the lack of a well-defined therapeutic range. Routine monitoring of clinical laboratory parameters during gabapentin therapy is not indicated.

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<table>
<thead>
<tr>
<th>CLcr (ML/Min)</th>
<th>Dosage Regimen</th>
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<tbody>
<tr>
<td>&gt;60</td>
<td>400 mg tid</td>
</tr>
<tr>
<td>30–60</td>
<td>300 mg bid</td>
</tr>
<tr>
<td>15–30</td>
<td>300 mg daily</td>
</tr>
<tr>
<td>&lt;15</td>
<td>300 mg every other day</td>
</tr>
<tr>
<td>Hemodialysis</td>
<td>200–300 mg after dialysis</td>
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</tbody>
</table>
Notes. Gabapentin is indicated as adjunctive treatment for partial and secondarily generalized seizures in adults. Preliminary studies have indicated that the drug can be efficacious as an adjunct in children with refractory partial seizures. Preliminary studies have suggested efficacy in other painful conditions (eg, reflex sympathetic dystrophy), bipolar disorder, and other psychiatric conditions.

Lamotrigine

Pharmacology. Lamotrigine is a phenyltriazine derivative unrelated to other marketed antiepileptic drugs. Lamotrigine inhibits voltage-dependent sodium channels, thereby stabilizing neuronal membranes and reducing the release of excitatory neurotransmitters such as glutamate and aspartate.

Administration and Adult Dosage. Adjust starting dosages, titration schedules, and maintenance dosage based on concomitant therapy. PO for epilepsy in patients receiving enzyme-inducing antiepileptic drugs (eg, carbamazepine, phenytoin, phenobarbital, and primidone) 50 mg/day for 2 weeks and then increase to 50 mg bid for 2 more weeks. Thereafter, increase dosage in 100 mg/day increments at weekly intervals to a maintenance dosage of 300–500 mg/day in 2 divided doses. PO for epilepsy in patients receiving enzyme-inducing antiepileptic drugs with valproic acid 25 mg every other day for 2 weeks and then increase to 25 mg/day for 2 more weeks. Thereafter, increase dosage in 25–50 mg/day increments at 1- to 2-week intervals to a maintenance dosage of 100–200 mg/day in 2 divided doses. Dosage recommendations are not available for patients receiving valproic acid alone, but dosages are expected to be lower due to prolonged half-life.

Special Populations. Pediatric Dosage. PO for Lennox–Gastaut syndrome in patients receiving enzyme-inducing antiepileptic drugs with valproic acid 0.15 mg/kg/day in 1–2 divided doses for 2 weeks and then increase to 0.3 mg/kg/day in 1–2 divided doses. Increase by 0.3 mg/kg/day at weekly intervals if needed. PO for epilepsy (Lennox–Gastaut syndrome only) in patients receiving enzyme-inducing antiepileptic drugs 0.6 mg/kg/day in 1–2 divided doses for 2 weeks and then increase to 1.2 mg/kg/day in 1–2 divided doses. Increase in increments of 1.2 mg/kg/day at weekly intervals, if needed.

Geriatric Dosage. Dosage reduction might be required in patients with reduced hepatic or renal function and should be guided by clinical response.

Other Conditions. Patients with chronic renal failure or liver disease might require lower dosages of lamotrigine; specific dosage guidelines are not available.

Dosage Forms. Chew Tab 5, 25 mg; Tab 25, 100, 150, 200 mg.

Patient Instructions. (See Anticonvulsants Class Instructions.) Inform your physician immediately if a skin rash develops.

Missed Doses. Take this drug at regular intervals. If you miss a dose, take it as soon as you remember. If it is about time for the next dose, take that dose only. Do not double the dose or take extra.
Pharmacokinetics. **Onset and Duration.** Steady-state serum levels are attained in 2–3 days.²

**Serum Levels.** A therapeutic range has not been established. In most clinical trials, trough serum concentrations of lamotrigine were 1–3 mg/L.⁴⁴

**Fate.** The drug is rapidly absorbed, with a bioavailability of 98 ± 5%; peak serum levels occur 2.8 ± 1.3 hr after an oral dose. Food does not affect absorption. Lamotrigine is 56% bound to plasma proteins. $V_d$ is 1.2 ± 0.12 L/kg.⁴⁵ $Cl$ is 0.049 ± 0.028 L/hr/kg in adults.⁴⁶ From 7% to 30% is excreted unchanged in urine; 80–90% is excreted as the inactive glucuronide conjugate.⁴⁵

$t_{1/2}$, 24.1 ± 5.7 hr in normal volunteers taking no other medications;⁴⁵ 14.3 ± 6.9 hr in patients taking enzyme-inducing antiepileptic drugs; 29.6 ± 10 hr in patients taking enzyme-inducing antiepileptic drugs with valproic acid;⁴⁶ 59 hr in patients taking valproic acid alone.⁴⁵

**Adverse Reactions.** Dose-related dizziness, ataxia, somnolence, headache, diplopia, nausea, vomiting, and rash occur frequently. Rash occurs in about 10% of patients, usually within 4–6 weeks of treatment initiation. The rash is usually maculopapular and erythematous. Potentially life-threatening rashes (including Stevens–Johnson syndrome and toxic epidermal necrolysis) are reported in 1:1000 adults and as many as 1:100 children. Risk factors for rash include concomitant valproic acid therapy, high initial dosage of lamotrigine, and rapid escalation of lamotrigine dosage. Discontinue lamotrigine at the first sign of rash.

**Contraindications.** None known.

**Precautions.** Initiate lamotrigine cautiously in patients taking valproic acid because of a higher risk of rash. (See Administration and Adult Dosage.)

**Drug Interactions.** Dizziness, diplopia, and ataxia are more common in patients taking carbamazepine concomitantly and appear to be the result of a pharmacodynamic interaction.⁴⁷ Lamotrigine has no important effect on blood levels of phenytoin, carbamazepine, or its metabolite CBZ-E. Lamotrigine reduces steady-state valproic acid levels by 25%. Valproic acid increases lamotrigine levels by about 2-fold, and carbamazepine, phenobarbital, primidone, and phenytoin each decrease lamotrigine serum levels.

**Parameters to Monitor.** Serum level monitoring is of limited value because of the lack of a well-defined therapeutic range. Routine monitoring of clinical laboratory parameters during lamotrigine therapy is not necessary.

**Notes.** Lamotrigine is indicated as adjunctive treatment for partial and secondarily generalized seizures in adults and for the treatment of the Lennox–Gastaut syndrome in adults and children.⁴⁸ Lamotrigine can be effective as monotherapy and appears to be better tolerated than carbamazepine monotherapy at dosages that are equally effective for the treatment of partial epilepsy.⁴⁹ (See Anticonvulsants Comparison Chart.)

**LEVETIRACETAM** Keppra

**Pharmacology.** Levetiracetam is a pyrrollidine derivative that is structurally unrelated to other antiepileptic drugs. Its mechanism of action is unclear and does not relate to any known mechanisms of neuronal excitation or inhibition. The action
of levetiracetam in animal models of seizures and epilepsy is unique from other antiepileptic drugs.50

Administration and Adult Dosage. PO for epilepsy 500 mg bid initially, increasing in increments of 1 g/day at 2-week intervals as needed. Usual maintenance dosage 2–3 g/day in 2 divided doses. Higher dosages have been used, but there is little evidence of increased effectiveness above 3 g/day.

Special Populations. Pediatric Dosage. Safety and efficacy not established.

Geriatric Dosage. Lower dosages might be required because of age-related decreases in renal function.51

Other Conditions. Dosage reduction is not necessary for patients with hepatic impairment. Reduce dosage in patients with compromised renal function, as indicated below:

<table>
<thead>
<tr>
<th>CLcr (ML/Min)</th>
<th>Dosage Regimen</th>
</tr>
</thead>
<tbody>
<tr>
<td>&gt;80</td>
<td>0.5–1.5 g bid</td>
</tr>
<tr>
<td>50–80</td>
<td>0.5–1 g bid</td>
</tr>
<tr>
<td>30–50</td>
<td>0.25–0.75 g bid</td>
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<tr>
<td>&lt;30</td>
<td>0.25–0.5 g bid</td>
</tr>
<tr>
<td>ESRD with hemodialysis</td>
<td>0.5–1 g/day*</td>
</tr>
</tbody>
</table>

*Supplemental doses of 250–500 mg recommended after dialysis.

Dosage Forms. Tab 250, 500, 750 mg.

Patient Instructions. (See Anticonvulsants Class Instructions.)

Missed Doses. Take this drug at regular intervals. If you miss a dose, take it as soon as you remember. If it is about time for the next dose, take that dose only. Do not double the dose or take extra.

Pharmacokinetics. Onset and Duration. Steady-state serum levels are attained in 2 days.51

Serum Levels. Not established.

Fate. Rapidly and completely absorbed within 1–1.5 hr; food has no effect on bioavailability. The drug is largely unbound to plasma proteins (<10% bound). Vd is 0.5–0.7 L/kg in adults; Cl is 0.96 mL/kg/min in adults with normal renal function; reduced by 38% in elderly patients with Clcr of 30–74 mL/min. Levetiracetam is eliminated primarily as unchanged drug in urine (66% of an administered dose). Metabolism is a minor route of elimination; three inactive metabolites have been identified. CYP pathways are not involved.51

t1/2. (Adults with normal renal function) 7 ± 1 hr.

Adverse Reactions. Somnolence, asthenia (lack of energy), infection, and dizziness occur frequently. Symptoms are of mild to moderate severity and usually occur within the first 4 weeks of treatment. Skin rash is rare.

Contraindications. None known.
Precautions. Minor decreases in RBC and WBC counts, hemoglobin, and hematocrit have been seen. The clinical importance of these findings appears to be minimal.

Drug Interactions. Levetiracetam does not induce or inhibit hepatic microsomal (CYP) enzymes and does not affect the metabolism of other antiepileptic drugs, oral contraceptives, warfarin, or digoxin. Antacids have no effect on levetiracetam bioavailability.

Parameters to Monitor. Serum level monitoring is not of value because of the lack of a defined therapeutic range. Routine monitoring of clinical laboratory parameters during levetiracetam therapy is not required.

Notes. Levetiracetam is indicated as adjunctive treatment for partial and secondarily generalized tonic-clonic seizures in adults. Seizures are reduced in frequency by ≥50% in 20–40% of patients taking 1–3 g/day.

Oxcarbazepine

Pharmacology. Oxcarbazepine is a 10-keto analogue of carbamazepine that exerts its anticonvulsant effect through an active 10-monohydroxy metabolite (MHD). Its mechanism of action is not known but likely involves blockade of voltage-dependent sodium channels and inhibition of repetitive neuronal firing.

Administration and Adult Dosage. PO for epilepsy 300 mg bid initially, increasing in increments of 300 mg/day at weekly intervals to effective dosage. Usual maintenance dosage is 1200–2400 mg/day in 2 divided doses.52

Special Populations. Pediatric Dosage. PO for epilepsy 8–10 mg/kg/day in 2 divided doses initially, increasing at weekly intervals as needed. Usual maintenance dosage in 2 divided doses: (20–29 kg) 900 mg/day; (29.1–39 kg) 1200 mg/day; (>39 kg) 1800 mg/day.

Geriatric Dosage. Clearance of the active MHD is reduced in some elderly patients, so lower maintenance dosages might be required.52

Other Conditions. No dosage adjustment is required in patients with mild to moderate hepatic impairment. Begin oxcarbazepine at one-half the usual starting dosage in patients with Clcr <30 mL/min and reduce the rate of titration.

Dosage Forms. Tab 150, 300, 600 mg.

Patient Instructions. (See Anticonvulsants Class Instructions.) Report symptoms of nausea, malaise, headache, lethargy, or confusion.

Missed Doses. Take this drug at regular intervals. If you miss a dose, take it as soon as you remember. If it is about time for the next dose, take that dose only. Do not double the dose or take extra.

Pharmacokinetics. Onset and Duration. Steady-state serum levels of MHD are attained in 2 days.53

Serum Levels. A therapeutic range has not been established.

Fate. Completely absorbed; food has no effect. Oxcarbazepine is converted into MHD, which is primarily responsible for the anticonvulsant activity of oxcarbazepine. MHD is 40% bound to plasma proteins, primarily albumin. Vd (MHD)
is 0.7–0.8 L/kg in adults; Cl (MHD) is 2.5 ± 0.1 L/kg/hr after a single dose in epilepsy patients taking other antiepileptic drugs. MHD is eliminated primarily by glucuronidation to inactive products (47%) and renal excretion of unchanged MHD (27%).

$t_{1/2}$. (Oxcarbazepine) 2.4 ± 1.1 hr; (MHD) 9.3 ± 1.8 hr in healthy adult volunteers.

**Adverse Reactions.** Dizziness, somnolence, diplopia, fatigue, and nausea occur frequently. Symptoms are more common with rapid dosage titration. Rash occurs in 2.8% of patients. Among patients with histories of hypersensitivity to carbamazepine, 25–30% experience hypersensitivity to oxcarbazepine. Hyponatremia (Na <125 mEq/L) occurs in 2.5% of patients.

**Contraindications.** None known.

**Precautions.** Patients with histories of severe hypersensitivity reactions to carbamazepine (eg, exfoliative dermatitis) appear to be at high risk for similar reactions to oxcarbazepine. Patients should report symptoms of nausea, malaise, headache, lethargy, or confusion, which might indicate hyponatremia.

**Drug Interactions.** Oxcarbazepine does not induce its own metabolism but does reduce estrogen and progesterin levels by 50%. Thus, the efficacy of oral contraceptives might be reduced. Oxcarbamazepine can increase phenytoin levels by up to 40% in adults; no effects on the metabolism of other antiepileptic drugs are reported. Carbamazepine, phenytoin, and phenobarbital increase the metabolism of MHD. Cimetidine, erythromycin, and propoxyphene do not affect MHD levels. Verapamil reduces MHD concentrations by 20%.

**Parameters to Monitor.** Consider measuring serum sodium levels during oxcarbazepine therapy, particularly in patients taking other drugs known to reduce sodium concentrations or those who develop signs or symptoms of hyponatremia. (See Precautions.)

**Notes.** Oxcarbazepine is indicated as monotherapy and adjunctive therapy for the treatment of partial and secondarily generalized seizures in adults and as adjunctive therapy for the treatment of partial-onset seizures in children (4–16 yr).

**PHENOBARBITAL**

**Pharmacology.** Phenobarbital is a barbiturate that exerts an anticonvulsant effect by depressing excitatory postsynaptic seizure discharge and increasing the convulsive threshold for electric and chemical stimulation.

**Administration and Adult Dosage.** PO or IM for epilepsy 60–90 mg/day initially, increasing in 30–60 mg/day increments q 7–14 days to an effective dosage. **Usual maintenance dosage** 90–240 mg/day or 1–3 mg/kg/day hs.1 **IV for status epilepticus** 20 mg/kg at a rate of 100 mg/min.1 **Rectal administration** has been reported. (See Fate.) (See also Adverse Reactions and Notes.)

**Special Populations.** **Pediatric Dosage.** PO or IM for epilepsy 0.5 mg/kg/day initially, increasing q 7–14 days to minimize sedation. **Usual maintenance dosage** is 2–5 mg/kg/day or 125 mg/m²/day given at bedtime.1 **IV for status epilepticus** 20 mg/kg at a rate of 50–100 mg/min.1 (See Adverse Reactions and Notes.)
Geriatric Dosage. Clearance of phenobarbital is reduced in the elderly, so lower maintenance dosages might be required.

Other Conditions. During pregnancy, phenobarbital clearance can increase. Dosage increases might be necessary and should be guided by serum levels and patient status.

Dosage Forms. Cap 16 mg; Tab 15, 16, 30, 60, 90, 100 mg; Elxr 3, 4 mg/mL; Inj 30, 60, 65, 130 mg/mL.

Patient Instructions. (See Anticonvulsants Class Instructions.)

Missed Doses. Take this drug at regular intervals. If you miss a dose, take it as soon as you remember. If it is about time for the next dose, take that dose only. Do not double the dose or take extra.

Pharmacokinetics. Onset and Duration. Steady-state serum levels are attained in about 21 days.

Serum Levels. (Anticonvulsant) 15–35 mg/L (65–150 μmol/L); dysarthria, ataxia, and nystagmus appear as serum level approaches 40 mg/L (172 μmol/L).1,2,58

Fate. The drug is slowly absorbed orally with 95–100% bioavailability; peak serum level occurs 2–4 hr after a PO or IM dose.2 Rectal bioavailability is 90%, with a peak of 7.2 ± 0.8 mg/L (31 ± 3.4 μmol/L) 4.4 ± 0.6 hr after rectal administration of a 5 mg/kg dose of parenteral phenobarbital sodium solution.2,59 The drug is 45–60% bound to plasma proteins. Vd is 0.61 ± 0.05 L/kg; Cl is 0.004 ± 0.0008 L/hr/kg in adults,60 with 50–80% metabolized in the liver to p-hydroxyphenobarbital (inactive). The drug is 20–50% excreted unchanged in urine; alkalization of urine increases renal phenobarbital clearance.2,6

\[ t_{1/2} \] (Adults) 100 ± 17 hr;61 (cirrhosis) 130 ± 15 hr;60 (children 1–5 yr) 69 ± 3.2 hr.

Adverse Reactions. (See Serum Levels.) Sedation is frequent and dose related; tolerance usually develops with long-term administration. In adults, phenobarbital can impair cognition, reaction time, and motor performance.63 Loss of concentration, mental dulling, depression of affect, insomnia, and hyperkinetic activity occur frequently with long-term therapy in children and the elderly.64 Connective tissue disorders associated with barbiturates occur in 6% of patients, usually within the first year of treatment.64 Occasionally, skin rashes or folate deficiency occur. Rarely, megaloblastic anemia, hepatitis, exfoliative dermatitis, or Stevens–Johnson syndrome is reported. Patients can be at risk for similar hypersensitivity reactions if rechallenged with phenytoin or carbamazepine.65 Neonatal hemorrhage has been reported in newborns whose mothers were taking phenobarbital. SC or intra-arterial injection can produce tissue necrosis. IV administration, especially when given after IV benzodiazepines, can produce severe respiratory depression and provision for respiratory support should be made.

Contraindications. History of porphyria or severe respiratory disease where dyspnea or obstruction is present.

Precautions. Pregnancy; lactation. Use with caution in patients with marked liver or renal disease because drug clearance is slowed. Abrupt withdrawal of the drug in patients with epilepsy can precipitate status epilepticus.
**Drug Interactions.** Concurrent use with other CNS depressants can potentiate the sedation caused by phenobarbital. Numerous drugs can increase phenobarbital serum levels, possibly requiring phenobarbital dosage reduction; phenobarbital can stimulate CYP2D6 and CYP3A and increase the metabolism of many drugs.

**Parameters to Monitor.** Periodic serum level monitoring, after attaining steady state (about 21 days), is useful in guiding dosage changes or evaluating adverse effects. Monitor CBC and liver function tests periodically during long-term therapy.

**Notes.** In tonic-clonic status epilepticus, phenobarbital is usually considered a third agent after IV phenytoin plus IV diazepam or lorazepam have failed to control seizures.\(^5\) Considering clinical efficacy and patient tolerance, phenobarbital is a third- or fourth-line choice for single-drug therapy of partial or generalized tonic-clonic seizures compared with the drugs of first choice: carbamazepine, phenytoin, or valproic acid.\(^5\) (See Anticonvulsants Comparison Chart.)

**Pharmacology.** Phenytoin is a hydantoin that suppresses the spread of seizure activity mainly by inhibiting synaptic post-tetanic potentiation and blocking the propagation of electric discharge. Phenytoin might decrease sodium transport and block calcium channels at the cellular level to produce these actions.

**Administration and Adult Dosage.** PO maintenance dosage 300 mg/day in 1–3 doses initially. Using serum levels as a guide, increase in 30–100 mg/day increments q 10–21 days to effective dosage.\(^2\) Because of dose-dependent saturable metabolism, small increases in dosage can produce disproportionate increases in serum levels. Usual maintenance dosage 300–400 mg/day or 4–8 mg/kg/day in 1 or 2 doses.\(^1\) Only extended-release phenytoin sodium capsules are approved for once-daily administration. PO loading dosage 15 mg/kg in 3 divided doses, administered at 2-hr intervals. Using serum levels as a guide, a maintenance dosage can be initiated within 24 hr of starting the loading dosage. IV loading dose 15–20 mg/kg by direct IV injection, at a rate not greater than 50 mg/min or 0.75 mg/kg/min in adults.\(^58\) Therapeutic serum levels persist for 12–24 hr in most patients.\(^66\) Alternatively, dilute the loading dose in 50–150 mL of 0.45% or 0.9% NaCl and infuse through an IV volume control set with an in-line filter at a rate not greater than 50 mg/min.\(^2\)\(^67\) In nonemergency situations, an IV dose of 5 mg/kg q 2 hr for 3 doses at a rate of 50 mg/min results in phenytoin serum levels of 10–20 mg/L 12 hr after the third dose.\(^68\) (See Adverse Reactions and Notes.) IM administration is painful and results in slow, but complete, absorption because of deposition of phenytoin crystals in muscle.\(^2\) The IM route is not recommended. The IV route is preferred in patients unable to take phenytoin by mouth. (See Fosphenytoin.)

**Special Populations.** Pediatric Dosage. PO maintenance dosage 5 mg/kg/day initially in 2–3 divided doses. Increase initial dosage in small increments q 7–10 days to effective dosage.\(^2\) Because of dose-dependent metabolism, small increases in dosage can produce disproportionate increases in serum levels. Usual maintenance dosage 4–8 mg/kg/day in 2–3 divided doses.\(^1\) IV loading dose (neonates)
15–20 mg/kg given at a rate of 0.5 mg/kg/min; (older infants and children) same as adult dosage.

**Geriatric Dosage.** PO maintenance dosage 200–300 mg/day in 1–3 doses. Advanced age can be associated with a decrease in phenytoin clearance and a reduction in albumin concentration.69 Dosage adjustment should be guided by phenytoin levels and patient status. (See Serum Levels.)

**Other Conditions.** During pregnancy or febrile illness or after acute traumatic injury, phenytoin clearance can increase; dosage adjustment might be necessary and should be guided by serum levels and patient status.70–72 Renal disease and hypoalbuminemia can alter phenytoin binding to plasma proteins, resulting in a change in the usual ratio of free to total phenytoin levels; renal disease alters phenytoin protein binding because of decreased affinity of plasma proteins. Increases in fraction unbound are most pronounced in patients with Clcr <25 mL/min. Ideally, adjust dosage guided by patient status and actual measurement of unbound and total phenytoin levels. (See Serum Levels.)

**Dosage Forms.** (Phenytoin) Chew Tab 50 mg; Susp 25 mg/mL. (Phenytoin sodium) Cap (extended or prompt) 30, 100 mg; Inj 50 mg/mL. Phenytoin sodium is 92% phenytoin.

**Patient Instructions.** (See Anticonvulsants Class Instructions.) Good dental hygiene and regular dental visits can minimize gum tenderness, bleeding, or enlargement (especially in children). Shake oral suspension well before each dose and use a calibrated measuring device. (See Notes.) Call physician if skin rash develops.

**Missed Doses.** Take this drug at regular intervals. If you miss a dose, take it as soon as you remember. If it is about time for the next dose, take that dose only. When taking multiple daily doses, leave a minimum of 4–6 hours between doses. If you are taking the drug only once daily and you do not remember until the next day, skip the missed dose and return to your normal schedule. Do not double the dose or take extra. If doses are missed for 2 or more days in a row, consult your physician.

**Pharmacokinetics. Onset and Duration.** Time to steady state increases with increasing dosage and serum level. Steady state is usually attained within 7–14 days but can take as long as 28 days.2

**Serum Levels.** 10–20 mg/L (40–80 μmol/L) in patients with normal renal function and serum albumin concentration. Nystagmus, slurred speech, ataxia, or dizziness appear in most patients as serum levels approach 20 mg/L; drowsiness, diplopia, behavioral changes, and cognitive impairment occur with serum levels above 30 mg/L (120 μmol/L).2 The equation \( C_{normal} = C_{observed} / ([0.2 \times \text{albumin}] + 0.1) \) estimates the concentration of phenytoin that would be expected if the albumin concentration were normal \( (C_{normal}) \) from the measured total phenytoin concentration in a hypoalbuminemia patient \( (C_{observed}) \) and the patient’s albumin concentration in g/dL (albumin). In patients with end-stage renal disease \( (\text{Clcr} < 10 \text{ mL/min}) \), the equation \( C_{normal} = C_{observed} / ([0.1 \times \text{albumin}] + 0.1) \) is used.73 (See also Special Populations, Other Conditions.)

**Fate.** Oral phenytoin absorption is very slow and incomplete in infants <3 months of age.2 Bioavailability of the suspension is decreased in patients receiving con-
comitant enteral feedings. (See Precautions.) IM injection is slowly absorbed over several days because of deposition of phenytoin crystals in muscle. Peak serum levels occur 4–8 hr after a single dose of an extended-release capsule; after oral loading given in divided doses, the times to reach a serum level of 10 mg/L are 4.5 ± 2.1 hr for prompt-release capsules and 9.6 ± 2.5 hr for extended-release capsules. Time to peak serum level increases with increasing oral dosages. About 90% is bound to plasma proteins. Hypoalbuminemia, chronic liver or renal disease, nephrotic syndrome, AIDS, or acute traumatic injury alter protein binding and increase the fraction of unbound phenytoin. $V_d$ is 0.83 ± 0.2 L/kg in adults with acute seizures and 0.79 ± 0.25 L/kg in critically ill adults after trauma. Hepatic metabolism is capacity limited, exhibiting Michaelis–Menden pharmacokinetics; therefore, Cl decreases as serum level increases. Mean apparent $V_{max}$ is 0.45 mg/L/hr; mean apparent $K_m$ is 6.2 mg/L in adults. About 70% of phenytoin is excreted in urine as the inactive metabolite, 5-(p-hydroxyphenyl)-5-phenyl hydantoin. Less than 5% of the parent drug is excreted unchanged in urine.

**Adverse Reactions.** (See Serum Levels.) Erythematous morbilliform rash occurs frequently. Do not resume phenytoin if rash is exfoliative, purpuric, bullous, or accompanied by fever. With long-term administration, hirsutism, gingival hypertrophy (especially in children and adolescents), coarsening of facial features, acniform eruption, osteomalacia, and folate deficiency with mild macrocytosis occur frequently. Bradycardia or hypotension caused by rapid IV administration are reported occasionally; slowing the rate of administration can minimize these complications. Severe soft tissue injury after IV phenytoin is more likely in elderly (>70 yr) women who receive 2 or more infusions through small (<20 gauge) IV devices. Hepatotoxicity occurs occasionally, usually within the first 6 weeks, and presents with fever, rash, lymphadenopathy, and hepatomegaly. Other idiosyncratic reactions are rare, can occur together within the first 2 months, and include fever, lymphoid hyperplasia, eosinophilia, erythema multiforme, exfoliative dermatitis, Stevens–Johnson syndrome, leukopenia, anemia, thrombocytopenia, serum sickness, and SLE. These patients are at risk for similar hypersensitivity reactions if rechallenged with phenobarbital or carbamazepine. Concurrent cranial irradiation predisposes patients to the development of erythema multiforme. A syndrome of anomalies in infants of phenytoin-exposed mothers has been described (fetal hydantoin syndrome).

**Contraindications.** (Parenteral phenytoin) sinus bradycardia; sinoatrial block; second- and third-degree AV blocks; Adams–Stokes syndrome.

**Precautions.** Pregnancy; lactation. Use with caution in patients with severe liver disease or diabetes or with histories of severe hypersensitivity reactions to carbamazepine or phenobarbital. Abrupt withdrawal of the drug in patients with epilepsy can precipitate status epilepticus. If the patient’s nutritional status allows, interrupt tube feeding 2 hr before and after the dose and irrigate the feeding tube to improve absorption; nevertheless, the patient might require an increase in phenytoin dosage. If the feedings are discontinued after the phenytoin dosage is
increased, the dosage must be adjusted to prevent toxic serum levels from occurring.\textsuperscript{30}

**Drug Interactions.** Chronic alcohol use, barbiturates, rifampin, and some other drugs can stimulate phenytoin metabolism and increase phenytoin dosage requirements. Numerous drugs can increase phenytoin serum levels, possibly requiring phenytoin dosage reduction; phenytoin can stimulate CYP2D6 and CYP3A and increase the metabolism of many drugs. IV phenytoin can produce hypotension in severely ill patients receiving IV dopamine. The antiparkinson effect of levodopa can be inhibited by phenytoin.

**Parameters to Monitor.** Serum level monitoring, after attaining steady state (10–21 days), is useful in evaluating therapeutic efficacy or potential for adverse effects.\textsuperscript{2} Patient and serum level monitoring are recommended when changing phenytoin dosage form or brand; monitor serum levels q 5–7 days to assess trend in concentrations. Monitor CBC and liver function tests periodically with long-term therapy.

**Notes.** Agitation or shaking is needed to resuspend phenytoin suspension; settling occurs 5 weeks after resuspension.\textsuperscript{81} In tonic-clonic status epilepticus, the anticonvulsant effect of phenytoin appears 20–30 min after start of the infusion. Thus, in this situation, concurrent use of phenytoin with a rapidly acting injectable benzodiazepine (diazepam or lorazepam) is recommended.\textsuperscript{56} Phenytoin is recommended, as is carbamazepine, as a drug of first choice for single-drug therapy of partial or generalized tonic-clonic seizures.\textsuperscript{82} (See Anticonvulsants Comparison Chart.)

**PRIMIDONE**

**Pharmacology.** Primidone (desoxyphenobarbital) is structurally related to the barbiturates. Primidone and its metabolites, phenylethylmalonamide (PEMA) and phenobarbital, exert anticonvulsant activity. (See Phenobarbital.)

**Administration and Adult Dosage.** PO for epilepsy 50–100 mg hs initially; increase in 100–125 mg/day increments q 2–3 days to effective dosage, to a maximum of 2 g/day. **Usual maintenance dosage** 250–500 mg tid.\textsuperscript{1}

**Special Populations.** Pediatric Dosage. PO for epilepsy (<8 yr) 50 mg hs initially, increasing by 50 mg in 3 days and thereafter in 100–125 mg/day increments q 3 days to effective dosage; **usual maintenance dosage** is 125–250 mg tid or 10–20 mg/kg/day in 3 divided doses; (≥8 yr) same as adult dosage.

**Geriatric Dosage.** Clearance of primidone is unchanged in elderly patients, but phenobarbital clearance is reduced. Lower maintenance dosages might be required.

**Dosage Forms.** **Susp** 50 mg/mL; **Tab** 50, 250 mg.

**Patient Instructions.** (See Anticonvulsants Class Instructions.)

**Missed Doses.** Take this drug at regular intervals. If you miss a dose, take it as soon as you remember. If it is about time for the next dose, take that dose only. Do not double the dose or take extra.
Pharmacokinetics. Onset and Duration. Steady-state serum levels are attained in about 3 days.  

Serum Levels. (Primidone) 6–12 mg/L (28–55 μmol/L).1,2,58 (See also Phenobarbital.) During monotherapy, the serum level ratio of phenobarbital to primidone is about 1:1.2 During polytherapy with enzyme-inducing agents, this ratio increases to 4:1.2 During monotherapy, the serum level ratio of PEMA to primidone at steady state is 0.74 ± 0.38 for samples drawn before the first morning dose.2 

Fate. The drug is rapidly absorbed with 90–100% bioavailability; peak serum levels occur 2–6 hr after an oral dose. The drug is 0–20% bound to plasma proteins; \( V_d \) is 0.86 ± 0.22 L/kg;83 \( Cl \) with monotherapy is 0.035 ± 0.02 L/hr/kg; with concomitant anticonvulsants, it is 0.052 ± 0.02 L/hr/kg; in monotherapy with concomitant acute viral hepatitis, it is 0.042 ± 0.14 L/hr/kg.83 Primidone is metabolized in the liver to PEMA and phenobarbital; 76% of the drug is excreted into the urine within 5 days after a single dose as 64% primidone, 7% PEMA, 2% phenobarbital, and 3% unidentified products.5 

\( t_{1/2} \). (Primidone) 15.2 ± 4.8 hr (monotherapy); 8.3 ± 2.9 hr (with concomitant enzyme-inducing anticonvulsants); 18 ± 3.1 hr (with acute viral hepatitis).83 (PEMA) 21 ± 3 hr (primidone monotherapy); 17 ± 4.3 hr (with concomitant anticonvulsants).84 (See also Phenobarbital.) 

Adverse Reactions. Drowsiness, ataxia, nausea, weakness, and dizziness occur frequently during the first month of therapy and might become tolerable with time. Behavioral disturbances, depression of affect, and cognitive impairment occur frequently with long-term therapy in children and the elderly.2 Occasionally, skin rashes and, rarely, impotence, leukopenia, thrombocytopenia, megaloblastic anemia, or lymphadenopathy occur.2,58 

Contraindications. History of porphyria; hypersensitivity to phenobarbital. 

Precautions. Pregnancy; lactation. Use with caution in patients with severe liver or renal disease. Abrupt withdrawal of the drug can precipitate status epilepticus. (See Phenobarbital Notes.) 

Drug Interactions. Concurrent use with other CNS depressants can potentiate the sedation caused by phenobarbital. Primidone levels can be decreased by concurrent acetazolamide, carbamazepine, or succinimides. Primidone levels can be increased by concurrent hydantoins, isoniazid, or niacinamide. 

Parameters to Monitor. Periodic serum level monitoring of primidone and phe- nobarbital, after attaining steady state (primidone, 3 days; phenobarbital, 21 days), is useful in guiding dosage changes, detecting noncompliance, or evaluating adverse effects.2 Monitor CBC, electrolytes, and liver function tests periodically during long-term therapy. 

Notes. Considering comparative efficacy and good patient tolerance of carbamazepine, phenytoin, and valproic acid, primidone is a fourth- or fifth-line choice for single-drug therapy of generalized tonic-clonic seizures. In a large, multicenter trial comparing carbamazepine, phenytoin, phenobarbital, and primidone, primidone was least successful in controlling seizures with acceptable adverse effects.52
Pharmacology. Tiagabine is a nipecotic acid derivative unrelated to other marketed antiepileptic drugs. It interacts with the GABA uptake carrier and is thought to enhance the inhibitory effect of GABA by preventing its reuptake into neurons. Tiagabine is indicated in adults and adolescents (>12 yr) as adjunctive therapy for patients with partial-onset seizures.85,86

Administration and Adult Dosage. PO for epilepsy in patients taking enzyme-inducing antiepileptic drugs (eg, carbamazepine, phenytoin, phenobarbital, primidone), initiate at 4 mg/day for 1 week, increasing in increments of 4–8 mg/day at weekly intervals according to clinical response. **Usual maintenance dosage** is 32–56 mg/day. The daily dosage is given in 2–4 divided doses; with dosages >32 mg/day, tid or qid administration might be required. Patients taking only non–enzyme-inducing antiepileptic drugs (eg, gabapentin, lamotrigine, valproate) might require lower doses or a slower titration schedule.

Special Populations. **Pediatric Dosage.** (≤12 yr) safety and efficacy not established; (>12 yr) same as adult dose.

**Geriatric Dosage.** Same as adult dosage.

**Other Conditions.** No apparent need to adjust tiagabine dosage in renal impairment. Patients with liver disease might require lower dosages of tiagabine; however, specific dosage guidelines are not available.

**Dosage Forms.** Tab 2, 4, 12, 16, 20 mg.

**Patient Instructions.** (See Anticonvulsants Class Instructions) Take this medication with food.

**Missed Doses.** Take this drug at regular intervals. If you miss a dose, take it as soon as you remember. If it is about time for the next dose, take that dose only. Do not double the dose or take extra.

**Pharmacokinetics. Onset and Duration.** Steady-state serum levels are attained within 2 days.

**Serum Levels.** A therapeutic range has not been established.

**Fate.** Rapidly absorbed with a bioavailability of 89.9 ± 7.7%.87,88 Food slows the rate, but not the extent, of tiagabine absorption. Peak serum concentrations of 241 ± 79 μg/L occurred 1.3 ± 1 hr after a single 12 mg dose in healthy volunteers.89 Tiagabine is 96% bound to plasma proteins, primarily albumin and α1-acid glycoprotein. Vd is 1.07 ± 0.22 L/kg89 and Cl is 6.5 ± 1.5 L/hr in healthy volunteers.88 Cl is increased in patients taking enzyme-inducing antiepileptic drugs. Tiagabine is metabolized by oxidation (primarily CYP3A) and glucuronidation; about 2% is excreted unchanged in urine.

**t½.** 7–9 hr in healthy subjects taking no other medications; reduced by 50–65% in patients with epilepsy taking enzyme-inducing antiepileptic drugs.88,89

**Adverse Reactions.** Dizziness, asthenia/lack of energy, somnolence, nausea, nervousness, tremor, abdominal pain, and difficulty with concentration occur frequently. Tremor, difficulty with concentration, and asthenia appear to be dose related. The most common reasons for discontinuation are dizziness, somnolence,
depression, confusion, and asthenia. Moderate to severe generalized weakness occurs occasionally. Tiagabine rarely induces absence status.

**Contraindications.** None known.

**Precautions.** Dosage reduction might be required in hepatic impairment.

**Drug Interactions.** Carbamazepine, phenytoin, and phenobarbital reduce tiagabine levels by 60% compared with noninduced patients. Valproate has no important effect on tiagabine levels. Tiagabine has no effect on serum concentrations of phenytoin, carbamazepine, phenobarbital, or primidone. Valproate concentrations decrease approximately 10% during tiagabine therapy. Tiagabine has no effect on the pharmacokinetics of warfarin, theophylline, digoxin, or oral contraceptives.

**Parameters to Monitor.** Serum level monitoring is of limited value because of the lack of a well-defined therapeutic range. No routine laboratory test monitoring is required during therapy.

### TOPIRAMATE

**Pharmacology.** Topiramate, a derivative of the naturally occurring monosaccharide D-fructose, reduces the frequency of action potentials elicited by depolarizing currents in a manner suggestive of sodium-channel blocking action. Topiramate also increases GABA-induced chloride flux, although the drug has no direct effect on GABA binding sites. It also inhibits kainate activation of a subtype of the excitatory glutamate receptor. Topiramate inhibits carbonic anhydrase, but this action might not contribute to the drug’s anticonvulsant effect. *(See Notes.)*

**Administration and Adult Dosage.** PO for epilepsy 25–50 mg/day initially, increasing in 25–50 mg/day increments at weekly intervals to 200–400 mg/day in 2 divided doses. **Usual maintenance dosage** 400 mg/day. Higher dosages have not been shown to be more effective; however, individual patients might require ≥1 g/day. *(See Notes.)*

**Special Populations.** **Pediatric Dosage.** PO for epilepsy (<2 yr) Safety and efficacy not established; (2–16 yr) 5–9 mg/kg in 2 divided doses. Increase in increments of 1–3 mg/kg/day at weekly intervals, if needed.

**Geriatric Dosage.** Same as adult dosage.

**Other Conditions.** With Clcr <70 mL/min/1.73 m², reduce dosage by 50%. Topiramate clearance increases during hemodialysis to a rate 4–6 times greater than in a normal person. Additional doses of topiramate might be required depending on dialysis method and duration.

**Dosage Forms.** Cap 15, 25, 50 mg; Tab 25, 100, 200 mg. *(See Notes.)*

**Patient Instructions.** *(See Anticonvulsants Class Instructions.)* Maintain adequate fluid intake (6 to 8 glasses of water daily) to minimize the formation of kidney stones. If you are taking oral contraceptives, report any change in menstrual bleeding patterns to your health care provider.

**Missed Doses.** Take this drug at regular intervals. If you miss a dose, take it as soon as you remember. If it is about time for the next dose, take that dose only. Do not double the dose or take extra.
Pharmacokinetics. Onset and Duration. Steady-state serum concentrations are attained in 4 days in patients with normal renal function.

Serum Levels. A therapeutic range has not been established.\(^9\)

Fate. The bioavailability of oral tablets is 80% compared with oral solution and peak concentrations occur 3.5 ± 0.6 hr after 400 mg.\(^2\) Administration with food delays absorption (by approximately 2 hr) but does not affect extent of absorption.\(^2\) Topiramate is 13–17% bound to plasma proteins and binds to a saturable, low-capacity binding site on or in erythrocytes.\(^8\) V\(_d\) is 0.6–0.8 L/kg in healthy volunteers.\(^2\) Cl is 0.021 ± 0.004 L/kg/hr in adults on topiramate monotherapy after tapering of valproic acid.\(^9\) Approximately 70% is eliminated unchanged in urine. Six metabolites (each <5% of the administered dose) have been identified.

\(t_{\frac{1}{2}}\). 23 hr after a single 400 mg dose in healthy adults.\(^2\)

Adverse Reactions. Somnolence, dizziness, ataxia, speech problems, psychomotor slowing, nystagmus, and paresthesias occur commonly and are not dose related. Common dose-related adverse effects include fatigue, nervousness, difficulty with concentration or attention, confusion, depression, weight loss, and tremor. These adverse effects are minimized by slow dose titration. Psychomotor slowing and difficulty with concentration are the most common reasons for topiramate discontinuation. Kidney stones occur in 1.5% of patients and might be related to carbonic anhydrase inhibition.

Contraindications. None known.

Precautions. Avoid concomitant use of other carbonic anhydrase inhibitors (eg, acetazolamide) because of the potential increased risk of kidney stones.

Drug Interactions. Concomitant phenytoin and carbamazepine reduce topiramate concentrations by 48% and 40%, respectively. Concomitant valproate reduces topiramate concentrations by 17%.\(^9\) Topiramate variably affects phenytoin concentrations (0–25% decrease) and has no important effect on other anticonvulsants. Topiramate can reduce the effectiveness of oral contraceptives; consider using products containing ≥35 µg of ethinyl estradiol.\(^9\)

Parameters to Monitor. Serum level monitoring is of limited value because of the lack of a well-defined therapeutic range. Routine monitoring of clinical laboratory parameters during topiramate therapy is not indicated.

Notes. Topiramate is indicated for adult and pediatric patients as adjunctive treatment for partial-onset and primary generalized tonic-clonic seizures. Preliminary evidence suggests that topiramate also might be effective as monotherapy for partial-onset seizures\(^9\) and as adjunctive treatment of Lennox–Gastaut syndrome.\(^9\) Capsules can be opened and sprinkled on food.

### VALPROIC ACID

Depakene, Depacon, Various

### DIVALPROEX SODIUM

Depakote

Pharmacology. Valproic acid is a carboxylic acid compound whose anticonvulsant activity might be mediated by an inhibitory neurotransmitter, GABA. Valproic acid might increase GABA levels by inhibiting GABA metabolism or enhancing postsynaptic GABA activity. Valproic acid also limits repetitive neuronal
firing through voltage- and usage-dependent sodium channels. Divalproex is comprised of sodium valproate and valproic acid. (See Notes.)

**Administration and Adult Dosage.** PO for epilepsy (valproic acid) 15 mg/kg/day in 2–3 divided doses initially, increasing in 5–10 mg/kg/day increments at weekly intervals to an effective dosage, to a maximum of 60 mg/kg/day. **Usual maintenance dosage** 15–40 mg/kg/day in 3 divided doses. In patients receiving valproic acid, divalproex can be substituted at the same daily dosage; in selected patients, it can be given bid. PO for migraine prophylaxis (divalproex) 250 mg bid, to a maximum of 1 g/day. PO for mania (divalproex) 750 mg/day in divided doses, increasing as rapidly as possible to the lowest dosage that produces the desired effect, to a maximum of 60 mg/kg/day. (See Serum Levels.) Long-term experience with this use is minimal and characterized by a high drop-out rate. IV for epilepsy (valproic acid) same as oral dosage. Administer infusion over 60 min or at a rate of ≤20 mg/min. Rectal administration has been reported. (See Fate.)

**Special Populations.** **Pediatric Dosage.** Same as adult dosage.

**Geriatric Dosage.** Reduce the starting dosage in the elderly. Protein binding and unbound clearance of valproic acid are reduced in the elderly, and the desired clinical response can be achieved with lower dosages than in younger adults. Adjust dosage guided by valproic acid levels (preferably free levels in patients with low serum albumin) and patient status.3

**Dosage Forms.** (Valproic acid) Cap 250 mg; Syrup 50 mg/mL; Inj 100 mg/mL; (divalproex) EC Tab 125, 250, 500 mg; Cap (EC granules) 125 mg; SR Tab 500 mg.

**Patient Instructions.** (See Anticonvulsants Class Instructions). This drug can be taken with food or milk to minimize stomach upset. Do not chew, break, or crush the tablet or capsule because this may irritate your mouth or throat. Sprinkle capsule can be swallowed whole or administered by sprinkling the entire contents on small amount (1 teaspoonful) of soft food such as pudding or applesauce; swallow the drug/food mixture immediately (avoid chewing). Polymer from the sprinkles might appear in the stools, but does not indicate a lack of absorption. Immediately report weakness, tiredness, repeated vomiting, or loss of seizure control, which might be early signs of severe, but rare, liver disorder.

**Missed Doses.** Take this drug at regular intervals. If you miss a dose, take it as soon as you remember. If it is about time for the next dose, take that dose only. Leave a minimum of 6 hours between doses. Do not double the dose or take extra.

**Pharmacokinetics.** **Onset and Duration.** Steady-state serum levels are attained in 2–4 days. Several weeks might be required to attain maximal therapeutic effect. For mania, levels should be above 45 mg/L for efficacy and below 125 mg/L to minimize adverse effects.

**Serum Levels.** 50–120 mg/L (350–830 μmol/L) for epilepsy and mania. Some patients require and can tolerate serum levels up to 150 mg/L. Tremor, irritability, confusion, and restlessness might be observed with levels >100–150 mg/L.2

**Fate.** The bioavailability of the oral capsule is 93 ± 13%, with peak levels occurring 1–2 hr after the dose. The bioavailability of the EC divalproex tablet is
90 ± 14%, with peak levels in 4 hr.\textsuperscript{2,96} The peak time of both is delayed by food: (Cap) 5.2 ± 1.7 hr; (EC divalproex tab) 8.1 ± 3.6 hr.\textsuperscript{97} Bioavailability of the SR tab is 80–90% of the EC product. Bioavailability is 80 ± 7% after a 250 mg suppository.\textsuperscript{96} Peak serum levels of 40–50 mg/L (280–350 \(\mu\)mol/L) occur 2–4 hr after a 15–20 mg/kg dose of syrup diluted 1:1 with water as a retention enema.\textsuperscript{11} \(V_d\) is 0.19 ± 0.05 L/kg in adults and 0.26 ± 0.09 L/kg in children.\textsuperscript{2} Plasma protein binding is about 90%.\textsuperscript{96} Increasing serum concentrations, hypoalbuminemia, severe liver disease, renal disease, or pregnancy reportedly increases the unbound fraction and might alter clearance. Cl is (healthy adults) 0.0066 ± 0.0005 L/hr/kg; (epileptic adults) 0.018 ± 0.011 L/hr/kg; (children) 0.027 ± 0.015 L/hr/kg.\textsuperscript{2} Over 96% is metabolized to at least 10 metabolites. Only 1.8–3.2% of drug is excreted unchanged in urine.\textsuperscript{2}

\(t_{\frac{1}{2}}\). (Healthy adults) 13.9 ± 3.4 hr; (epileptic adults) 8.5 ± 3.3 hr; (children) 7.2 ± 2.3 hr.\textsuperscript{2}

\textbf{Adverse Reactions.} (See Serum Levels.) Nausea, vomiting, diarrhea, and abdominal cramps occur frequently during initiation of therapy and are minimized by slow titration of valproic acid or substitution of EC divalproex for valproic acid. Transient elevations in liver function tests occur frequently. The risk of valproate-exposed women having children with spina bifida is approximately 1–2%. Drowsiness, ataxia, tremor, behavioral disturbances, transient hair loss, asymptomatic hyperammonemia, or weight gain occurs occasionally. Drowsiness and ataxia are more prominent in patients taking valproic acid with other anticonvulsants.\textsuperscript{2} Rarely, thrombocytopenia, acute pancreatitis, abnormal coagulation parameters, or hyperglycinemia occurs. Liver failure occurs rarely; the greatest risk is during the first 6 months of therapy and in children <2 yr who receive multiple anticonvulsants.\textsuperscript{99}

\textbf{Contraindications.} Hepatic dysfunction or disease.

\textbf{Precautions.} Pregnancy; lactation. The drug can alter results of urine ketone tests.

\textbf{Drug Interactions.} Valproate levels can be decreased by concurrent carbamazepine, lamotrigine, phenytoin, or rifampin. Valproate levels can be increased by concurrent aspirin, chlorpromazine, cimetidine, or felbamate. Lamotrigine and phenobarbital levels can be increased by valproate.

\textbf{Parameters to Monitor.} Baseline liver function tests and platelets; repeat liver function tests frequently, especially during the first 6 months. Monitor platelet count and coagulation tests before surgery. Periodic serum level monitoring is useful for guiding dosage changes and evaluating potential adverse effects. Serum levels fluctuate considerably over 24 hr, making a single random measurement of limited value. Predose blood sampling at standard times is recommended.\textsuperscript{3,96}

\textbf{Notes.} Valproic acid and ethosuximide are equally effective for treating absence seizures, although ethosuximide is sometimes preferred as a first-line agent because of its lower risk of serious toxicity. Valproic acid is preferred for patients with absence and generalized tonic-clonic seizures. Many clinicians in the United States use valproic acid as a second-line agent (after phenytoin or carbamazepine) for the treatment of partial seizures. Valproic acid is as effective as
phenytoin and carbamazepine for tonic-clonic seizures and is a drug of choice for atonic and myoclonic seizures.\(^2\) See Anticonvulsants Comparison Chart.

Divalproex is equivalent to lithium in bipolar disorder and is more effective than lithium for rapid-cycling bipolar patients (four or more episodes in 1 yr) and comorbid substance abuse. Carbamazepine is more effective as an adjunctive treatment with lithium to enhance partial efficacy than with lithium alone.\(^{101-103}\)

For migraine prophylaxis, divalproex is effective and well tolerated. It might be more effective in those having frequent migraines than in those characterized as having tension headaches.\(^{104,105}\) The sustained-release formulation is approved for migraine prophylaxis only.

### Zonisamide

**Pharmacology.** Zonisamide is a 1,2-benzisoxazole sulfonamide derivative that is chemically unrelated to other antiepileptic drugs. It blocks seizure spread and inhibits epileptic foci in animals. The anticonvulsant effect is likely related to blockade of voltage-sensitive sodium and T-type calcium channels. It is also a weak carbonic anhydrase inhibitor.\(^{106}\)

**Administration and Adult Dosage.** PO as adjunctive therapy for partial seizures 100 mg/day initially, increasing in 100 mg/day increments at intervals of 2 weeks as needed. **Usual maintenance dosage** is 200–400 mg/day in 1–2 divided doses. Some patients might require dosages of 600 mg/day; there is little evidence of increased effectiveness above 400 mg/day.

**Special Populations.** **Pediatric Dosage.** Safety and efficacy not established. Zonisamide is used in Japan for the treatment of epilepsy in children. The recommended dosage is 2–4 mg/kg/day initially, increasing at 2-week intervals to 4–12 mg/kg/day as needed.

**Geriatric Dosage.** Advanced age has no effect on zonisamide pharmacokinetics. No dosage adjustment is necessary.

**Other Conditions.** Zonisamide clearance is reduced in patients with renal disease. These patients might require slower titration because of the prolonged half-life of the drug. The effect of liver disease on zonisamide pharmacokinetics is unknown.

**Dosage Forms.** Cap 100 mg.

**Patient Instructions.** (See Anticonvulsants Class Instructions). Drink 6–8 glasses of water daily to lessen the likelihood of kidney stone formation.

**Missed Doses.** Take this drug at regular intervals. If you miss a dose, take it as soon as you remember. If it is about time for the next dose, take that dose only. Do not double the dose or take extra.

**Pharmacokinetics.** **Onset and Duration.** Steady-state serum levels are attained in 14 days in patients with normal renal function.

**Serum Levels.** Therapeutic range not established.

**Fate.** Rapidly absorbed after oral administration with peak serum concentrations occurring 2.8 ± 1.4 hr after a dose in healthy volunteers. Food delays the rate but has no effect on the extent of zonisamide absorption. Zonisamide is 40% bound to plasma proteins, mainly albumin. In healthy volunteers, \(V_d/F\) is 1.47 ± 0.39 L/kg.
Cl/F is 0.019 ± 0.004 L/kg/hr. Cl is increased 30–40% during concomitant therapy with enzyme-induced antiepileptic drugs. Elimination is mainly by urinary excretion of unchanged drug and glucuronide metabolite. Other metabolic pathways include acetylation and reduction of an acetylated metabolite (via CYP3A4).

\[ t_{1/2} \] (Adults with normal renal function) 63 hr.

**Adverse Reactions.** Frequent adverse effects include somnolence, ataxia, anorexia, confusion, abnormal thinking, and nervousness. Kidney stones occur in 2.6% of patients.²

**Contraindications.** Allergy to sulfonamides.

**Drug Interactions.** Enzyme-inducing antiepileptic drugs (carbamazepine, phenytoin, barbiturates) enhance zonisamide’s metabolism and reduce its half-life to 27–36 hr. Zonisamide has no apparent effect on the pharmacokinetics of other antiepileptic drugs.⁸⁸

**Notes.** Zonisamide is indicated as adjunctive treatment for partial and secondarily generalized tonic-clonic seizures in adults. It reduces the frequency of seizures by ≥50% in 30–40% of patients as adjunctive therapy. The drug also appears to be effective for generalized and progressive myoclonic epilepsies.²
## ANTICONVULSANTS COMPARISON CHART

### CHOICE OF ANTICONVULSANT FOR CLINICAL SEIZURE TYPE

<table>
<thead>
<tr>
<th>DRUG</th>
<th>Generalized Seizures</th>
<th>Partial Seizures&lt;sup&gt;b&lt;/sup&gt;</th>
<th>Dosage Range (mg/kg/day)</th>
<th>Therapeutic Serum Levels (mg/L) (µmol/L)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Tonic-Clonic</td>
<td>Absence</td>
<td>Myoclonic</td>
<td>Atonic</td>
</tr>
<tr>
<td>Carbamazepine</td>
<td>1</td>
<td>W</td>
<td>—</td>
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<tr>
<td>Clonazepam</td>
<td>W</td>
<td>3</td>
<td>2</td>
<td>1</td>
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<tr>
<td>Ethosuximide</td>
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<td>Gabapentin</td>
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<td>Lamotrigine</td>
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<td>Levetiracetam</td>
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<td>Oxcarbazepine</td>
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<tr>
<td>Phenytoin</td>
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<td>W</td>
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<td>Tiagabine</td>
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<td>Topiramate</td>
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<tr>
<td>Valproic Acid&lt;sup&gt;c&lt;/sup&gt;</td>
<td>1</td>
<td>2</td>
<td>1</td>
<td>1</td>
</tr>
<tr>
<td>Zonisamide</td>
<td>—</td>
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</tbody>
</table>

1 = Drug of first choice; initial agent; given as monotherapy.
2 = Drug of second choice; alternative to first choice; given as monotherapy or in combination with agent of first choice; 3 = Drug of third choice; alternative to first or second choice; given as monotherapy or in combination with another agent; 4 = Useful as adjunctive therapy after failure of monotherapy with preferred agents; W = May worsen clinical seizure type.

<sup>a</sup>Choice of anticonvulsant based on relative and comparative efficacy and potential for adverse effects. Choice of agent should consider individual patient factors. (See references 1, 82, 95, and 100.)

<sup>b</sup>Includes simple-partial, complex-partial, and secondarily generalized tonic-clonic seizures.

<sup>c</sup>Drug of first choice when both generalized tonic-clonic and absence seizures are present.
Antidepressants

**Class Instructions.** Antidepressants. This drug can cause drowsiness. Until the extent of this effect is known, use caution when driving, operating machinery, or performing other tasks requiring mental alertness. Avoid excessive concurrent use of alcohol or other drugs that cause drowsiness.

**BUPROPION**

**Pharmacology.** Bupropion is a monocyclic antidepressant, unique as a mild dopamine and norepinephrine uptake inhibitor with no direct effect on serotonin receptors or MAO. It is essentially devoid of anticholinergic, antihistaminic, and peripheral adrenergic effects. In contrast with heterocyclic antidepressants, bupropion produces no clinically important effect on cardiac conduction, no orthostatic hypotension, minimal anticholinergic effects, and it is not associated with weight gain. Compared with SSIs, bupropion offers a similar side effect profile without sexual dysfunction. Its lack of sedation and its activating effect can be advantageous for patients with decreased psychomotor activity and lethargy. Disadvantages of bupropion include seizures and the necessity of multiple daily doses. 107–109

**Administration and Adult Dosage.** Small initial doses and gradual dosage escalation is necessary to minimize the risk of seizures. PO for depression (immediate-release) 100 mg bid initially, increasing to 100 mg tid no sooner than 3 days after the start of therapy. The maximum daily dosage is 450 mg, with a maximum single dose of 150 mg; (sustained-release) 150 mg given in the morning initially, increasing to 150 mg bid no sooner than 4 days after the start of therapy, to a maximum of 200 mg bid. PO as an aid to smoking cessation (sustained-release) Zyban is identical to Wellbutrin SR and is given in the same dosage regimen as Wellbutrin SR for depression for 7–12 weeks. 110

**Dosage Forms.** Tab 75, 100 mg (Wellbutrin); SR Tab 100, 150 mg (Wellbutrin SR, Zyban).

**Pharmacokinetics.** Elimination half-life is 11–14 hr, but an active hydroxy metabolite has a half-life longer than 24 hr.

**Adverse Reactions.** Frequent adverse effects include insomnia, agitation, headache, and nausea. With dosages of ≤450 mg/day, seizures occur in 0.4% of patients, with a 1-yr cumulative incidence of 0.5%. Bupropion is contraindicated in patients with psychotic disorders (its dopamine agonist effect can increase psychotic symptoms), seizure disorders, anorexia, or bulimia, and in those receiving MAOIs. Bupropion seems to offer better safety in overdose than heterocyclic antidepressants.

**Drug Interactions.** Bupropion can increase levodopa side effects. Phenelzine can increase bupropion’s acute adverse reactions.

**CLOMIPRAMINE**

**Pharmacology.** Clomipramine is a 3-chloro analogue of imipramine that is a potent inhibitor of serotonin reuptake and, unlike other tricyclic antidepressants, antagonizes dopaminergic neurotransmission. It has a specific indication for treat-
ment of obsessive-compulsive disorder (OCD). Few patients experience complete OCD symptom relief; typically, about 40–50% of patients have marked symptom improvement. Although it is an effective antidepressant, its adverse effect profile makes other antidepressants preferred for this indication.111–113 (See Antidepressants Comparison Chart.)

**Adult Dosage.** PO for OCD 25 mg/day initially, increasing to 100 mg/day during the first 2 weeks, and then gradually increasing over several weeks to a maximum of 250 mg/day. **PO as an antidepressant** 100–150 mg/day. Clomipramine can be given safely once daily at bedtime.

**Dosage Forms.** Cap 25, 50, 75 mg.

**Pharmacokinetics.** Antiobsessional effects are first seen at week 4, with maximum effects between weeks 10 and 18. Clomipramine is a highly lipophilic drug with a large first-pass effect and oral bioavailability of 36–62%. The major route of elimination is metabolism by demethylation and then hydroxylation and conjugation, with an elimination half-life of 20–24 hr.

**Adverse Reactions.** Clomipramine’s adverse effect profile is similar to that of amitriptyline (ie, frequent sedation, anticholinergic effects, orthostatic hypotension, tremor, nausea, and sweating), but it has a much higher prevalence of sexual dysfunction and seizures. In controlled studies, 42% of patients experienced ejaculatory failure and 20% were impotent. Frequency of sexual dysfunction increases to >90% of patients when asked directly rather than relying on self-reporting.114 Seizures occur in 0.5% of patients receiving 250 mg/day or less, and 2% of patients experience seizures with dosages above 250 mg/day. Clomipramine is contraindicated in patients who have received MAOIs within the past 14 days. Use with caution in patients with cardiovascular disease (eg, arrhythmias, angina, MI).

**Drug Interactions.** Drug interactions are the same as other tricyclic antidepressants (TCAs). (See Heterocyclic Antidepressants.)

**FLUOXETINE**

**Pharmacology.** Fluoxetine is a bicyclic antidepressant that is a selective and potent inhibitor of presynaptic reuptake of serotonin (an SSRI). It does not affect reuptake of norepinephrine or dopamine and has a relative lack of affinity for muscarinic, histamine, α1- and α2-adrenergic, and serotonin receptors.115

**Administration and Adult Dosage.** (See Antidepressants Comparison Chart.) **PO for depression or OCD** 20 mg/day initially, administered in the morning. Increase dosage no more frequently than q 3–5 weeks. Divide higher dosages, with the last dose given in early afternoon. Although the maximum labeled dosage is 80 mg/day, 20 mg is equal in efficacy for major depression to higher dosages with the benefit of fewer adverse effects.115,116 For depression maintenance, Prozac Weekly 90 mg once/week can be started one week after the last 20 mg/day dose. **PO for bulimia** 60 mg/day in the morning. **PO for premenstrual dysphoric disorder** 20 mg/day; higher dosages appear to have no increased efficacy. Administration for the 14 days before menses can be as effective as continuous use.117

**Special Populations.** Pediatric Dosage. (<18 yr) safety and efficacy not established.
Geriatric Dosage. Reduce initial dosage and rate of dosage increase in the elderly. Single-dose studies suggest no difference in maintenance dosage in the elderly, but data from multiple-dose studies are needed. (See Notes.)

Other Conditions. Reduce initial dosage and rate of dosage increase in patients with hepatic impairment. Dosage adjustment in renal impairment is unnecessary.115

Dosage Forms. Cap 10, 20, 40 mg; SR cap 90 mg (Prozac Weekly); Soln 4 mg/mL; Tab 10 mg.

Patient Instructions. This drug requires at least 2 weeks for a noticeable response in mood and up to 4 weeks for full therapeutic benefit. Take fluoxetine in the morning or early afternoon. Inform your physician of any other medications you are taking.

Pharmacokinetics. Onset and Duration. Onset is delayed 2–4 weeks, which is similar to other antidepressants.

Serum Levels. Not established.

Fate. Oral bioavailability is 95% with all dosage forms. It is 94% bound to plasma proteins, with a $V_d$ of 35 ± 21 L/kg; CI is 0.58 ± 0.41 L/hr/kg, decreasing with repeated administration. The primary active metabolite is norfluoxetine; the metabolic rate is possibly under polygenic control.

$t_1/2$. (Fluoxetine) 1–3 days after a single oral dose, increasing with multiple doses to 4–5 days; (norfluoxetine) 7–15 days. Half-lives do not appear to be altered in the elderly or in patients with renal impairment. Patients with alcohol-induced cirrhosis have fluoxetine half-life increased by 100% and norfluoxetine half-life increased by 60% compared with controls.115,118

Adverse Reactions. Nausea, anxiety, insomnia, nervousness, diarrhea, anorexia, dry mouth, headache, and tremor occur with a frequency greater than 10%. Delayed ejaculation and anorgasmia occurs with fluoxetine and all SSRIs in at least 30–55% of patients.119,120 Unlike TCAs, which typically cause weight gain, fluoxetine dosages over 40 mg/day cause a weight loss of 1–2 kg within the first 6 weeks of treatment.121 Fluoxetine rarely causes sedation except at dosages over 40 mg/day and has no adverse cardiovascular or anticholinergic effects.122 Initial case reports of patients developing new and intense suicidal preoccupation, agitation, and impulsiveness after several weeks of fluoxetine therapy have been adequately evaluated and found not to be directly related to the drug.123

Contraindications. Pregnancy. Concurrent use of an MAOI; 5 weeks must elapse between discontinuation of fluoxetine and starting an MAOI.124

Precautions. Use cautiously in the elderly and in patients with hepatic impairment. Use fluoxetine with caution in depressed patients with psychomotor agitation and anxiety or with anorexia and weight loss.

Drug Interactions. Fluoxetine is a potent inhibitor of CYP2D6, causing decreased metabolism and increased serum levels and adverse effects of many drugs, including most other antidepressants, antipsychotics, β-blockers, and type Ic antiarrhythmics. Fluoxetine’s effect on other P450 isoenzymes has not been well defined.
Parameters to Monitor. Monitor liver function tests periodically during long-term therapy.

Notes. Fluoxetine is a useful alternative to TCAs because of its greater safety in overdose and relative lack of anticholinergic and cardiovascular effects. For severely depressed elderly patients, fluoxetine is less effective than nortriptyline. Fluoxetine and other SSRIs have demonstrated efficacy for OCD and panic disorder. (See Antidepressants Comparison Chart.)

**FLUVOXAMINE**

**Pharmacology.** Fluvoxamine has a selective and potent inhibitory effect on serotonergic presynaptic reuptake, similar to fluoxetine. Although it is also an effective antidepressant, fluvoxamine has been marketed for use in OCD. Fluvoxamine is equal in efficacy to clomipramine in OCD and causes fewer anticholinergic effects and sexual dysfunction but more headache and insomnia. (See Antidepressants Comparison Chart.)

**Adult Dosage.** PO for OCD 50 mg/day initially, with a maintenance dosage of 100–300 mg/day. Give dosages over 100 mg/day in 2 divided doses. The elderly and those with hepatic impairment might require a lower starting dosage and slower dosage titration.

**Pediatric Dosage.** PO for OCD (<8 yr) safety and efficacy not established; (8–17 yr) 25 mg hs initially, increasing in 25 mg/day increments q 4–7 days to a usual maintenance dosage of 50–200 mg/day. Divide dosages >50 mg/day into 2 doses, either equal or with a greater portion given hs.

**Dosage Forms.** Tab 25, 50, 100 mg.

**Pharmacokinetics.** Bioavailability is about 50% and not affected by food. Fluvoxamine is the least protein bound of the SSRIs (77%). On the same dosage, elderly patients have 40% higher serum concentrations than younger patients. Fluvoxamine is metabolized to inactive metabolites. Elimination half-life is about 16 hr in adults and 26 hr in the elderly.

**Adverse Reactions.** Frequent adverse effects include nausea, somnolence or insomnia, dry mouth, and sexual dysfunction.

**Drug Interactions.** Unlike other SSRIs, fluvoxamine is a potent inhibitor of CYP1A2, so increased levels and adverse effects are possible with warfarin, propranolol, metoprolol, caffeine, and theophylline. As with all SSRIs, do not administer fluvoxamine with MAOIs.

**HETEROCYCLIC ANTIDEPRESSANTS**

**Pharmacology.** Heterocyclic antidepressants (tricyclic antidepressants, amoxapine, and maprotiline) have specific effects on neurotransmitters and receptor sensitivity. The primary pharmacologic effect of heterocyclic antidepressants is blockade of presynaptic reuptake of norepinephrine, with subsequent downregulation of adrenergic receptors. Amoxapine, a metabolite of loxapine, retains some postsynaptic dopamine reuptake inhibition. Heterocyclic antidepressants have less effect on serotonergic activity than on other neurotransmitters.
Administration and Adult Dosage. (See Antidepressants Comparison Chart for dosage ranges.) PO for depression initiate dosage at lower limit of range. Administer in divided doses to assess tolerance to side effects and then once-daily hs can be used.\textsuperscript{129,131} Maintenance dosage should be the same as the dosage necessary to treat the acute depressive episode.\textsuperscript{132} IM rarely used (eg, surgical patient NPO for 1–2 days). PO for chronic pain (amitriptyline or imipramine) 10–25 mg/day initially; most patients respond to a dosage of 25–75 mg/day, although dosages up to 200 mg/day have been used.\textsuperscript{133,134} (See Notes.)

Special Populations. Pediatric Dosage. Not recommended <12 yr except for childhood enuresis. PO for enuresis (imipramine) (<12 yr) 25–50 mg/day; (≥12 yr) up to 75 mg/day.\textsuperscript{135} Imipramine maximum dosage in children is 2.5 mg/kg/day; however, use in prepubertal major depression disorder often requires up to 5 mg/kg/day with serum levels over 150 μg/L.\textsuperscript{136}

Geriatric Dosage. (>65 yr) reduce initial dosage by at least 50% of adult dosage and increase the dosage slowly.\textsuperscript{137}

Other Conditions. Reduce initial dosage and rate of titration in patients with cardiovascular or hepatic disease.\textsuperscript{138} During the last trimester of pregnancy, the mean dosage of TCAs required is 1.6 times that of nonpregnant women.\textsuperscript{139}

Dosage Forms. (See Antidepressants Comparison Chart.)

Patient Instructions. (See Antidepressants Class Instructions.) These drugs usually take 2 weeks for a noticeable response in mood and up to 4 weeks for full therapeutic benefit. If you have small children, be sure to keep this medication in a secure place.

Pharmacokinetics. Onset and Duration. Physiologic symptoms of depression (eg, sleep and appetite disturbance, decreased energy) should improve after 1 week, but mood (pessimism, hopelessness, anhedonia) often requires 2–4 weeks for response.

Serum Levels. Nortriptyline has a well-established therapeutic range and a curvilinear relationship of serum levels and response (“therapeutic window”). Other antidepressants show a linear response relationship.\textsuperscript{140} (See Antidepressants Comparison Chart.)

Fate. Bioavailability is variable (30–70%) because of first-pass metabolism. Major metabolites for TCAs are desmethyl (for tertiary amines) and hydroxy compounds; rate is possibly genetically determined and can result in 30-fold variation in steady-state levels in patients given the same dosage.\textsuperscript{141}

\( t_{1/2} \). (Tertiary amine TCAs) 10–25 hr; (secondary amine TCAs) 12–44 hr.\textsuperscript{140}

Adverse Reactions. Sedation, postural hypotension, anticholinergic effects (dry mouth, blurred near vision, constipation, urinary retention, aggravation of narrow-angle glaucoma, and prostatic hypertrophy), weight gain, and cardiac effects (ECG changes and slowed AV conduction) are frequent. Nortriptyline is least likely of the TCAs to cause postural hypotension. (See Antidepressants Comparison Chart for relative differences in frequency of common adverse reactions.) Fine hand tremors, seizures, cardiac arrhythmia, or cholestasis can occur, as can
hypomanic or manic episodes in bipolar patients. Seizures and blood dyscrasias are rare.\textsuperscript{142,143}

**Contraindications.** Cardiac arrhythmias, especially bundle-branch block.

**Precautions.** Use with caution in the elderly, in pregnancy, or in patients with CHF and angina pectoris, epilepsy, glaucoma, prostatic hypertrophy, or renal or liver disease. When discontinuing therapy, taper the heterocyclic antidepressant dosage to prevent cholinergic rebound. Cases of sudden cardiac death have been reported in children with attention deficit disorder who received desipramine in therapeutic or subtherapeutic dosages.\textsuperscript{144} Ingestion of $\geq 1$ g of a heterocyclic antidepressant constitutes a life-threatening medical emergency. Limit the quantities dispensed to depressed patients with suicidal ideation. Maprotiline has an increased frequency of seizures at dosages above 225 mg/day. Amoxapine has a metabolite with dopamine-blocking activity, resulting in possible extrapyramidal effects, tardive dyskinesia, endocrine effects, and neuroleptic malignant syndrome. Neither amoxapine nor maprotiline offers greater efficacy or safety in overdose than TCAs.

**Drug Interactions.** Many drug interactions occur. Use with caution with MAOIs. The antihypertensive effects of guanethidine, clonidine, and closely related drugs might be reduced.

**Parameters to Monitor.** Monitor hepatic and renal function tests periodically during long-term therapy. Obtain ECG in the elderly, children, and those with pre-existing heart disease. With amoxapine, monitor carefully for signs of tardive dyskinesia.

**Notes.** TCAs are commonly used in treating pain associated with diabetic neuropathy and postherpetic neuralgia; amitriptyline, desipramine, and nortriptyline have proven efficacy, but SSRIs are less effective.\textsuperscript{133,134} (See Antidepressants Comparison Chart.)

**MIRTAZAPINE**

**Pharmacology.** Mirtazapine is an antidepressant that antagonizes presynaptic $\alpha_2$-adrenergic auto- and heteroreceptors that are responsible for controlling the release of norepinephrine and serotonin (5-HT). It is also a potent antagonist of postsynaptic 5-HT$_2$ and 5-HT$_3$ receptors. The net outcome of these effects is increased noradrenergic activity and enhanced 5-HT activity, especially at 5-HT$_1A$ receptors. This unique mechanism of action preserves antidepressant efficacy but minimizes many of the adverse effects common to heterocyclic antidepressants and SSRIs. Mirtazapine is effective in moderate and severe major depression.\textsuperscript{145,146} (See Antidepressants Comparison Chart.)

**Administration and Adult Dosage.** PO for depression 15 mg/day at bedtime initially, increasing at 1–2-week intervals to a maximum of 45 mg/day.

**Dosage Forms.** Tab (conventional and rapidly dissolving) 15, 30, 45 mg.

**Pharmacokinetics.** Mirtazapine has an onset of clinical effect in 2–4 weeks, similar to other antidepressants. It has an elimination half-life of 20–40 hr, allowing once-daily administration at bedtime.

**Adverse Reactions.** Sedation, increased appetite, and weight gain are the most frequent side effects. Sedation is most frequent at lower doses (15 mg) and de-
creases in frequency with increasing dosage. Although two cases of agranulocytosis occurred in clinical trials, no specific or additional blood count monitoring is required. Mirtazapine has minimal cardiovascular and anticholinergic effects and essentially lacks adverse GI effects, insomnia, and sexual dysfunction. Do not use mirtazapine within 14 days of an MAOI. Overdose up to 975 mg in combination with a benzodiazepine has caused marked sedation but no difficulty with cardiovascular or respiratory effects.

**MONOAMINE OXIDASE INHIBITORS**

**Pharmacology.** MAOIs are thought to exert their antidepressant action because of alterations in adrenergic and serotonergic receptor sensitivity. The most consistent findings during long-term MAOI therapy include downregulation of β-adrenergic and adenyl cyclase activities. Isoxcarboxazid and phenelzine are hydrazine derivatives; tranylcypromine is a nonhydrazine.

**Administration and Adult Dosage.** PO for depression (isocarboxazid) 20–30 mg/day; (phenelzine) 45–90 mg/day; (tranylcypromine) 30–60 mg/day. Initiate dosage at the lower limit and titrate upward depending on tolerance to side effects. Dosage schedule should remain divided, usually bid or tid. Avoid bedtime administration because MAOIs can delay onset of sleep.

**Special Populations.** Pediatric Dosage. (<16 yr) not recommended.

Geriatric Dosage. Limited information, but decrease initial dosage by 50% because of orthostatic hypotension. Contraindicated in patients older than 60 yr.

Other Conditions. Reduce the initial dosage and rate of upward titration if the patient has taken a heterocyclic antidepressant within 7–10 days.

Dosage Forms. (Isocarboxazid) Tab 10 mg; (phenelzine) Tab 15 mg; (tranylcypromine) Tab 10 mg.

Patient Instructions. (See Antidepressants Class Instructions.) This drug usually takes 2 weeks for noticeable response in mood and up to 4 weeks for full therapeutic benefit to occur. This drug can cause faintness or dizziness, especially after rising suddenly or standing for prolonged periods, or after exertion or alcohol intake. Immediately report nausea, vomiting, sweating, severe occipital headache, and stiff neck, which might be signs of a serious adverse effect. Avoid concurrent use of diet pills and cough and cold remedies and restrict consumption of aged foods high in tyramine. (See Foods That Interact with MAO Inhibitors Chart.)

Pharmacokinetics. Onset and Duration. Onset 2 weeks; maximum improvement occurs after 3–4 weeks.  

Serum Levels. Not used clinically.

Fate. Termination of drug action is dependent on MAO regeneration because the drugs or their active metabolites chemically combine with the MAO enzyme.

Adverse Reactions. Autonomic effects are frequent and not necessarily dose dependent; these include postural hypotension, dry mouth, and constipation. Drowsiness is more frequent with phenelzine, whereas overstimulation and agitation are more likely with tranylcypromine; isocarboxazid is mildly stimulating. Occasionally, delayed ejaculation, edema, skin rash, urinary retention, and blurred vision
occur. MAOIs are much less likely than TCAs to cause weight gain, with tranylcypromine the least likely.\textsuperscript{148}

**Contraindications.** Patients older than 60 yr; patients with confirmed or suspected cerebrovascular defect; cardiovascular disease; pheochromocytoma; history of liver disease or abnormal liver function tests.

**Precautions.** Always consider the possibility of suicide in depressed patients and take adequate precautions. Like other antidepressant drugs, MAOIs can switch bipolar patients to a hypomanic or manic state.

**Drug Interactions.** Postural hypotension can increase with co-administration of antipsychotic, heterocyclic antidepressant, or antihypertensive drugs, and in patients with CHF. Avoid concurrent use with buspirone, heterocyclic antidepressants, meperidine, sympathomimetic drugs, SSRIs, and other MAOIs. A 1–2-week drug-free interval is necessary when switching from an MAOI to a TCA, but a drug-free interval is not necessary when switching from a TCA to an MAOI.\textsuperscript{149} Although uncommon, hypertensive crisis can result from concurrent use of sympathomimetic amines or ingestion of food and drinks high in tyramine.\textsuperscript{150,151} Avoid diets high in tyramine content. (See Foods That Interact with MAO Inhibitors Chart.)

**Parameters to Monitor.** Monitor blood pressure frequently.

**Notes.** MAOIs are excellent alternatives to heterocyclic antidepressants in major depressive disorder, are very effective in panic disorder, and are drugs of choice for atypical depression.\textsuperscript{150,152}

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**FOODS THAT INTERACT WITH MAO INHIBITORS**

Many fermented foods contain tyramine as a byproduct formed by the bacterial breakdown of the amino acid tyrosine; it also can be formed by parahydroxylation of phenylethylamine or dehydroxylation of dihydroxyphenylalanine (DOPA) and dopamine. Tyramine and some other amines found in food can cause hypertensive reactions in patients taking MAO inhibitors. MAO found in the GI tract inactivates tyramine; when drugs prevent this, exogenous tyramine and other monoamines are absorbed and release norepinephrine from sympathetic nerve endings and epinephrine from the adrenal gland. If sufficient quantities of these pressor compounds are released, palpitations, severe headache, and hypertensive crisis can result.

**FOODS THAT CONTAIN TYRAMINE**

<table>
<thead>
<tr>
<th>Foods</th>
<th>Notes</th>
</tr>
</thead>
<tbody>
<tr>
<td>Avocados</td>
<td>Particularly if overripe.</td>
</tr>
<tr>
<td>Bananas</td>
<td>Reactions can occur if eaten in large amounts; tyramine levels are high in peel.</td>
</tr>
<tr>
<td>Bean curd</td>
<td>Fermented bean curd, fermented soya bean, soya bean pastes, soy sauces, and miso soup, prepared from fermented bean curd, contain tyramine in large amounts; miso soup has caused reactions.</td>
</tr>
<tr>
<td>Beer and ale</td>
<td>Major domestic brands do not contain appreciable amounts; some imported brands have had high levels. Nonalcoholic beer might contain tyramine and should be avoided.</td>
</tr>
<tr>
<td>Caviar</td>
<td>Safe if vacuum-packed and eaten fresh or refrigerated only briefly.</td>
</tr>
</tbody>
</table>
## FOODS THAT CONTAIN TYRAMINE

<table>
<thead>
<tr>
<th>Item</th>
<th>Notes</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cheese</td>
<td>Reactions possible with most, except unfermented varieties such as cottage cheese. In others, tyramine concentration is higher near the rind and close to fermentation holes.</td>
</tr>
<tr>
<td>Figs</td>
<td>Particularly if overripe.</td>
</tr>
<tr>
<td>Fish</td>
<td>Safe if fresh; avoid dried products. Caution required in restaurants.Vacuum-packed products are safe if eaten promptly or refrigerated only briefly.</td>
</tr>
<tr>
<td>Liver</td>
<td>Safe if very fresh, but rapidly accumulates tyramine; caution required in restaurants.</td>
</tr>
<tr>
<td>Meat</td>
<td>Safe if known to be fresh; caution required in restaurants.</td>
</tr>
<tr>
<td>Milk products</td>
<td>Milk and yogurt appear to be safe.</td>
</tr>
<tr>
<td>Protein extracts</td>
<td>See also soups; avoid liquid and powdered protein dietary supplements.</td>
</tr>
<tr>
<td>Sausage</td>
<td>Fermented varieties such as bologna, pepperoni, and salami have a high tyramine content.</td>
</tr>
<tr>
<td>Shrimp paste</td>
<td>Contains large amounts of tyramine.</td>
</tr>
<tr>
<td>Soups</td>
<td>Might contain protein extracts and should be avoided.</td>
</tr>
<tr>
<td>Soy sauce</td>
<td>Contains large amounts of tyramine; reactions have occurred with teriyaki.</td>
</tr>
<tr>
<td>Wines</td>
<td>Generally do not contain tyramine, but many reactions have been reported with Chianti, champagne, and other wines.</td>
</tr>
<tr>
<td>Yeast extracts</td>
<td>Dietary supplements (eg, Marmite) contain large amounts; yeast in baked goods, is safe.</td>
</tr>
</tbody>
</table>

## FOODS THAT DO NOT CONTAIN TYRAMINE

<table>
<thead>
<tr>
<th>Item</th>
<th>Notes</th>
</tr>
</thead>
<tbody>
<tr>
<td>Caffeine</td>
<td>A weak pressor agent; large amounts can cause reactions.</td>
</tr>
<tr>
<td>Chocolate</td>
<td>Contains phenylethylamine, a pressor agent that can cause reactions in large amounts.</td>
</tr>
<tr>
<td>Fava beans</td>
<td>(Broad beans, &quot;Italian&quot; green beans) Contain dopamine, a pressor amine, particularly when overripe.</td>
</tr>
<tr>
<td>Ginseng</td>
<td>Some preparations have caused headache, tremulousness, and manic-like symptoms.</td>
</tr>
<tr>
<td>Liqueurs</td>
<td>Reactions reported with some (eg, Chartreuse, Drambuie); cause unknown.</td>
</tr>
<tr>
<td>New Zealand prickly spinach</td>
<td>Single case report; patient ate large amounts.</td>
</tr>
<tr>
<td>Whiskey</td>
<td>Reactions have occurred; cause unknown.</td>
</tr>
</tbody>
</table>


Pharmacology. Nefazodone is a postsynaptic serotonin 5-HT₂A antagonist and presynaptic serotonin reuptake inhibitor. These two serotonergic effects make it different from SSRIs and TCAs.153–156 (See Antidepressants Comparison Chart.)

Administration and Adult Dosage. PO for depression 100 mg bid initially (50 mg bid in the elderly), increasing q 4–7 days to the effective dosage range of 150–300 mg bid. After initial dosage titration, once-daily bedtime administration is preferred to minimize daytime sedation.157

Dosage Forms. Tab 50, 100, 150, 200, 250 mg.

Pharmacokinetics. Nefazodone has an oral bioavailability of about 20%. Single-dose studies in the elderly have shown a 100% larger AUC; with multiple doses, the AUC differences decreased to 10–20% above those in younger populations. It is >99% protein bound and extensively metabolized, with a dose-dependent elimination half-life of about 1–2.3 hr in young patients, modestly prolonged in the elderly, and 2–3 times longer in hepatic disease. The major active metabolite, hydroxynefazodone, has a half-life of 1.2–1.6 hr in young and elderly patients, increasing to 2–4 hr with hepatic disease. Renal impairment does not markedly affect nefazodone pharmacokinetics.

Adverse Reactions. Although chemically similar to trazodone, it causes less sedation and orthostatic hypotension, and its lower α-adrenergic blockade makes priapism much less likely (no cases reported). Frequent adverse effects include sedation, dry mouth, nausea, and dizziness. Unlike SSRIs, nefazodone’s effects on sexual function, agitation, tremor, insomnia, and weight are no different from placebo.

Drug Interactions. Nefazodone is a potent inhibitor of the CYP3A4 isoenzyme and a weak inhibitor of the CYP2D6 isoenzyme. Drug interactions of particular concern include the triazolobenzodiazepines (ie, alprazolam, triazolam, midazolam). A 1- to 2-week washout period is recommended when converting a patient to or from a MAOI and nefazodone.

Paroxetine is a highly selective and potent inhibitor of serotonin reuptake (an SSRI) similar to fluoxetine.126,158–164 (See Antidepressants Comparison Chart.)

Administration and Adult Dosage. PO for depression 20 mg/day; a few patients require 30–50 mg/day for full efficacy. PO for social anxiety disorder and panic disorder 10 mg/day initially; usual maintenance dosage is 20–60 mg/day. PO for OCD 20 mg/day initially; maintenance dosage is 40 mg/day to a maximum of 60 mg/day, preferably as a single dose in the morning or evening. The starting dosage for all uses in elderly patients and those with marked renal or hepatic impairment is 10 mg/day. For the elderly or those with severe renal or hepatic impairment, the maximum dosage is 40 mg/day.

Dosage Forms. Tab 10, 20, 30, 40 mg; SR Tab 12.5, 25 mg; Susp 2 mg/mL.
Pharmacokinetics. Paroxetine is completely orally bioavailable; protein binding is 93–95%. Unlike fluoxetine, paroxetine is metabolized to inactive metabolites and has an elimination half-life of 24 hr.

Adverse Reactions. Paroxetine causes the typical SSRI adverse effects of nausea, sexual dysfunction, and headache but is more likely to cause sedation than insomnia and can cause more delay of orgasm or ejaculation and more impotence than other SSRIs. Like the other SSRIs, it is much safer in overdose than TCAs.

Drug Interactions. Paroxetine is a potent inhibitor of CYP2D6, so most other antidepressants, antipsychotics, β-blockers, and type Ic antiarrhythmics can have increased serum levels and adverse effects when paroxetine is combined with these drugs. Do not use paroxetine within 14 days of using an MAOI.

Pharmacology. Reboxetine is the first in a new class of selective norepinephrine reuptake inhibitors with no affinity for serotonin or dopamine reuptake sites. It has negligible affinity for muscarinic, histaminic, or adrenergic receptors. This noradrenergic mechanism for antidepressant efficacy is similar to TCAs such as desipramine without the potential for appreciable adverse anticholinergic, cardiovascular, and sedative effects. It has efficacy for major depression equal to fluoxetine and desipramine.

Administration and Adult Dosage. PO for depression 8–10 mg/day given bid, 4–6 mg/day given bid in the elderly.

Dosage Forms. Tab 4 mg (investigational).

Pharmacokinetics. Reboxetine is rapidly absorbed. Metabolism occurs through three oxidative pathways: hydroxylation, dealkylation, and oxidation. The CYP450 isoenzymes responsible for metabolism have not been identified, and the degree of activity of the metabolites is unknown. Reboxetine has no inhibitory effect on CYP450 isoenzymes. Elimination half-life is 13 hr.

Adverse Reactions. The most common adverse effects include dry mouth, constipation, increased sweating, insomnia, and urinary hesitancy, which are greater than placebo, but less frequent than imipramine. These “anticholinergic-like” effects are believed to result from increased norepinephrine levels. Side effects commonly associated with serotonin reuptake inhibitors such as nausea, anxiety or agitation, and daytime somnolence were no more common with reboxetine than with placebo.

No information is available regarding reboxetine overdose in humans.

Pharmacology. Sertraline is an SSRI similar to fluoxetine, which indirectly results in a downregulation of β-adrenergic receptors. It has no clinically important effect on noradrenergic or histamine receptors and no effect on MAO. It lacks stimulant, cardiovascular, anticholinergic, and convulsant effects. Sertraline has antidepressant effects equal to TCAs and fluoxetine and might have anorectic effects and efficacy in OCD. (See Antidepressants Comparison Chart.)

Administration and Adult Dosage. PO for depression, panic disorder, OCD, and posttraumatic stress disorder 50 mg/day initially, increasing if necessary at weekly intervals to a maximum of 200 mg/day in a single dose in the morning or evening.
Dosage Forms. Tab 25, 50, 100 mg; Soln 20 mg/mL.

Pharmacokinetics. Sertraline has an oral bioavailability of 36%, and, when it is taken with food, peak serum concentrations and bioavailability increase by 30–40%. Peak serum concentrations are reached in 6–8 hr. Sertraline concentrations in breast milk are the lowest of the SSRIs and produce minimal serum levels in the breast-fed infant. Its primary metabolite is N-desmethylsertraline, which has 5–10 times less activity than sertraline as an SSRI and has no demonstrated antidepressant activity. CI is decreased by up to 40% in the elderly. Steady-state half-life is 27 hr.

Adverse Reactions. Frequent adverse effects include nausea, diarrhea, ejaculatory delay, tremor, and increased sweating. It causes less agitation, anxiety, and insomnia than fluoxetine and is a less potent inhibitor of the CYP2D6 isoenzyme at a dosage of 50 mg/day. Use with caution in patients with renal or hepatic impairment and do not use it within 14 days of using an MAOI. SIADH has been reported.

Pharmacology. Venlafaxine is a potent reuptake inhibitor of serotonin and noradrenaline, like many TCAs, but lacks effects on muscarinic, α-adrenergic, or histamine receptors. (See Antidepressants Comparison Chart.)

Administration and Adult Dosage. PO for depression (immediate-release) 75 mg bid or tid initially, increasing q 4–7 days to an effective antidepressant dosage of 225–375 mg/day in 2 or 3 divided doses; (sustained-release) 75 mg once daily initially, increasing in increments of up to 75 mg/day at intervals of 4 or more days to a maximum of 225 mg/day. The sustained-release preparation does not reduce side effects but allows once-daily administration. PO for generalized anxiety disorder 75–225 mg/day in 2–3 divided doses. Patients with renal impairment or on hemodialysis require a 25–50% dosage reduction.

Dosage Forms. Tab 25, 37.5, 50, 75, 100 mg; SR Cap 37.5, 75, 150 mg (Effexor XR).

Pharmacokinetics. Venlafaxine is well absorbed orally; food has no effect on absorption. Serum concentrations in elderly patients are no different from those in younger patients. Unlike SSRIs, venlafaxine has minimal protein binding (27–30%). It undergoes extensive hepatic metabolism. Venlafaxine has an elimination half-life of 5 hr, and one major active metabolite has an 11-hr half-life. Venlafaxine exhibits linear pharmacokinetics over the recommended dosage range, and steady state is reached in 3 days.

Adverse Reactions. Frequent adverse effects include expected serotonin-related effects (eg, nausea, headache, insomnia or somnolence, and sexual dysfunction). At higher dosages (375 mg/day), venlafaxine is unique in causing a consistent but mild elevation in diastolic blood pressure (6 mm Hg). Regular blood pressure monitoring is required for all patients.

Drug Interactions. Venlafaxine is not a potent inhibitor of the cytochrome P450 enzyme system, making it different from most of the SSRIs. Avoid it in patients who have received an MAOI within the past 14 days.
## ANTIDEPRESSANTS COMPARISON CHART

<table>
<thead>
<tr>
<th>CLASS AND DRUG</th>
<th>DOSAGE FORMS</th>
<th>USUAL DAILY ADULT DOSAGE RANGE (MG)</th>
<th>THERAPEUTIC SERUM LEVELS (µG/L)</th>
<th>RELATIVE FREQUENCY OF SIDE EFFECTS</th>
<th>SEDATION</th>
<th>ANTICHOLINERGIC</th>
<th>ORTHOSTATIC HYPOTENSION</th>
</tr>
</thead>
<tbody>
<tr>
<td>α₂-ADRENERGIC BLOCKERS</td>
<td></td>
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</tr>
<tr>
<td>Mirtazapine</td>
<td>Tab (conventional and rapidly dissolving) 15, 30, 45 mg.</td>
<td>15–45</td>
<td>b</td>
<td>Moderate</td>
<td>None</td>
<td>None</td>
<td></td>
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<tr>
<td>Remeron tab</td>
<td></td>
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<tr>
<td>CHLOROPROPIOPHENONES</td>
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<tr>
<td>Bupropion</td>
<td>Tab 75, 100 mg SR Tab 100, 150 mg.</td>
<td>300–450</td>
<td>b</td>
<td>None</td>
<td>None</td>
<td>None</td>
<td></td>
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<tr>
<td>Wellbutrin</td>
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<tr>
<td>Zyban</td>
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<tr>
<td>DIBENZOXAZEPINES</td>
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</tr>
<tr>
<td>Amoxapine</td>
<td>Tab 25, 50, 100, 150 mg.</td>
<td>300–600</td>
<td>b</td>
<td>Low</td>
<td>Low</td>
<td>Low</td>
<td></td>
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<tr>
<td>Asendin</td>
<td></td>
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<tr>
<td>Various</td>
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<td></td>
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</tr>
<tr>
<td>MORPHOLINES</td>
<td></td>
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</tr>
<tr>
<td>Reboxetine</td>
<td>Tab 15 mg.</td>
<td>45–90</td>
<td>b</td>
<td>Moderate</td>
<td>Low</td>
<td>Very High</td>
<td></td>
</tr>
<tr>
<td>Vestra</td>
<td></td>
<td></td>
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</tr>
<tr>
<td>Tranproxyline</td>
<td>Tab 10 mg.</td>
<td>30–60</td>
<td>b</td>
<td>Low</td>
<td>Low</td>
<td>Very High</td>
<td></td>
</tr>
<tr>
<td>Nardil</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Parnate</td>
<td></td>
<td></td>
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</tr>
</tbody>
</table>

(continued)
<table>
<thead>
<tr>
<th>CLASS AND DRUG</th>
<th>DOSAGE FORMS</th>
<th>USUAL DAILY ADULT DOSAGE RANGE (MG)</th>
<th>THERAPEUTIC SERUM LEVELS (µG/L)</th>
<th>RELATIVE FREQUENCY OF SIDE EFFECTS</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>SELECTIVE SEROTONIN REUPTAKE INHIBITORS (SSRIs)</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Citalopram</td>
<td>Tab 20, 40 mg&lt;br&gt;Soln 2 mg/mL.</td>
<td>20–60</td>
<td>b</td>
<td>Very Low</td>
</tr>
<tr>
<td>Celexa</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Fluoxetine</td>
<td>Cap, Tab 10, 20, 40 mg&lt;br&gt;SR Cap 90 mg&lt;br&gt;Soln 4 mg/mL&lt;br&gt;Tab 10 mg.</td>
<td>10–80</td>
<td>b</td>
<td>None</td>
</tr>
<tr>
<td>Prozac</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Fluvoxamine</td>
<td>Tab 25, 50, 100 mg.</td>
<td>100–300&lt;sup&gt;f&lt;/sup&gt;</td>
<td>b</td>
<td>None</td>
</tr>
<tr>
<td>Luvox</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Paroxetine</td>
<td>Tab 10, 20, 30, 40 mg&lt;br&gt;SR Tab 12.5, 25 mg&lt;br&gt;Susp 2 mg/mL.</td>
<td>20–50</td>
<td>b</td>
<td>Low</td>
</tr>
<tr>
<td>Paxil</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Sertraline</td>
<td>Tab 25, 50, 100 mg&lt;br&gt;Soln 20 mg/mL.</td>
<td>50–200</td>
<td>b</td>
<td>None</td>
</tr>
<tr>
<td>Zoloft</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>SEROTONIN NOREPINEPHRINE REUPTAKE INHIBITORS (SNRIs)</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Venlafaxine</td>
<td>Tab 25, 37.5, 50, 75, 100 mg&lt;br&gt;Effexor&lt;br&gt;Effexor XR&lt;br&gt;SR Cap 37.5, 75, 150 mg.</td>
<td>225–375</td>
<td>b</td>
<td>Very Low</td>
</tr>
<tr>
<td><strong>TETRACYCLICS</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Maprotiline</td>
<td>Tab 25, 50, 75 mg.</td>
<td>150–225</td>
<td>200–300&lt;sup&gt;f&lt;/sup&gt;</td>
<td>Moderate</td>
</tr>
<tr>
<td>Ludiomil</td>
<td>Various</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Various</td>
<td></td>
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<td></td>
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</tr>
</tbody>
</table>
### Antidepressants Comparison Chart (continued)

<table>
<thead>
<tr>
<th>Class and Drug</th>
<th>Dosage Forms</th>
<th>Usual Daily Adult Dosage Range (mg)</th>
<th>Therapeutic Serum Levels (µg/L)</th>
<th>Relative Frequency of Side Effects</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>Sedation</td>
</tr>
<tr>
<td><strong>Triazolopyridines</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Trazodone</td>
<td>Tab 50, 100, 150, 300 mg.</td>
<td>50–100 (hypnotic) 200–400 (antidepressant)</td>
<td>b</td>
<td>High</td>
</tr>
<tr>
<td>Desyrel</td>
<td>Various</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Nefazodone</td>
<td>Tab 50, 100, 150, 200, 250 mg.</td>
<td>300–600</td>
<td>b</td>
<td>Moderate</td>
</tr>
<tr>
<td>Serzone</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Tricyclics (TCAs)</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Amitriptyline</td>
<td>Tab 10, 25, 50, 75, 100, 150 mg</td>
<td>150–300</td>
<td>75–175‡</td>
<td>High</td>
</tr>
<tr>
<td>Elavil</td>
<td>Inj 10 mg/mL.</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Clomipramine</td>
<td>Cap 25, 50, 75 mg.</td>
<td>100–250′</td>
<td>b</td>
<td>High</td>
</tr>
<tr>
<td>Anafranil</td>
<td>100–150′</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Various</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Desipramine</td>
<td>Tab 10, 25, 50, 75, 100, 150 mg</td>
<td>150–300</td>
<td>100–160</td>
<td>Low</td>
</tr>
<tr>
<td>Norpramin</td>
<td>Various</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Doxepin</td>
<td>Cap 10, 25, 50, 75, 100, 150 mg</td>
<td>150–300</td>
<td>110–250′</td>
<td>High</td>
</tr>
<tr>
<td>Adapin</td>
<td>Sinequan Soln 10 mg/mL.</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Various</td>
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<td></td>
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</tr>
</tbody>
</table>

(continued)
<table>
<thead>
<tr>
<th>CLASS AND DRUG</th>
<th>DOSAGE FORMS</th>
<th>USUAL DAILY ADULT DOSAGE RANGE (MG)</th>
<th>THERAPEUTIC SERUM LEVELS (MG/L)</th>
<th>RELATIVE FREQUENCY OF SIDE EFFECTS</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>Sedation</td>
</tr>
<tr>
<td>Imipramine</td>
<td>Tab 10, 25, 50 mg</td>
<td>150–300</td>
<td>&gt;200*</td>
<td>Moderate</td>
</tr>
<tr>
<td></td>
<td>Cap (as pamoate) 75, 100, 125, 150 mg</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Nortriptyline</td>
<td>Cap 10, 25, 50, 75 mg</td>
<td>100–200</td>
<td>50–150</td>
<td>Moderate</td>
</tr>
<tr>
<td>Aventyl</td>
<td>Soln 2 mg/mL.</td>
<td>Step 5, 10 mg.</td>
<td>30–60</td>
<td>70–260*</td>
</tr>
<tr>
<td>Parnetic</td>
<td>Various</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Protriptyline</td>
<td>Tab 5, 10 mg.</td>
<td>30–60</td>
<td>70–260*</td>
<td>Very Low</td>
</tr>
<tr>
<td>Aventyl</td>
<td>Various</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Trimepramine</td>
<td>Cap 25, 50, 100 mg</td>
<td>150–300</td>
<td>b</td>
<td>Moderate</td>
</tr>
<tr>
<td>Surmontil</td>
<td>Various</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
**Antipsychotic Drugs**

**Class Instructions.** Antipsychotics. This drug can cause drowsiness. Until the extent of this effect is known, use caution when driving, operating machinery, or performing other tasks requiring mental alertness. Avoid excessive concurrent use of alcohol or other drugs that cause drowsiness.

**Missed Doses.** If you miss a dose, take it as soon as you remember. If it is almost time for your next dose, skip it and resume your normal schedule. Do not double doses.

**Pharmacology.** Antipsychotic efficacy is most likely related to blockade of postsynaptic dopaminergic receptors in the mesolimbic and prefrontal cortices of the brain, although other neurotransmitter systems also are involved.\(^{181}\)

**Administration and Adult Dosage.** (See Antipsychotic Drugs Comparison Chart for oral dosage ranges.) Initiate therapy with divided doses until therapeutic dosage is found; then, for most patients, once-daily hs administration is preferred. For maintenance, decrease acute dosage by 25% q 3 months, with a target maintenance dosage being 50–67% of the acute treatment dosage.\(^{182}\) Recent concern has focused on the need to establish a minimum effective dosage for antipsychotic drugs, and treatment regimens at the low end of the dosage range are preferred. Oral dosages of high-potency antipsychotics (eg, fluphenazine, haloperidol) in the range of 5–20 mg/day are better tolerated and equal in efficacy to dosages >20 mg/day.\(^{183}\) Most patients can be given a maintenance dosage of 50% the acute dosage by the end of 1 yr, although 10–15% of chronically ill patients require a maintenance dosage >15 mg/day of haloperidol or its equivalent.\(^{184,185}\) For manic episodes, no additional benefit is achieved with dosages >10 mg/day of haloperidol.\(^{186}\) Mesoridazine and thioridazine are indicated only in patients who fail with other drugs because of inefficacy or intolerable side effects.

**Special Populations.** *Pediatric Dosage.* As with adults, dosage is determined primarily by titration to individual response. No precise dosage range exists, but in general the initial dosage is lower and increased more gradually in children.

**Geriatric Dosage.** Initial dosage is 20–25% of the dosage used in younger adults. Typical starting dosages in the elderly are haloperidol 0.5–2 mg/day. Dosage adjustments also must be done more slowly than in younger adults.\(^{187}\)

**Other Conditions.** Dosages in the lower range are sufficient for most elderly patients, and the rate of dosage titration is slower.

**Dosage Forms.** (See Antipsychotic Drugs Comparison Chart.)

**Patient Instructions.** (See Antipsychotics Class Instructions.) These drugs usually take several weeks for clinical response and up to 8 weeks for full therapeutic response.

**Pharmacokinetics. Onset and Duration.** Onset of antipsychotic activity is variable, with noticeable response requiring days to weeks.

**Serum Levels.** Correlation of serum levels with clinical response is not consistently established. The best evidence exists for haloperidol, with serum concen-
trations of 5–15 μg/L (13–40 nmol/L) correlating well with therapeutic effects in adult psychotic patients, and an increasing risk of adverse effects and decreased efficacy when steady-state concentrations exceed 15 μg/L.188,189

**Fate.** Haloperidol is well absorbed; peak serum levels are achieved 2–6 hr after liquid or tablets and within 30 min after IM. Oral bioavailability of haloperidol is 60–70%. Haloperidol is extensively metabolized, with one active hydroxy metabolite. Chlorpromazine and other phenothiazines are well absorbed but undergo extensive and variable presystemic metabolism in the gut wall and liver; more than 20 chlorpromazine metabolites with different activities have been identified in human plasma. SR formulations result in a greater first-pass effect.

\[ t_{1/2} \]

Serum half-lives have no clinical correlation with biologic half-lives for antipsychotic drugs. Chlorpromazine serum half-life is 30 hr, thioridazine 4–10 hr, thiothixene 34 hr, and haloperidol 12–24 hr. Of more clinical importance is that steady-state CNS levels and tissue saturation allow once-daily administration of all antipsychotic drugs.141

**Adverse Reactions.** (See Antipsychotic Drugs Comparison Chart for relative frequency of common adverse reactions.) Frequently, sedation, extrapyramidal effects (eg, parkinsonism, dystonic reactions, akathisia), tardive dyskinesia, anticholinergic effects (eg, dry mouth, blurred vision, constipation, urinary retention), photosensitivity, and postural hypotension occur. Occasionally, weight gain, amenorrhea, galactorrhea, ejaculatory disturbance, neuroleptic malignant syndrome, agranulocytosis, skin rash, cholestatic jaundice, and skin or eye pigmentation occur. Rarely, seizures, thermoregulatory impairment, and slowed AV conduction occur. Mesoridazine and thioridazine can prolong QTc interval, leading to torsades de pointes and sudden death. Low-potency drugs are more likely to cause sedation, anticholinergic effects, and orthostatic hypotension, whereas high-potency drugs cause more extrapyramidal effects. Tardive dyskinesia is a long-term adverse effect, untreatable, and sometimes irreversible. Tardive dyskinesia occurs at a 4% yearly incidence for at least the first 5–6 yr of treatment. Neuroleptic malignant syndrome (ie, fever, extrapyramidal rigidity, autonomic instability, alterations in consciousness) occurs more frequently with high-potency antipsychotics, with a prevalence of 1.4% and a fatality rate of 4%.183,190,191

**Contraindications.** Coma; circulatory collapse or severe hypotension; bone marrow depression; history of blood dyscrasia. (Mesoridazine and thioridazine) concurrent use with drugs that prolong QTc interval; baseline QTc >450 msec.

**Precautions.** Use cautiously in patients with myasthenia gravis, Parkinson’s disease, seizure disorders, or hepatic disease.

**Drug Interactions.** Barbiturates can enhance phenothiazine metabolism; carbamazepine can enhance haloperidol metabolism. Phenothiazines can decrease efficacy of guanethidine or guanadrel or have additive hypotensive effects with hypotensive drugs. Phenothiazines can inhibit the antiparkinson activity of levodopa. Haloperidol can increase the CNS toxicity of lithium. Combined use of haloperidol and methyldopa can result in dementia.

**Parameters to Monitor.** (Mesoridazine and thioridazine) obtain baseline and periodic ECGs and serum potassium.
Notes. (See also Prochlorperazine Salts in the Antiemetics section for antiemetic uses.)

**CLOZAPINE**

**Pharmacology.** Clozapine is an atypical antipsychotic drug that is chemically similar to loxapine and has unique pharmacologic effects and indications, as well as very serious adverse effects. Whereas typical antipsychotic drugs exert their effects primarily with a blockade of dopamine-D₂ receptors, clozapine affects several dopamine and serotonin receptors. Its high serotonin-5HT₂ to dopamine-D₂ ratio is the likely explanation for its unique efficacy. Compared with traditional antipsychotic drugs, clozapine is more effective for negative symptoms of schizophrenia, is more effective in treatment-resistant patients, and rarely causes extrapyramidal effects.¹⁹²⁻¹⁹⁵

**Adult Dosage.** PO 100–200 mg tid is effective for most patients, but some might require up to 900 mg/day. A therapeutic trial of 12–24 weeks is required for the full therapeutic effect to become apparent.

**Dosage Forms.** Tab 25, 100 mg.

**Pharmacokinetics.** Clozapine is nearly completely absorbed after oral administration, with about 30% oral bioavailability because of extensive first-pass metabolism. Clozapine is 95% bound to plasma proteins; with multiple doses, its elimination half-life is 12 hr.¹⁹⁶

**Adverse Reactions.** Frequent adverse effects include sedation, orthostatic hypotension, anticholinergic effects, fever, and excessive salivation. Seizures are dose related, with a frequency up to 5% in the therapeutic dosage range and a 1-yr cumulative incidence of 10%. Agranulocytosis is the major adverse effect of concern, occurring in 0.8% of patients after 1 yr.¹⁹⁷ Most cases of agranulocytosis occur within the first 3 months of therapy. Substantial weight gain has been reported in most patients receiving clozapine.¹⁹⁸ (See Antipsychotic Drugs Comparison Chart.)

**Parameters to Monitor.** Patients must have a baseline WBC count and differential before initiating therapy, mandatory weekly WBC monitoring for the first 6 months, and then q 2 weeks throughout treatment and for 4 weeks after discontinuation.

**HALOPERIDOL DECANOATE**

**Pharmacology.** Haloperidol decanoate (HD) is the preferred long-acting depot antipsychotic drug. Depot antipsychotics are indicated only for patients who demonstrate good response but are consistently drug-noncompliant with resultant frequent psychotic relapses. Depot antipsychotics provide fewer relapses and hospitalizations, stable serum drug levels, and side effects equal to oral antipsychotic drugs. HD can be given q 4 weeks; fluphenazine decanoate (FD) is similar in efficacy and adverse effects, but it must be administered q 2 weeks. Do not use HD or FD to treat acute psychotic symptoms; rather, use the drug only after a patient has been stabilized on an oral antipsychotic drug.¹⁹⁹⁻²⁰³
Adult Dosage. IM do not exceed an initial HD dosage of 100 mg, with a target monthly dosage 20 times the oral haloperidol daily dosage. An IM loading dose technique has been described that gives 20 times the daily oral dosage, using 100–200 mg of depot q 3–7 days to reach the calculated amount, with a maximum of 450 mg. In geriatric or hepatically impaired patients, use a monthly HD dose of 15 times the oral haloperidol dosage. Experience with HD doses greater than 500 mg is limited; divide injections >5 mL into 2 equal portions given at 2 sites. Oral haloperidol supplementation might be necessary between monthly injections to treat re-emergence of psychotic symptoms until steady-state concentrations are reached.

Dosage Forms. Inj 50, 100 mg/mL.

Pharmacokinetics. After IM administration of HD, esterases cleave the decanoate chain to release the active drug. Peak serum concentrations of haloperidol occur in 3–9 days, with an apparent half-life of 3 weeks; steady-state levels are reached after 12–16 weeks.

Adverse Reactions. There is no evidence that HD causes adverse effects with a frequency different from that of oral haloperidol.

Olanzapine

Pharmacology. Olanzapine is an atypical antipsychotic agent that is a potent serotonin-5HT2 and dopamine-D2 antagonist. It also has anticholinergic and histamine H1-receptor antagonistic effects that might account for some of its side effects.\textsuperscript{204,205} (See Antipsychotic Drugs Comparison Chart.)

Adult Dosage. PO for psychotic disorders 5–10 mg/day initially (5 mg/day in patients >65 yr, debilitated patients, or those with a predisposition to hypotensive reactions). Increase in 5 mg/day increments at ≥7-day intervals. Usual maintenance dosage 10–15 mg/day, to a maximum of 20 mg/day, although dosages >10 mg/day are generally no more effective than 10 mg/day. IM for acute psychosis 2.5–10 mg/dose has been used investigationally.

Dosage Forms. Tab (conventional) 2.5, 5, 7.5, 10, 15 mg; (rapidly dissolving) 5, 10, 15, 20 mg (Zyprexa Zydis); Inj (investigational).

Pharmacokinetics. Olanzapine is well absorbed orally; food has no effect, but bioavailability is about 60% because of a first-pass effect. It is 93% bound to plasma proteins and has a \( V_d \) of about 1000 L. The drug is hepatically metabolized, probably by CYP1A2 and CYP2D6. Only 7% is excreted unchanged in urine. Its half-life is 30 hr.

Adverse Reactions. Frequent adverse effects include drowsiness, agitation, nervousness, orthostatic hypotension, dizziness, tachycardia, headache, rhinitis, constipation, akathisia, and weight gain. As with other atypical antipsychotic drugs, weight gain is the most troublesome long-term adverse effect, often affecting compliance.

Drug Interactions. Inducers of CYP2D6 may decrease olanzapine serum levels. Olanzapine does not appear to affect cytochrome P450 enzymes.
Pimozide is indicated for the treatment of Tourette’s disorder. Although structurally different from other antipsychotic drugs, pimozide shares their ability to block dopaminergic receptors. Its lack of effect on norepinephrine receptors led to the hope that pimozide would have a more favorable adverse effect profile than other antipsychotic drugs. Haloperidol is the drug of choice for Tourette’s disorder.

Adult Dosage. PO 1–2 mg/day in divided doses initially, with dosage increased every other day up to a maximum of 20 mg/day. Most patients who respond require ≤10 mg/day. Periodically decrease the dosage and attempt to withdraw treatment.

Pediatric Dosage. PO 0.05 mg/kg/day initially (preferably hs), increasing q 3 days to a maximum of 0.2 mg/kg or 10 mg daily.

Dosage Forms. Tab 2 mg.

Pharmacokinetics. Pimozide is about 50% absorbed orally. It undergoes extensive first-pass metabolism in the liver to two metabolites with unknown activity. The elimination half-life averages 55 hr.

Adverse Reactions. The relative frequencies of adverse effects of pimozide and haloperidol are similar, and pimozide remains an alternative to haloperidol for treating Tourette’s disorder.

Risperidone is a potent serotonin-5-HT₂ antagonist with dopamine-D₂ antagonism. Whereas typical antipsychotics are dopamine antagonists, the additional serotonin antagonism increases efficacy for negative symptoms of schizophrenia and reduces the likelihood of extrapyramidal symptoms. Initial evidence also suggests that risperidone is more effective than traditional antipsychotic drugs for treatment-resistant schizophrenic patients. (See Antipsychotic Drugs Comparison Chart.)

Adult Dosage. PO 1 mg bid initially (0.5 mg bid in the elderly or patients with severe renal or hepatic impairment), increasing q 2–4 days to the usual effective dosage of 4–6 mg/day in 1 or 2 doses. Occasionally, dosages above 6 mg/day might be necessary, but adverse effects increase and efficacy can be less. The solution can be mixed with water, coffee, orange juice, or low-fat milk; do not mix with cola or tea.

Dosage Forms. Tab 0.25, 0.5, 1, 2, 3, 4 mg; Soln 1 mg/mL.

Pharmacokinetics. Risperidone is well absorbed orally. The free fraction of risperidone in serum increases in hepatic disease, necessitating lower dosages. It is metabolized by CYP2D6 to an active metabolite. Risperidone’s elimination half-life is 3 hr; its active metabolite has a half-life of 24 hr. The half-lives of one or both are prolonged in patients with renal disease.

Adverse Reactions. Frequent dose-related adverse effects are extrapyramidal effects, orthostatic hypotension, headache, rhinitis, and insomnia.
Drug Interactions. Inhibitors of CYP2D6 can increase risperidone levels and have adverse effects.

**ZIPRASIDONE HYDROCHLORIDE**

**Pharmacology.** Ziprasidone is an atypical antipsychotic drug with a very high ratio of 5-HT$_2$A to dopamine-2 blockade, suggesting a very low risk of extrapyramidal effects. In addition, it is a 5-HT$_1$A agonist like buspirone, and inhibits reuptake of both serotonin and norepinephrine like antidepressants. The clinical value of the latter two effects are not established.\textsuperscript{213}

**Adult Dosage.** PO as an antipsychotic 20 mg bid with food initially, increasing as necessary at intervals of at least 2 days to a maximum of 80 mg bid. Maintenance dosage may be as low as 40 mg/day. IM ziprasidone for acute agitation in a psychotic patient 10–20 mg, may repeat in 2–4 hr.\textsuperscript{213}

**Dosage Forms.** Cap 20, 40, 60, 80 mg; Inj investigational.

**Patient Instructions.** (See Antipsychotics Class Instructions.) Take this medication with food.

**Pharmacokinetics.** Oral bioavailability is 60% when taken with food. With oral twice daily administration, peak blood levels occur at 6–8 hr. Ziprasidone is metabolized by aldehyde oxidase and to a lesser extent by CYP3A4 to inactive metabolites. The elimination half-life is 5–10 hr (range 3–18) for oral ziprasidone and 3 hr for IM ziprasidone. The pharmacokinetics are unaffected by sex, age, or moderate renal or hepatic disease.

**Adverse Reactions.** Extrapyramidal effects are minimal, but comparative data with other atypical antipsychotic drugs are not available. A major potential advantage of ziprasidone is that it is the least likely atypical antipsychotic drug to cause weight gain.\textsuperscript{214} Compared to placebo, the only side effect greater with ziprasidone is sedation. Ziprasidone increases the QT$_c$ interval by up to 14 msec. Ziprasidone should be avoided in patients with pre-existing QT$_c$ prolongation, after acute MI, in severe CHF, and in patients taking other drugs that prolong the QT$_c$ interval. The drug should be discontinued if the QT$_c$ interval is persistently >500 msec.
### ANTIPSYCHOTIC DRUGS COMPARISON CHART

<table>
<thead>
<tr>
<th>DRUG AND CLASS</th>
<th>DOSAGE FORMS</th>
<th>ADULT ORAL DOSAGE RANGE (MG/DAY)</th>
<th>ORAL EQUIVALENT ANTIPSYCHOTIC DOSE (MG)</th>
<th>USUAL SINGLE IM DOSE (MG)</th>
<th>RELATIVE FREQUENCY OF SIDE EFFECTS</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>LOW POTENCY</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>Sedation</td>
</tr>
<tr>
<td>Chlorpromazine</td>
<td>Soln 30, 100 mg/mL, Syrup 2 mg/mL, Tab 10, 25, 50, 100, 200 mg, Inj 25 mg/mL, Supp 25, 100 mg SR Cap not recommended.</td>
<td>50–1200</td>
<td>100</td>
<td>25–50</td>
<td>High</td>
</tr>
<tr>
<td>Thorazine</td>
<td>Various</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Thioridazine</td>
<td>Soln 30, 100 mg/mL, Susp 5, 20 mg/mL, Tab 10, 15, 25, 50, 100, 150, 200 mg.</td>
<td>50–800</td>
<td>100</td>
<td>—</td>
<td>High</td>
</tr>
<tr>
<td>Mellaril</td>
<td>Various</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

| **INTERMEDIATE POTENCY** | | | | | | |
| Loxapine          | Cap 5, 10, 25, 50 mg | 20–250 | 10 | 12.5–50 | Low | Low | Moderate | Low |
| Loxitane          | Soln 25 mg/mL, Inj 50 mg/mL. | | | | | | | |
| Various           |                          | | | | | | | |

(continued)
<table>
<thead>
<tr>
<th>DRUG AND CLASS</th>
<th>DOSAGE FORMS</th>
<th>ADULT ORAL DOSAGE RANGE (MG/DAY)</th>
<th>ORAL EQUIVALENT ANTIPSYCHOTIC DOSE (MG)</th>
<th>USUAL SINGLE IM DOSE (MG)</th>
<th>RELATIVE FREQUENCY OF SIDE EFFECTS</th>
<th>Orthostatic Hypotension</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Molindone</strong></td>
<td>Tab 5, 10, 25, 50, 100 mg, Soln 20 mg/mL.</td>
<td>25–225</td>
<td>10</td>
<td>—</td>
<td>Very Low</td>
<td>Low</td>
</tr>
<tr>
<td><strong>Moban</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>Low</td>
<td>Low</td>
</tr>
<tr>
<td><strong>Clozapine</strong></td>
<td>Tab 25, 100 mg.</td>
<td>300–900</td>
<td>50</td>
<td>—</td>
<td>High</td>
<td>High</td>
</tr>
<tr>
<td><strong>Clozaril</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>High</td>
<td>High</td>
</tr>
<tr>
<td><strong>Olanzapine</strong></td>
<td>Tab 2.5, 5, 7.5, 10, 15 mg.</td>
<td>10–15</td>
<td>—</td>
<td>2.5–10</td>
<td>Low</td>
<td>Low</td>
</tr>
<tr>
<td><strong>Zyprexa</strong></td>
<td>Inj (Investigational)</td>
<td></td>
<td></td>
<td></td>
<td>Low</td>
<td>Low</td>
</tr>
<tr>
<td><strong>Quetiapine</strong></td>
<td>Tab 25, 100, 200 mg.</td>
<td>150–500</td>
<td>—</td>
<td>—</td>
<td>Moderate</td>
<td>Low</td>
</tr>
<tr>
<td><strong>Seroquel</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>Low</td>
<td>Low</td>
</tr>
<tr>
<td><strong>Risperidone</strong></td>
<td>Tab 0.25, 0.5, 1, 2, 3, 4 mg.</td>
<td>4–6&lt;sup&gt;a&lt;/sup&gt;</td>
<td>—</td>
<td>—</td>
<td>Very Low</td>
<td>Low&lt;sup&gt;a&lt;/sup&gt;</td>
</tr>
<tr>
<td><strong>Risperdal</strong></td>
<td>Soln 1 mg/mL.</td>
<td></td>
<td></td>
<td></td>
<td>Very Low</td>
<td>Low</td>
</tr>
<tr>
<td><strong>Ziprasidone</strong></td>
<td>Cap 20, 40, 60, 80 mg.</td>
<td>40–160</td>
<td>—</td>
<td>10</td>
<td>Moderate</td>
<td>Very Low</td>
</tr>
<tr>
<td><strong>Geodon</strong></td>
<td>Inj (Investigational)</td>
<td></td>
<td></td>
<td></td>
<td>Very Low</td>
<td>Very Low</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>Low</td>
<td>Low</td>
</tr>
</tbody>
</table>

(continued)
### Antipsychotic Drugs Comparison Chart (continued)

<table>
<thead>
<tr>
<th>Drug and Class</th>
<th>Dosage Forms</th>
<th>Adult Oral Dosage Range (mg/day)</th>
<th>Oral Equivalent Antipsychotic Dose (mg)</th>
<th>Usual Single IM Dose (mg)</th>
<th>Relative Frequency of Side Effects</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>High Potency</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Fluphenazine</td>
<td>Elxr 0.5 mg/mL</td>
<td>2–40</td>
<td>2</td>
<td>2–5</td>
<td>Low</td>
</tr>
<tr>
<td></td>
<td>Prolixin Tab 1, 2.5, 5, 10 mg</td>
<td></td>
<td></td>
<td></td>
<td>Low</td>
</tr>
<tr>
<td></td>
<td>Inj 2.5 mg/mL.</td>
<td></td>
<td></td>
<td></td>
<td>Very High</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>Low</td>
</tr>
<tr>
<td>Fluphenazine Decanoate</td>
<td>Inj 25 mg/mL.</td>
<td>—</td>
<td>—</td>
<td>12.5–75 q 2 weeks</td>
<td>Low</td>
</tr>
<tr>
<td>Haloperidol</td>
<td>Soln 2 mg/mL.</td>
<td>2–100</td>
<td>2</td>
<td>2–5</td>
<td>Very Low</td>
</tr>
<tr>
<td></td>
<td>Haldol Tab 0.5, 1, 2, 5, 10, 20 mg</td>
<td></td>
<td></td>
<td></td>
<td>Very Low</td>
</tr>
<tr>
<td></td>
<td>Inj 5 mg/mL.</td>
<td></td>
<td></td>
<td></td>
<td>Very High</td>
</tr>
<tr>
<td>Haloperidol Decanoate</td>
<td>Inj 50, 100 mg/mL.</td>
<td>—</td>
<td>—</td>
<td>50–450 monthly</td>
<td>Very Low</td>
</tr>
</tbody>
</table>

(continued)
## Antipsychotic Drugs Comparison Chart (continued)

<table>
<thead>
<tr>
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<th>Adult Oral Dosage Range (mg/day)</th>
<th>Oral Equivalent Antipsychotic Dose (mg)</th>
<th>Usual Single IM Dose (mg)</th>
<th>Relative Frequency of Side Effects</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Perphenazine</strong></td>
<td>Soln 3.2 mg/mL, Tab 2, 4, 8, 16 mg, Inj 5 mg/mL</td>
<td>12–64</td>
<td>8</td>
<td>5–10</td>
<td>Low, Low, High, Low</td>
</tr>
<tr>
<td><strong>Trilafon</strong></td>
<td>Various</td>
<td>Trilafon Tab 2, 4, 8, 16 mg, Inj 5 mg/mL</td>
<td>5–40</td>
<td>5</td>
<td>1–2</td>
</tr>
<tr>
<td><strong>Trifluoperazine</strong></td>
<td>Soln 10 mg/mL, Tab 1, 2, 5, 10 mg, Inj 2 mg/mL</td>
<td>5–40</td>
<td>5</td>
<td>1–2</td>
<td>Low, Low, High, Low</td>
</tr>
<tr>
<td><strong>Stelazine</strong></td>
<td>Various</td>
<td>Stelazine Tab 1, 2, 5, 10 mg, Inj 2 mg/mL</td>
<td>5–60</td>
<td>4</td>
<td>2–4</td>
</tr>
<tr>
<td><strong>Thiothixene</strong></td>
<td>Cap 1, 2, 5, 10, 20 mg, Soln 5 mg/mL, Inj 5 mg/mL</td>
<td>Cap 1, 2, 5, 10, 20 mg, Soln 5 mg/mL, Inj 5 mg/mL</td>
<td>5–60</td>
<td>4</td>
<td>2–4</td>
</tr>
</tbody>
</table>

*At dosages over 6 mg/day, nausea and insomnia are limiting side effects; extrapyramidal symptoms markedly increase at dosages over 6 mg/day. From references 181–183, 192, 196, 200, 201, 209, and 215–219.*
Anxiolytics, Sedatives, and Hypnotics

Class Instructions. Sedatives and Hypnotics. This drug causes drowsiness and can produce sleep. Do not exceed the prescribed dosage and use caution when driving, operating machinery, or performing other tasks requiring mental alertness. Avoid concurrent use of alcohol or other drugs that cause drowsiness or sleep. Do not abruptly stop taking this medication; the dosage must be decreased slowly.

Missed Doses. If you miss a dose, take it as soon as you remember. If it is almost time for your next dose, skip it and resume your normal schedule. Do not double doses.

Pharmacology. Alprazolam is a triazolobenzodiazepine that is equal in efficacy to other benzodiazepines for generalized anxiety disorder but more effective in the treatment of panic disorder. Although alprazolam has some efficacy in major depression, it is less effective than heterocyclic antidepressants.220,221 (See also Clonazepam, and the Benzodiazepines and Related Drugs Comparison Chart.)

Adult Dosage. PO for generalized anxiety disorder 0.25 mg tid initially, increasing gradually to 4 mg/day. PO for panic disorder 0.5 mg tid is recommended initially; most panic patients require 5–6 mg/day, and occasionally 10 mg/day can be needed for full response. SL alprazolam tablets can be administered SL with no difference from oral administration in onset, peak, or pharmacokinetics.222 Discontinuation decrease the daily dosage by no more than 0.5 mg/day q 3 days until the daily dosage reaches 2 mg and then decrease dosage in 0.25 mg/day increments q 3 days.

Dosage Forms. Tab 0.25, 0.5, 1, 2 mg; Soln 0.1, 1 mg/mL.

Pharmacokinetics. Like diazepam, alprazolam has a rapid onset of effect after oral administration, but its shorter half-life requires tid administration. The half-life is 11 hr in adults; the elderly might have decreased clearance and an increased half-life of 21 hr.221–223

Adverse Reactions. (See Benzodiazepines.) Patients do not show complete cross-tolerance between triazolobenzodiazepines and other benzodiazepines, but clonazepam has been shown to be an effective long-half-life substitute drug for use in alprazolam withdrawal.224

BENZODIAZEPINES

Pharmacology. Benzodiazepines have a more specific anxiolytic effect than other sedatives. Benzodiazepines facilitate the inhibitory effect of GABA on neuronal excitability by increasing membrane permeability to chloride ions.225

Administration and Adult Dosage. (See Benzodiazepines and Related Drugs Comparison Chart.) Optimal oral use requires individual dosage titration to clinical response. The long-acting drugs can be administered once daily hs; the short-acting drugs require multiple daily doses. (See Benzodiazepines and Related Drugs Comparison Chart.) Determine the dosage schedule by the individual pa-
tient’s relative degree of dysfunction from daytime anxiety compared with insomnia. Despite physiologic dependence, benzodiazepines might need to be used for months and sometimes years for treatment of panic disorder and generalized anxiety disorder; situational anxiety, adjustment disorders, and anxiety secondary to other causes require only days to weeks of drug treatment.²²⁶ **PO for alcohol withdrawal** evidence suggests no superiority of any benzodiazepine in alcohol withdrawal, although **chlordiazepoxide** has been most adequately studied; (chlordiazepoxide) 25–100 mg for agitation, anxiety, and tremor; on the first day, up to 400 mg can be given in divided doses, with gradual dosage reductions over 4 days; **(diazepam)** 5–20 mg for agitation, anxiety, and tremor; alternatively, it can be given in 20 mg doses q 2 hr until complete suppression of signs and symptoms is achieved. After this loading dose, further administration is unnecessary;²²⁷ (oxazepam) 15–60 mg q 4–6 hr for agitation, anxiety, and tremor. Oxazepam is preferred in patients with severe liver disease. IM chlordiazepoxide is not recommended because of slow, erratic absorption; however, **lorazepam** is suitable for IM administration.¹⁴¹,²²⁷ Diazepam injectable solution (Valium, various) can be administered IM or IV; the injectable emulsion (Dizac) is for IV use only (do not administer IM or SC); neither the solution nor the emulsion should be administered faster than 5 mg/min into a peripheral vein, and small veins should be avoided; neither product is recommended to be added to other drugs or solutions. (See Fate.) **PR diazepam for seizures** 0.2 mg/kg of Diastat rectal gel, rounded up to the next dosage size (2.5, 5, 10, 15, 20 mg); an additional dose can be given 4–12 hr after the first dose. Treat no more than 1 episode q 5 days or 5 episodes/month with Diastat.

**Special Populations. Pediatric Dosage.** PO (diazepam, >6 months) 1–2.5 mg tid or qid. Most benzodiazepines are not recommended in children because of insufficient clinical experience and concern about the stimulating and paradoxical effects that occur because of disinhibition. **Midazolam** is commonly used in children for preanesthetic sedation. (See Midazolam.) **PR diazepam for seizures** (2–5 yr) 0.5 mg/kg of Diastat rectal gel, rounded up to the next dosage size (2.5, 5, 10, 15, 20 mg); (6–11 yr) 0.3 mg/kg of Diastat rectal gel, rounded up to the next dosage size. An additional dose may be given 4–12 hr after the first dose. Treat no more than 1 episode q 5 days or 5 episodes/month with Diastat.

**Geriatric Dosage.** The elderly might have reduced clearance and enhanced CNS sensitivity, which requires initial dosage to be reduced by 33–50%.²²⁸

**Other Conditions.** Higher dosages might be needed in heavy smokers. Patients with liver disease might have reduced clearance and/or enhanced CNS sensitivity, which requires reduction of initial and subsequent doses. Alcoholic patients with reduced plasma proteins might require a lower dosage because of decreased protein binding.

**Dosage Forms.** (See Benzodiazepines and Related Drugs Comparison Chart.)

**Patient Instructions.** (See Sedatives and Hypnotics Class Instructions.)

**Pharmacokinetics. Serum Levels.** Not used clinically.
Fate. Diazepam and chlordiazepoxide are absorbed faster and more completely orally than intramuscularly. Lorazepam and midazolam have rapid and reliable IM absorption.\textsuperscript{141,229} (See Benzodiazepines and Related Drugs Comparison Chart.)

Adverse Reactions. Frequent effects include drowsiness, dizziness, ataxia, and disorientation; these effects rarely require drug discontinuation and are easily managed by dosage reduction. Anterograde amnesia is frequent.\textsuperscript{141} Occasionally, agitation and excitement occur.\textsuperscript{230} With parenteral therapy, hypotension and respiratory depression occur occasionally. Rarely, hepatotoxicity or blood dyscrasias occur. Diazepam emulsion is associated with less venous thrombosis and phlebitis than the solution, which can be very irritating to veins.

Contraindications. Acute narrow-angle glaucoma; (diazepam emulsion injection) hypersensitivity to soy protein.

Precautions. Pregnancy; impaired hepatic function. Abrupt drug withdrawal can result in rebound insomnia, abstinence syndrome similar to barbiturate withdrawal, seizures, or, rarely, psychosis. Patients do not show complete cross-tolerance between triazolobenzodiazepines and other benzodiazepines. History of substance abuse can indicate increased likelihood of benzodiazepine misuse.\textsuperscript{225}

Drug Interactions. Concurrent use with other CNS depressants can potentiate the sedation caused by benzodiazepines. Nefazodone inhibits alprazolam and triazolam metabolism; fluoxetine and fluvoxamine increase levels of alprazolam and diazepam; omeprazole increases serum diazepam levels.

Parameters to Monitor. Periodically reassess the need for therapy during long-term use.

BUSPIRONE HYDROCHLORIDE BuSpar, Various

Pharmacology. Buspirone is the first of a class of selective serotonin-5-HT\textsubscript{1A} receptor partial agonists. It also has some effect on dopamine-D\textsubscript{2} autoreceptors and, like antidepressants, can downregulate \(\beta\)-adrenergic receptors. Unlike benzodiazepines, it lacks amnestic, anticonvulsant, muscle relaxant, and hypnotic effects. Its exact anxiolytic mechanism of action is complex and not clearly defined.\textsuperscript{231,232}

Administration and Adult Dosage. PO for anxiety 5 mg tid or 7.5 mg bid for 1 week, increasing in 5 mg/day increments q 2–3 days to a maximum of 60 mg/day in 2 or 3 divided doses. Most patients require 20–30 mg/day in divided doses.

Special Populations. Pediatric Dosage. Safety and efficacy not established.

Geriatric Dosage. Same as adult dosage.

Other Conditions. Decrease the initial dose to 5 mg bid in patients with hepatic or renal impairment.\textsuperscript{231,233}

Dosage Forms. Tab 5, 7.5, 10, 15 mg.

Patient Instructions. This drug requires several weeks of continuous use for therapeutic effect and is not effective when used intermittently.

Pharmacokinetics. Onset and Duration. Onset of anxiolytic effect can take several weeks.
**Fate.** The drug is well absorbed; oral bioavailability is 3.9 ± 4.3%. Administration after meals increases bioavailability by 80%. It is extensively metabolized by oxidative dealkylation pathways. \( t_{1/2} \) 2.1 ± 1.2 hr.\(^{234}\)

**Adverse Reactions.** Dosages >60 mg/day can cause dysphoria.\(^{225}\) Frequent nausea, dizziness, headache, and insomnia occur. Unlike benzodiazepines, buspirone does not cause dependence or withdrawal effects.\(^{231,235}\)

**Contraindications.** None known.

**Precautions.** Buspirone has no cross-tolerance with benzodiazepines, so patients being switched from a benzodiazepine should have their dosages of the benzodiazepine decreased slowly.

**Drug Interactions.** Unlike benzodiazepines, buspirone does not interact with alcohol.\(^{231,235}\) Buspirone can increase haloperidol serum levels. Avoid concurrent buspirone and an MAOI because the combination can cause hypertension.

**Parameters to Monitor.** Monitor renal and hepatic function initially and periodically during long-term therapy.

**Notes.** Buspirone is indicated only for the treatment of generalized anxiety disorder and is not effective as a prn medication or hypnotic. Buspirone’s anxiolytic effect without sedation or respiratory depression has led to its use in agitation and anxiety, dementia, mental retardation, and spinal cord injury. Its unique effect on the 5-HT\(_{1A}\) receptor has led to uncontrolled studies and clinical use for premenstrual tension syndrome and to decrease craving in smoking cessation.\(^{236}\)

**FLUMAZENIL**

**Pharmacology.** Flumazenil is a selective inhibitor of the CNS effects of benzodiazepine sedatives. It competitively blocks the effect of benzodiazepines and zolpidem on GABA-mediated inhibitory pathways within the CNS. Flumazenil finds its greatest use in the reversal of benzodiazepine sedation after medical and surgical procedures and occasionally in the management of benzodiazepine overdose.\(^{237,239}\)

**Adult Dosage.** **IV for reversal of conscious sedation** 0.2 mg over 15 sec; this dose can be repeated after 45 sec and every minute thereafter prn, to a total dosage of 1 mg. **IV for benzodiazepine overdose** 0.2 mg over 30 sec, followed, if necessary, by 0.3 mg after 30 sec. Further doses of 0.5 mg over 30 sec can be given at 1-min intervals to a cumulative dosage of 3 mg. Rarely, patients who respond partially to 3 mg respond more completely to a dosage of 5 mg. If resedation occurs after either use, additional doses of up to 1 mg can be given at 20-min intervals to a maximum of 3 mg/hr. Flumazenil does not consistently reverse benzodiazepine amnesia, so give patients written instructions to avoid operation of motor vehicles or hazardous equipment, or ingestion of alcohol or nonprescription medications for 18–24 hr, or longer if benzodiazepine effects persist.

**Dosage Forms.** **Inj** 0.1 mg/mL.

**Pharmacokinetics.** Reversal of benzodiazepine coma can occur within 1–2 min and last 1–5 hr, depending on the dosages of the benzodiazepine and flumazenil.
474  CENTRAL NERVOUS SYSTEM DRUGS

First-pass hepatic metabolism limits the bioavailability of oral flumazenil, so the drug is administered by IV injection. It is rapidly hydroxylated in the liver to inactive metabolites. $V_d$ is 0.6–1.6 L/kg; its elimination half-life is 0.7–1.3 hr.

**Adverse Reactions.** Frequent side effects have been minimal and are usually limited to nausea and vomiting, anxiety, and agitation. However, seizures have occurred, most often in patients on long-term benzodiazepine therapy or after overdose with heterocyclic antidepressants or other potentially convulsant drugs (eg, bupropion, cocaine, cyclosporine, isoniazid, lithium, methylxanthines, MAOIs, propoxyphene). Be prepared to manage seizures before giving flumazenil.240

**MIDAZOLAM HYDROCHLORIDE  Versed**

**Pharmacology.** Midazolam is a short-acting triazolobenzodiazepine for use in anesthesia. It is unique in its physicochemical properties; at a pH under 4 the drug exists as a highly water-soluble, stable compound, but at physiologic pH, it becomes lipophilic. This allows IV administration of a water-soluble, rapidly acting drug with a very low frequency of venous irritation. Midazolam is given IM for preoperative sedation and IV for induction of anesthesia or for conscious sedation for endoscopy and other procedures.241-243

**Adult Dosage.** IM for preoperative sedation 0.07–0.08 mg/kg (about 5 mg) 1 hr before surgery. IV for endoscopy and other conscious sedation procedures dosage must be individualized and not administered by rapid bolus. Titrate slowly to desired effect; some patients might respond to as little as 1 mg. Give no more than 2.5 mg over at least 2 min as the 1 mg/mL (or more dilute) solution; in elderly, debilitated, or chronically ill patients, limit the initial dose to 1.5 mg. Further small doses can be given after waiting at least 2 min. Do not give the drug IV without oxygen and resuscitation equipment immediately available.

**Pediatric Dosage.** PO for sedation (6 mo–16 yr) 0.25–1 mg/kg (usually 0.5 mg/kg) to a maximum of 20 mg. PR for preanesthetic sedation 0.3 mg/kg as a solution diluted in 5 mL of saline is a safe and effective alternative to IM administration.244

**Dosage Forms.** Inj 1, 5 mg/mL; Syrup 2 mg/mL.

**Pharmacokinetics.** Midazolam is >90% absorbed after IM injection; peak serum levels occur within 30 min. Peak levels after IM administration are about 50% of IV levels. PO onset is 10–20 min; IM onset is about 15 min. The drug is 97% bound to plasma proteins and has a $V_d$ of 1–3 L/kg; Cl is 0.25–0.54 L/hr/kg. Midazolam is hepatically metabolized via CYP3A4 to the 1-hydroxy- and 4-hydroxy-metabolites; the 1-hydroxy-metabolite is at least as active as midazolam. Midazolam’s half-life is 1.8–6.4 hr.

**Adverse Reactions.** Respiratory depression and respiratory arrest occur frequently. Impairment of psychomotor skills continues after acute sedation has passed, so patients should not drive or operate machinery until it is clear that they have recovered fully.
Pharmacology. Triazolam is a triazolobenzodiazepine hypnotic whose effect is likely related to its facilitation of GABA-mediated neurotransmission, but its exact mechanism is unknown.

Administration and Adult Dosage. (See Benzodiazepines and Related Drugs Comparison Chart.) PO as a hypnotic 0.25 mg hs initially; do not exceed 0.5 mg.

Special Populations. Pediatric Dosage. (<18 yr) safety and efficacy not established.

Geriatric Dosage. PO decrease initial dose to 0.125 mg, increase if necessary to 0.25 mg hs.245,246

Other Conditions. PO 0.125 mg initially in debilitated patients and those with low body weights or with hepatic impairment.

Dosage Forms. Tab 0.125, 0.25 mg.

Patient Instructions. (See Sedatives and Hypnotics Class Instructions.)

Pharmacokinetics. Onset and Duration. Onset of hypnotic effect is 0.5–1 hr, with peak serum levels achieved within 2 hr.

Fate. Oral bioavailability 44%, SL 53%, because of nonhepatic presystemic metabolism. Vdₐ is 1.2 ± 0.5 L/kg; Cl is 0.34 ± 0.2 L/hr/kg. Cl decreases with advancing age (attributed to reduced hepatic oxidizing capacity in the elderly). Triazolam undergoes hydroxylation and rapid conjugation. Smoking does not affect elimination.118,247 Accumulation does not occur with multiple doses.

t¹⁄₂. 2.6 ± 1 hr. Half-life is not affected by end-stage renal disease or liver disease.118,247,248

Adverse Reactions. Frequently, anterograde amnesia,249 daytime anxiety, and ataxia occur. Occasionally, agitation, confusion, or mood disturbance occur. Rarely, respiratory depression, depersonalization, and derealization, or psychosis occur. Unlike other benzodiazepines, several fatalities have been reported in elderly patients who overdosed on triazolam.250,251

Contraindications. Pregnancy.

Precautions. Pregnancy; impaired hepatic function. Abrupt drug withdrawal can result in rebound insomnia, abstinence syndrome similar to barbiturate withdrawal, seizures, or, rarely, psychosis. Patients do not show complete cross-tolerance between triazolam and other benzodiazepines. History of substance abuse can indicate an increased likelihood of triazolam misuse.225 Do not prescribe the drug for more than 7–10 days of consecutive therapy or in quantities larger than a 30-day supply.

Drug Interactions. Concurrent use with other CNS depressants can potentiate the sedation caused by benzodiazepines. Nefazodone inhibits triazolam metabolism.

Notes. Compared with other benzodiazepine hypnotics, triazolam is equally effective in reducing sleep latency and less likely to cause daytime sedation; however, it is less likely to prevent early morning awakening and more likely to cause rebound insomnia. Hypnotic drugs are most effective when used to treat transient situational insomnia (1–3 days) and short-term insomnia (1–3 weeks maximum).251,252
Pharmacology. Zaleplon is a nonbenzodiazepine hypnotic that, like zolpidem, selectively binds only to the \( \omega_1 \) receptor. This selectivity suggests a sedative effect with less potential for memory impairment, interaction with alcohol, and psychomotor effects than with benzodiazepines.\(^{253}\)

Administration and Adult Dosage. PO as a hypnotic 10 mg hs initially (5 mg in the elderly). Because of its very rapid onset and offset, zaleplon can be given during the night after the patient experiences difficulty falling asleep rather than being given before bedtime in anticipation of sleep difficulty. Zaleplon can be given during the night without morning hangover as long as there are 4 hr remaining in bed after administration.

Dosage Forms. Cap 5, 10 mg.

Pharmacokinetics. After oral administration, zaleplon reaches peak serum concentrations in 1.1 hr. Zaleplon is metabolized by CYP3A4 but has no active metabolites. Its half-life is 0.8–1.4 hr (average 1).\(^{254}\)

Adverse Reactions. Dose-related side effects include dizziness, headache, and somnolence. Symptoms begin to appear approximately 30 min after a dose, peak at 1–2 hr, and are no longer evident at 4 hr. After a 10-mg dose, zaleplon has no residual effects on performance and memory tests after 2 hr; in contrast, residual effects persist for up to 5 hr with zolpidem.\(^{255}\)

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ZOLPIDEM TARTRATE

Pharmacology. Zolpidem is a short-acting nonbenzodiazepine hypnotic indicated for the short-term treatment of insomnia. Most benzodiazepines bind to all GABA-benzodiazepine (\( \omega \)) receptor complexes, but zolpidem selectively binds only to the \( \omega_1 \) receptor. This difference suggests a more selective sedative–hypnotic effect without anxiolytic, anticonvulsant, or muscle relaxant effects.\(^{256–258}\) (See Benzodiazepines and Related Drugs Comparison Chart.)

Adult Dosage. PO as a hypnotic 10 mg immediately before bedtime. In the elderly, patients with hepatic impairment, or patients taking other CNS depressants, the dose is 5 mg.

Dosage Forms. Tab 5, 10 mg.

Pharmacokinetics. After oral administration, zolpidem reaches peak serum concentrations in 1.6 hr, is 93% bound to plasma proteins, has no active metabolites, and has an elimination half-life of 1.5–4 hr (average 2.5). Half-life is increased by one-third in the elderly and greatly increased in patients with hepatic impairment (9.9 hr).

Adverse Reactions. Dose-related side effects include daytime drowsiness, dizziness, and diarrhea. Clinical trials with 20 mg doses have reported headache, nausea, memory problems, and CNS stimulation. Tolerance has not been reported, nor has rebound insomnia after therapeutic doses. Psychomotor performance is impaired when zolpidem is combined with alcohol. Efficacy has been demonstrated for 35 nights at doses of 10 mg without affecting sleep stages or psychomotor performance.
### Benzodiazepines and Related Drugs Comparison Chart

<table>
<thead>
<tr>
<th>DRUG AND SCHEDULEA</th>
<th>DOSAGE FORMS</th>
<th>ADULT ORAL DOSAGE RANGE</th>
<th>PEAK ORAL SERUM LEVELS (HR)</th>
<th>HALF-LIFE (HR)b</th>
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</thead>
<tbody>
<tr>
<td><strong>SHORT-ACTING ANXIOLYTICS</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Alprazolam (C-IV)</td>
<td>Tab 0.25, 0.5, 1, 2 mg</td>
<td>0.75–4 mg/day²</td>
<td>0.7–1.6</td>
<td>11–21</td>
</tr>
<tr>
<td>Xanax</td>
<td>Various</td>
<td>5–10 mg/day³</td>
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<tr>
<td>Lorazepam* (C-IV)</td>
<td>Tab 0.5, 1, 2 mg</td>
<td>2–10 mg/day</td>
<td>2</td>
<td>10–20</td>
</tr>
<tr>
<td>Ativan</td>
<td>Various</td>
<td>Soln 2 mg/mL</td>
<td>Inj 2, 4 mg/mL</td>
<td></td>
</tr>
<tr>
<td>Oxazepam (C-IV)</td>
<td>Cap 10, 15, 30 mg</td>
<td>30–120 mg/day</td>
<td>1–2</td>
<td>5–15</td>
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<tr>
<td>Serax</td>
<td>Various</td>
<td>Tab 15 mg</td>
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<td><strong>LONG-ACTING ANXIOLYTICS</strong></td>
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<tr>
<td>Chlordiazepoxide (C-IV)</td>
<td>Cap 5, 10, 25 mg</td>
<td>15–100 mg/day</td>
<td>2–4</td>
<td>&gt;24</td>
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<tr>
<td>Librium</td>
<td>Tab 10, 25 mg</td>
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<tr>
<td>Libritabs</td>
<td>Inj 100 mg</td>
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<tr>
<td>Clorazepate (C-IV)</td>
<td>Cap 3.75, 7.5, 15 mg</td>
<td>15–60 mg/day</td>
<td>1–2³</td>
<td>&gt;24</td>
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<td>Tranxene</td>
<td>Tab 3.75, 7.5, 15 mg</td>
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<td>Various</td>
<td>SR Tab 11.25, 22.5 mg</td>
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<td>Diazepam (C-IV)</td>
<td>Tab 2, 5, 10 mg</td>
<td>6–40 mg/day</td>
<td>1–2</td>
<td>&gt;24</td>
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<tr>
<td>Dizac</td>
<td>Soln 1, 5 mg/mL</td>
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<tr>
<td>Valium</td>
<td>Inj 5 mg/mL</td>
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<td>Various</td>
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<tr>
<td>Halazepam (C-IV)</td>
<td>Tab 20, 40 mg</td>
<td>60–160 mg/day</td>
<td>1–3</td>
<td>&gt;24</td>
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<tr>
<td>Paxipam</td>
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<tr>
<td><strong>SHORT-ACTING HYPNOTICS</strong></td>
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<tr>
<td>Midazolam* (C-IV)</td>
<td>Inj 1, 5 mg/mL</td>
<td>—</td>
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<td>Versed</td>
<td>Syrup 2 mg/mL</td>
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<td>Triazolam (C-IV)</td>
<td>Tab 0.125, 0.25 mg</td>
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<td>Halcion</td>
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<td>Zaleplon° (C-IV)</td>
<td>Cap 5, 10 mg</td>
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<tr>
<td>Zolpidem° (C-IV)</td>
<td>Tab 5, 10 mg</td>
<td>5–20 mg</td>
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<tr>
<td>Ambien</td>
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<thead>
<tr>
<th>DRUG AND SCHEDULE</th>
<th>DOSAGE FORMS</th>
<th>ADULT ORAL DOSAGE RANGE</th>
<th>PEAK ORAL SERUM LEVELS (HR)</th>
<th>HALF-LIFE (HR)</th>
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<tr>
<td><strong>INTERMEDIATE-ACTING HYPNOTICS</strong></td>
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<tr>
<td><em>Estazolam (C-IV)</em></td>
<td>Tab 1, 2 mg.</td>
<td>1–2 mg</td>
<td>1–2</td>
<td>12–15</td>
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<td>Pro-Som</td>
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<tr>
<td><strong>Temazepam (C-IV)</strong></td>
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<td>7.5–30 mg</td>
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<td>10–15</td>
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<td>Restoril</td>
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<td>Various</td>
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<tr>
<td><strong>LONG-ACTING HYPNOTICS</strong></td>
<td></td>
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<tr>
<td><em>Flurazepam (C-IV)</em></td>
<td>Cap 15, 30 mg.</td>
<td>15–30 mg</td>
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<td>&gt;24^a</td>
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<td>Dalmane</td>
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<tr>
<td><em>Quazepam (C-IV)</em></td>
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<td>7.5–15 mg</td>
<td>1–2</td>
<td>&gt;24^a</td>
</tr>
<tr>
<td>Doral</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

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^aControlled substance schedule designated after each drug (in parentheses).
^bParent drug plus active metabolites.
^cFor generalized anxiety disorder.
^dFor panic disorder.
^eAlso used as an IV anesthetic; well absorbed IM.
^fHydrolyzed to nordazepam (desmethyldiazepam) before absorption.
^gNot a benzodiazepine chemically, but an imidazopyridine, which is a selective benzodiazepine-1 receptor agonist.
^hRapidly and completely metabolized to desalkylflurazepam.

From references 221, 222, 225, 229, 241, 247, 249, and 258–262.
# Sedatives and Hypnotics Comparison Chart

<table>
<thead>
<tr>
<th>Drug and Schedule</th>
<th>Dosage Forms</th>
<th>Oral Dosage</th>
<th>Half-Life (HR)</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>SEDATIVES</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Meprobamate (C-IV)</td>
<td>Tab 200, 400, 600 mg</td>
<td>400 mg tid or qid or 600 mg bid</td>
<td>6–16</td>
</tr>
<tr>
<td>Equanil</td>
<td>SR Cap 200, 400 mg</td>
<td></td>
<td></td>
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<tr>
<td>Miltown</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Various</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Phenobarbital (C-IV)</td>
<td>Cap 16 mg Elixir 3, 4 mg/mL Tab 8, 15, 16, 30, 60, 90, 100 mg Inj 30, 60, 65, 130 mg/mL</td>
<td>15–30 mg bid-qid</td>
<td>48–120</td>
</tr>
<tr>
<td>Various</td>
<td></td>
<td></td>
<td></td>
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<tr>
<td><strong>HYPNOTICS</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Chloral Hydrate (C-IV)</td>
<td>Cap 500 mg Supp 325, 500, 650 mg Syrup 50, 100 mg/mL</td>
<td>500 mg–1.5 g (Trichloroethanol)</td>
<td>8</td>
</tr>
<tr>
<td>Noctec</td>
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<tr>
<td>Various</td>
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<td></td>
<td></td>
</tr>
<tr>
<td>Ethchlorvynol (C-IV)</td>
<td>Cap 200, 500, 750 mg</td>
<td>500 mg–1 g</td>
<td>6</td>
</tr>
<tr>
<td>Placidyl</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Various</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Glutethimide (C-II)</td>
<td>Tab 250 mg</td>
<td>250–500 mg</td>
<td>5–22</td>
</tr>
<tr>
<td>Various</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Pentobarbital (C-II)</td>
<td>Cap 50, 100 mg Elixir 4 mg/mL Supp 30, 60, 120, 200 mg (C-III) Inj 50 mg/mL</td>
<td>100–200 mg</td>
<td>21–42</td>
</tr>
<tr>
<td>Nembutal</td>
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<td></td>
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</tr>
<tr>
<td>Various</td>
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<td></td>
</tr>
<tr>
<td>Secobarbital (C-II)</td>
<td>Cap 100 mg</td>
<td>100–200 mg</td>
<td>19–34</td>
</tr>
<tr>
<td>Seconal</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Various</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

*Controlled substance schedule designated after each drug (in parentheses). From references 251, 252, 262, and 263.*
Lithium

Pharmacology. Lithium’s mechanism of antimanic effect is unknown; it alters the actions of several second-messenger systems (eg, adenylate cyclase and phosphoinositols).264,265

Administration and Adult Dosage. Individualize dosage according to serum levels and clinical response. Acute manic episodes typically require PO 1.2–2.4 g/day; maintenance therapy requires 900 mg–1.5 g/day. A loading dose of 30 mg/kg, in 3 divided doses, can be given to achieve the desired serum level within 12 hr.266 A number of predictive dosage techniques have been developed based on estimated steady state after one serum level.267,268

Special Populations. Pediatric Dosage. PO (<12 yr) 15–20 mg (0.4–0.5 mEq)/kg/day in 2–3 divided doses; (12–18 yr) same as adult dosage.269

Geriatric Dosage. (>65 yr) decrease adult dosage by 33–50%.270

Other Conditions. Adjust the dosage more carefully in patients with decreased renal function and in patients receiving thiazide diuretics or NSAIDs.

Dosage Forms. Cap 150, 300, 600 mg; Tab 300 mg; SR Tab 300, 450 mg; Syrup 1.6 mEq/mL (as citrate).

Patient Instructions. This drug can be taken with food, milk, or antacid to minimize stomach upset. Report immediately if signs of toxicity occur, such as persistent diarrhea, vomiting, coarse hand tremor, drowsiness, or slurred speech, or before beginning any diet. In hot weather, ensure adequate water and salt intake.

Pharmacokinetics. Onset and Duration. Onset 7–10 days for therapeutic effect.141

Serum Levels. (Acute mania or hypomania) 0.8–1.5 mEq/L; (prophylaxis) 0.6–1.2 mEq/L, although concern about long-term renal effects suggests most patients should be maintained <0.9 mEq/L. Levels >1.5 mEq/L are regularly associated with some signs of toxicity, and levels >2 mEq/L result in serious toxicity.271 (See Adverse Reactions.)

Fate. Absorption is virtually complete within 8 hr after oral administration, with peak levels occurring in 2–4 hr. Distribution is throughout total body water, but tissue uptake is not uniform. The drug is not protein bound or metabolized, but freely filtered through the glomerulus, with about 80% being reabsorbed.

t₁/₂. 18–20 hr; up to 36 hr in the elderly.141

Adverse Reactions. Frequent, dose-related effects with therapeutic serum levels include nausea, diarrhea, polyuria, polydipsia, fine hand tremor, and muscle weakness. Signs of toxicity include coarse hand tremor, persistent GI effects, muscle hyperirritability, slurred speech, confusion, stupor, seizures, increased deep tendon reflexes, irregular pulse, and coma. Frequent, non–dose-related effects include nontoxic goiter, hypothyroidism, nephrogenic diabetes insipidus-like syndrome,
foliculitis, aggravation of acne or psoriasis, leukocytosis, hypercalcemia, and weight gain.\textsuperscript{272,273}

**Contraindications.** Pregnancy; fluctuating renal function; severe renal or cardiovascular disease.

**Precautions.** Use with caution in patients with cardiac disease, dehydration, sodium depletion, diuretic therapy, or dementia, in nursing mothers, and in the elderly. (See Special Populations.)

**Drug Interactions.** ACE inhibitors can increase serum lithium concentrations. Theophylline or excess sodium enhance renal lithium clearance; sodium deficiency can promote lithium retention and increase risk of toxicity. Long-term diuretic or NSAID use can result in decreased lithium elimination. Haloperidol can increase the CNS toxicity of lithium; with methylxanthine or phenytoin, signs of lithium toxicity can occur without increased serum lithium.

**Parameters to Monitor.** Prelithium workup should include thyroid function tests, Cr, BUN, CBC (for baseline WBC count), urinalysis (for baseline specific gravity), electrolytes, and ECG (if patient is older than 40 yr). During therapy, obtain serum lithium levels (drawn 12 hr after last dose) weekly during initiation and monthly during maintenance.\textsuperscript{274,275}

**Notes.** Divalproex sodium is equivalent in efficacy to lithium for bipolar disorder and more effective than lithium for rapid-cycling bipolar illness.\textsuperscript{276–278}

## Neurodegenerative Disease Drugs

### AMANTADINE

**Pharmacology.** Amantadine is an antiviral compound that prevents the release of viral nucleic acid into the host cell. In Parkinson’s disease, the drug increases presynaptic dopamine release, blocks the reuptake of dopamine into the presynaptic neurons, and exerts anticholinergic effects. Amantadine also can reduce levodopa-induced dyskinesias in patients with Parkinson’s disease, possibly by acting as an N-methyl-D-aspartate receptor antagonist.\textsuperscript{279,280}

**Administration and Adult Dosage.** PO for Parkinson’s disease, give 100 mg/day initially, increasing in 100 mg/day increments q 7–14 days to effective dosage, or to a maximum of 300 mg/day in divided doses. **Usual maintenance dosage** 100 mg bid. PO for extrapyramidal reactions 100 mg bid, to a maximum of 300 mg/day in 3 divided doses. PO for prophylaxis of influenza A 200 mg/day in 1–2 divided doses continuing for at least 10 days after exposure, for 2–3 weeks after giving influenza A vaccine, or for up to 90 days when vaccine is unavailable or contraindicated. PO for treatment of influenza A 200 mg/day in 1–2 divided doses starting within 24–48 hr after onset of illness and continuing for 24–48 hr after symptoms disappear.

**Special Populations.** **Pediatric Dosage.** PO for prophylaxis or treatment of influenza A (<1 yr) safety and efficacy not established; (1–9 yr) 4.4–8.8 mg/kg/day in 2 divided doses, to a maximum of 150 mg/day; (9–12 yr) 100 mg bid. For prophylaxis, continue therapy for at least 10 days after exposure, for 2–3 days after
giving influenza A vaccine, or for up to 90 days when vaccine is unavailable or contraindicated. For treatment, continue for 24–48 hr after symptoms disappear.

**Geriatric Dosage.** PO for influenza prophylaxis or treatment (>65 yr) 100 mg/day. PO for Parkinson’s disease same as adult dosage, adjusting for renal impairment.

**Other Conditions.** Reduce dosage in renal impairment as follows: with Cl$_r$ of 30–50 mL/min, give 200 mg first day and then 100 mg/day; with Cl$_r$ of 15–29 mL/min, give 200 mg first day and then 100 mg every other day; with Cl$_r$ <15 mL/min or for patients on hemodialysis, give 200 mg q 7 days.

**Dosage Forms.** Cap 100 mg; Syrup 10 mg/mL.

**Patient Instructions.** This medication can cause dizziness, confusion, or difficulty in concentrating. Until the extent of these effects is known, use caution when driving, operating machinery, or performing other tasks requiring mental alertness. Avoid excessive concurrent use of alcohol. Parkinson’s disease Stopping this medication suddenly can cause your Parkinson’s disease to worsen.

**Missed Doses.** Take this drug at regular intervals. If you miss a dose, take it as soon as you remember. If it is about time for the next dose, take that dose only. Do not double the dose or take extra.

**Pharmacokinetics. Onset and Duration.** Antiparkinson effects disappear in most patients after 6–12 weeks of therapy.\(^{281}\)

**Serum Levels.** (Therapeutic trough, antiviral) 300 µg/L (2 µmol/L).

**Fate.** Peak serum levels occur in 1–4 hr in young adults, 4.5–7 hr in older adults. Steady-state serum levels occur in healthy volunteers and Parkinson’s patients within 4–7 days;\(^{282}\) V$_d$ is 6.6 ± 1.5 L/kg; Cl is 0.39 ± 0.13 L/hr/kg with normal renal function. From 78% to 88% is excreted unchanged in urine.\(^{283}\)

$\tau$/\(_{\text{1/2}}\). (Healthy young adults) 11.8 ± 2.1 hr,\(^{284}\) (elderly adults) 31 ± 7.2 hr,\(^{285}\) (during chronic hemodialysis) 8.3 ± 1.5 days.\(^{284}\)

**Adverse Reactions.** Nausea, dizziness, insomnia, confusion, hallucinations, anxiety, restlessness, depression, irritability, peripheral edema, orthostatic hypotension, or livedo reticularis occur frequently. Occasionally, CHF, psychosis, urinary retention, or reversible elevation of liver enzymes can occur. Seizures, corneal opacities, or leukopenia rarely occur.

**Drug Interactions.** Amantadine can potentiate the CNS effects of anticholinergic agents.

**Precautions.** Pregnancy; lactation. Use with caution in patients with CHF, seizures, renal or hepatic disease, peripheral edema, orthostatic hypotension, psychosis, or history of eczematoid rash or in those receiving CNS stimulants. Abrupt drug discontinuation in patients with Parkinson’s disease can result in rapid clinical deterioration. Observe patients carefully when dosages of amantadine are reduced abruptly or discontinued, especially if patients are receiving neuroleptics. Sporadic cases of neuroleptic malignant syndrome have been reported in association with amantadine withdrawal or dosage reduction. Suicide attempts have been reported in patients treated with amantadine, including short influenza treatment,
and in patients with and without psychiatric histories. Amantadine can exacerbate mental problems in patients with psychiatric disorders.

**Parameters to Monitor.** Monitor renal function and disease symptoms periodically in parkinsonian patients.

**Notes.** In Parkinson’s disease, amantadine is indicated as initial treatment alone or in combination with levodopa. Amantadine produces clinical improvements in akinesia and rigidity, but to a lesser degree than levodopa. Amantadine also can be beneficial in Parkinson’s disease patients with nighttime monoclonus, freezing, or dystonia. There is no evidence that amantadine alters the course of Parkinson’s disease. Anticholinergics appear to reduce tremor to a greater degree than amantadine.

Rimantadine (Flumadine) is an antiviral compound with efficacy similar to amantadine against influenza A. It appears to be slightly better tolerated than amantadine. Dosage is 100 mg bid in adults. In elderly nursing home patients or those with severe hepatic dysfunction or renal failure (Cl_{cr} \leq 10 \text{ mL/min}), reduce dosage to 100 mg/day. In children younger than 10 yr (prophylaxis only), give 5 mg/kg once daily, not to exceed 150 mg. For children 10 yr and older, use the adult dose. Available as 100 mg tablets and 10 mg/mL syrup.

**BENZTROPINE MESYLATE** Cogentin

**Pharmacology.** Benztropine is a synthetic competitive antagonist of acetylcholine. In Parkinson’s disease, the drug reduces the relative excess of cholinergic activity in the basal ganglia that develops because of absolute dopamine deficiency in this area.

**Administration and Adult Dosage.** PO, IM, or IV for Parkinson’s disease: 0.5–1 mg/day initially, increasing in 0.5 mg/day increments q 5–6 days to effective dosage, to a maximum of 6 mg/day. **Usual maintenance dosage** 1–2 mg/day in 2–3 divided doses. When used concurrently with levodopa, the dosages of both drugs might require reduction. PO, IM, or IV for drug-induced extrapyramidal disorders 1–4 mg/day in 1–2 doses.

**Special Populations. Pediatric Dosage.** (<3 yr) contraindicated; (>3 yr) 0.02–0.05 mg/kg/dose once or twice daily.

**Geriatric Dosage.** Same as adult dosage, although older patients often can be controlled with 1–2 mg/day. Some consider it best to avoid this drug in the elderly.

**Dosage Forms.** Tab 0.5, 1, 2 mg; Inj 1 mg/mL.

**Patient Instructions.** This drug can cause constipation, difficult or painful urination, dry mouth, blurred vision, or drowsiness. Use caution when driving, operating machinery, or performing other tasks requiring mental alertness. Avoid excessive concurrent use of alcohol and other drugs that cause drowsiness.

**Missed Doses.** Take this drug at regular intervals. If you miss a dose, take it as soon as you remember. If it is about time for the next dose, take that dose only. Do not double the dose or take extra.

**Pharmacokinetics. Onset and Duration.** Onset of resolution of drug-induced extrapyramidal symptoms is within 15 min after IV or IM administration and 1–2 hr after oral administration. Duration is 24 hr.
**Fate.** Benztrapine pharmacokinetics are not well studied, but the drug apparently is hepatically metabolized to conjugates and might undergo enterohepatic recycling.287

**Adverse Reactions.** Frequent adverse effects are dose related and include dry mouth, blurred vision, nausea, dizziness, constipation, nervousness, and urinary retention. Confusional states, impairment of recent memory, and hallucinations occur with use of high doses and in patients with advanced age and underlying dementia. Rarely, paralytic ileus, parotitis, hyperthermia, or skin rash occurs.

**Contraindications.** Children <3 yr; narrow-angle glaucoma; pyloric or duodenal obstruction; stenosing peptic ulcers; achalasia; bladder-neck obstructions; myasthenia gravis; cognitive disturbances.286

**Precautions.** Pregnancy; elderly patients. Use with caution in hot weather or during exercise and in patients with tachycardia, prostatic hypertrophy, open-angle glaucoma, or obstructive diseases of the GI tract.

**Drug Interactions.** Carefully observe patients given concomitant phenothiazines and/or heterocyclic antidepressants because intensification of mental symptoms, paralytic ileus, or hyperthermia can occur. Anticholinergics can decrease the effectiveness of phenothiazines. Use with amantadine can result in increased CNS anticholinergic effects. Anticholinergics can decrease digoxin absorption from digoxin tablets.

**Parameters to Monitor.** Intraocular pressure monitoring and gonioscope evaluation periodically. Monitor for Parkinson’s disease symptoms periodically.

**Notes.** Anticholinergic agents are considered useful for the initial treatment of parkinsonism in patients ≤60 yr with a rest tremor and without akinesia or rigidity.281,282 The drug does not alleviate the symptoms of tardive dyskinesia.

**Pharmacology.** Levodopa is centrally converted to dopamine by DOPA decarboxylase and replenishes dopamine, which is deficient in the basal ganglia of patients with Parkinson’s disease. Carbidopa, which does not cross the blood–brain barrier, inhibits peripheral DOPA decarboxylase, thereby increasing the amount of levodopa available to the brain for conversion to dopamine and limiting peripheral side effects. Addition of carbidopa decreases levodopa-induced nausea and vomiting but does not decrease adverse reactions caused by the central effects of levodopa.288

**Administration and Adult Dosage.** PO for Parkinson’s disease in patients not receiving levodopa (standard formulation) 25 mg carbidopa/100 mg levodopa tid initially, increasing in 1 tablet/day increments q 1–2 days to effective dosage, to a maximum of 8 tablets/day. Alternatively, 10 mg carbidopa/100 mg levodopa tid or qid initially, to a maximum of 8 tablets/day. Initial use of 10 mg carbidopa/100 mg levodopa can result in more nausea and vomiting because 70–100 mg/day of carbidopa is needed to saturate peripheral DOPA decarboxylase. If initial dosage maximum is reached with 10/100 tablets and further titration is necessary, substitute 25 mg carbidopa/250 mg levodopa tid or qid, increasing in 0.5–1 tablet/day increments q 1–2 days to effective dosage, to a maximum of
8 tablets/day. Some long-term users with advanced disease might need >2 g/day. SR Tab (patients already taking non-SR tablets) start with a dosage that provides 10% more levodopa daily. Initially, divide dosage bid or tid with an interval of 4–8 hr between doses while awake. Ultimately, dosages up to 30% greater might be needed, depending on patient response; (patients not receiving carbidopa/levodopa) 1 tablet bid initially, at least 6 hr apart, and allow 3 days between dosage adjustments. Usual dosage 2–8 tablets/day. If given in combination with a dopamine agonist or selegiline, lower dosages can be effective.

Special Populations. Pediatric Dosage. (<18 yr) safety and efficacy not established.

Geriatric Dosage. Same as adult dosage.

Dosage Forms. Tab 10 mg carbidopa/100 mg levodopa, 25 mg carbidopa/100 mg levodopa, 25 mg carbidopa/250 mg levodopa; SR Tab 25 mg carbidopa/100 mg levodopa, 50 mg carbidopa/200 mg levodopa. (See Notes.)

Patient Instructions. Stopping this medication suddenly can cause Parkinson’s disease to worsen quickly. Report bothersome or unexpected side effects. Unless prescribed, do not take levodopa in addition to this drug. Avoid pyridoxine (vitamin B₆) if you are taking levodopa alone, although it can be taken with carbidopa/levodopa. Avoid high-protein meals for maximum absorption. If you are taking the sustained-release tablet, swallow a whole or one-half tablet without chewing or crushing it. Onset of effect of the first morning dose of the sustained-release product could be delayed up to 1 hour compared with the quick-release product. A dark color (red, brown, or black) might appear in saliva, urine, or sweat and can stain clothing.

Missed Doses. Take this drug at regular intervals. If you miss a dose, take it as soon as you remember. If it is about time for the next dose, take that dose only. Do not double the dose or take extra.

Pharmacokinetics. Onset and Duration. Up to 50% of patients experience a reduction in efficacy after 5 yr. (See Notes.)

Fate. Carbidopa’s inhibition of peripheral levodopa decarboxylation doubles the oral bioavailability of levodopa and decreases its clearance by one-half. Dietary proteins compete with levodopa for intestinal absorption and decrease its effectiveness. (Rapid-release 50 mg carbidopa/200 mg levodopa) levodopa bioavailability is 99 ± 21%. A peak of 3.2 ± 1.1 mg/L occurs in 0.7 ± 0.3 hr. (SR 50 mg carbidopa/200 mg levodopa) levodopa bioavailability is 71 ± 24% and increases with food. A peak of 1.14 ± 0.42 mg/L occurs in 2.4 ± 1.2 hr. Levodopa Vₐ is 1.09 ± 0.59 L/kg; Cl is 0.28 ± 0.06 L/hr/kg; 90% of clearance is nonrenal. t₁/₂. (Carbidopa) 2.1 ± 0.6 hr; (levodopa alone) 1.4 ± 0.3 hr; (levodopa with carbidopa) 2 ± 1.3 hr.

Adverse Reactions. Anorexia, nausea, vomiting, and involuntary muscle movements (dyskinesias) occur frequently and are generally reversible with dosage reduction. Occasionally, mental changes, depression, dementia, palpitations, or orthostatic hypotensive episodes, increased libido, and bullous lesions occur. Rarely, psychosis, hemolytic anemia, leukopenia, or agranulocytosis is reported.
Compared with levodopa alone, carbidopa/levodopa has markedly reduced GI and cardiovascular side effects. However, mental disturbances are not eliminated and dyskinesias can appear earlier in therapy. These dyskinesias might require a decrease in dosage or dosage interval. Side effects can be more pronounced in patients receiving selegiline or a dopamine agonist as adjunctive therapy.

**Contraindications.** Lactation; nonselective MAO inhibitors concurrently or 2 weeks before carbidopa/levodopa; narrow-angle glaucoma; undiagnosed skin lesions; history of melanoma.

**Precautions.** Pregnancy. Use with caution in patients with histories of MI complicated by arrhythmias; peptic ulcer disease; severe cardiovascular, pulmonary, renal, hepatic, or endocrine disease; open-angle glaucoma; bronchial asthma; urinary retention; or psychosis. Also, use caution in patients receiving antihypertensives. Symptoms resembling neuroleptic malignant syndrome can occur when carbidopa/levodopa in combination with other antiparkinson agents is reduced abruptly or discontinued.

**Drug Interactions.** Iron salts, including low doses in multivitamins, can decrease levodopa absorption. Other agents for Parkinson’s disease, such as dopamine agonists, COMT inhibitors, and selegiline, can increase levodopa side effects when added to carbidopa/levodopa. Dosage of the levodopa product might need to be reduced by 10–30%. Metoclopramide and older neuroleptics (eg, chlorpromazine, haloperidol) have antidopaminergic effects and oppose the action of levodopa. Atypical neuroleptics have less antidopaminergic effect (clozapine has the least) but still can reduce the effectiveness of Parkinson’s disease therapy. Cholinergic agents such as tacrine, donepezil, and rivastigmine can worsen Parkinson’s symptoms by changing the dopamine–acetylcholine balance in the brain. Bupropion elicits a higher frequency of side effects in patients taking levodopa. Administer bupropion with caution, using small initial doses and small, gradual increases. There are rare reports of adverse reactions, including hypertension and dyskinesias, resulting from the concomitant use of TCAs and levodopa. Isoniazid, phenytoin, and papaverine can decrease the therapeutic effects of levodopa.

**Parameters to Monitor.** Monitor CBC, renal, cardiovascular, and liver functions periodically during long-term therapy. Monitor symptoms of Parkinson’s disease periodically. In patients with open-angle glaucoma, monitor intraocular pressure.

**Notes.** Levodopa produces sustained improvement in rigidity and bradykinesia in 50–60% of patients. Tremor is variably affected, and postural stability is unresponsive. Loss of therapeutic effect is manifested by fluctuations in motor performance. Patients can experience periods of lack of drug effect (“off” periods) alternating with periods of therapeutic efficacy (“on” periods). Response can be predictable, where the effect fades before the next dose (“wearing-off” or “end-of-dose”), or unpredictable (“yo-yo”), where there is no relation to the time of dose. SR carbidopa/levodopa reduces “off” time an average of 30–40 min/day and allows a mean 33% reduction in the frequency of administration; the lower bioavailability of the SR product necessitates a 25% median increase in the daily dosage of levodopa compared with non-SR carbidopa/levodopa. With disease progression, adjunctive therapy with an MAO-B inhibitor, a dopamine agonist, or...
a COMT inhibitor (eg, tolcapone, entacapone) might be required to decrease the frequency of fluctuations caused by dyskinesia or dystonia.

**DONEPEZIL**

**Pharmacology.** Donepezil enhances the action of acetylcholine by reversibly inhibiting acetylcholinesterase (AChE), the enzyme responsible for its hydrolysis. It has a high degree of selectivity for AChE in the CNS, which might explain the relative lack of peripheral side effects. Donepezil is indicated for the treatment of mild to moderate dementia of the Alzheimer’s type. No evidence suggests that donepezil alters the course of the disease.

**Administration and Adult Dosage.** PO for Alzheimer’s disease 5 mg once daily at bedtime, with or without food. The 10 mg dose is associated with a higher frequency of side effects but can provide extra benefit in some individuals. If a 10 mg dose is desired, first allow 4–6 weeks at 5 mg/day.

**Special Populations.** Geriatric Dosage. Same as adult dosage.

**Other Conditions.** No dosage adjustment is necessary in patients with renal or hepatic disease.

**Dosage Forms.** Tab 5, 10 mg.

**Patient Instructions.** This drug can be taken with or without food. Side effects can occur when you first start taking donepezil, but these frequently subside after 1 to 2 weeks. The maximum benefits of the drug might not occur until 4 to 8 weeks after starting the drug. Because there is variability in the way patients respond to donepezil, decide with your doctor how long to take donepezil. Do not abruptly discontinue donepezil on your own.

**Missed Doses.** Take this drug at regular intervals. If you miss a dose, take it as soon as you remember. If it is about time for the next dose, take that dose only. Do not double the dose or take extra.

**Pharmacokinetics.** **Onset and Duration.** Onset is in about 3 weeks, with maximum benefits occurring in 4–8 weeks.\(^{295}\) The manufacturer recommends waiting 3 months before evaluating the full effects of the drug.\(^{296}\)

**Fate.** Donepezil is completely absorbed and is 96% protein bound. \(V_{dss}\) is 12 L/kg; \(Cl\) is 0.13 L/hr/kg. About 60% is eliminated as hepatic metabolites including products of CYP2D6 and CYP3A4 and glucuronides. About 17% is excreted unchanged in urine.\(^{297,298}\)

\(t_{1/2}\) >70 hr.

**Adverse Reactions.** Occasionally, nausea, vomiting, muscle cramps, fatigue, anorexia, and headache occur.

**Contraindications.** Hypersensitivity to donepezil or piperidine derivatives (eg, biperiden, bupivacaine, methylphenidate, paroxetine, rifabutin, trihexyphenidyl).

**Precautions.** Use with caution in peptic ulcer disease, syncope, sick sinus syndrome, bradycardia, altered supraventricular cardiac conduction, asthma, seizures, or COPD.
Drug Interactions. There are few in vivo studies. In vitro, ketoconazole and quinidine decrease donepezil metabolism; enzyme inducers might increase its metabolism. Although extensively bound to plasma proteins, donepezil does not interact with warfarin or furosemide or with cimetidine or digoxin. Donepezil can increase the risk of GI side effects from NSAIDs because of the possible increase in stomach acid production.

Parameters to Monitor. Monitor mental status and improvements in activities of daily living initially and then periodically during therapy.

Notes. Because Alzheimer’s disease is a neurodegenerative disorder, patients might improve or show no change in their cognitive functions.

DOPAMINE AGONISTS:

<table>
<thead>
<tr>
<th>Bromocriptine</th>
<th>Parlodel</th>
</tr>
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<tbody>
<tr>
<td>Pergolide</td>
<td>Permax</td>
</tr>
<tr>
<td>Pramipexole</td>
<td>Mirapex</td>
</tr>
<tr>
<td>Ropinirole</td>
<td>Requip</td>
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</table>

Pharmacology. Bromocriptine and pergolide are ergot-derived dopamine agonists that stimulate dopamine-D<sub>2</sub> receptors; in addition, pergolide stimulates and bromocriptine partly antagonizes D<sub>1</sub> receptors. Pramipexole and ropinirole are non–ergot-derived dopamine subtype selective agonists that exert activity in the CNS at D<sub>2</sub> and D<sub>3</sub> receptors but have no activity at the D<sub>1</sub> receptor.<sup>299-301</sup> D<sub>2</sub> receptors are thought to play an important role in improving the akinesia, bradykinesia, rigidity, and gait disturbances of Parkinson’s disease. Pramipexole, unlike other dopamine agonists, binds with 7-fold greater affinity to D<sub>3</sub> receptors than to D<sub>2</sub> receptors and can affect mood. Although bromocriptine also can inhibit prolactin secretion, it is no longer indicated for the prevention of postpartum lactation. Other uses of bromocriptine are the treatment of acromegaly, prolactin-secreting pituitary adenomas, and amenorrhea/galactorrhea secondary to hyperprolactinemia without a primary tumor. (See Dopamine Agonists Comparison Chart.)

Administration and Adult Dosage. (Bromocriptine) PO for acromegaly 1.25 mg/day or bid with food initially and then increasing in 1.25–2.5 mg/day increments q 3–7 days to a usual maintenance dosage of 2.5 mg bid–tid. Maximum dosage is 100 mg/day. PO for hyperprolactinemia 2.5 mg tid. PO for Parkinson’s disease (see Dopamine Agonists Comparison Chart).

Special Populations. Pediatric Dosage. Safety and efficacy not established. PO for treatment of prolactin-secreting pituitary adenomas (≥11 yr) 1.25–2.5 mg daily; increase in 2.5 mg/day increments q 2–7 days as tolerated until therapeutic response is achieved.

Geriatric Dosage. Same as adult dosage.

Other Conditions. (Pramipexole) adjust for renal impairment as follows: Cl<sub>e</sub> 35–59 mL/min, give 1.5 mg bid initially, to a maximum of 1.5 mg bid; Cl<sub>e</sub> 15–34 mL/min, give 0.125 mg/day initially, to a maximum of 1.5 mg/day.
Dosage Forms. (See Dopamine Agonists Comparison Chart.)

Patient Instructions. This medication might improve the symptoms of Parkinson’s disease but will not cure it. Take this drug with food to minimize stomach upset. This drug can cause dizziness, drowsiness, or fainting, especially after the first dose. Until the extent of these effects is known, use caution when driving, operating machinery, or performing tasks requiring mental alertness. Mental disturbances, including vivid dreams, confusion, and paranoid delusions, can occur even with low doses, especially when added to levodopa therapy. Avoid concurrent use of alcohol. Inform your physician and pharmacist of any other prescription or over-the-counter medications you might be taking because these can interact with your antiparkinsonian medications. Do not abruptly stop taking this medication or change your dosage without medical supervision. (Bromocriptine) women taking this drug to induce ovulation should use a barrier contraceptive. (Pramipexole and ropinirole) some patients have reported sudden excessive drowsiness, causing them to fall asleep during activities of daily living, including driving. Notify your doctor immediately if you notice significant daytime drowsiness. Avoid use of other sedating medications.

Missed Doses. Take this drug at regular intervals. If you miss a dose, take it as soon as you remember. If it is about time for the next dose, take that dose only. Do not double the dose or take extra.

Pharmacokinetics. Onset and Duration. Onset (bromocriptine) 0.5–1 hr;302,303 (pergolide and pramipexole) 1–2 hr; (ropinirole) 1–2 hr on an empty stomach, 3–4 hr with food.304 Duration (bromocriptine, ropinirole) 3–6 hr; (pergolide, pramipexole) 8–12 hr.305 (Bromocriptine in amenorrhea) normal menstrual function usually returns within 6–8 weeks.

Fate. (Bromocriptine) bioavailability is about 28%; it is 90% bound to plasma proteins. Peak serum concentrations occur in 1.2 ± 0.4 hr, and detectable concentrations are found for up to 12 hr after discontinuation of drug.302 Cl is 4.4 ± 2.6 L/hr/kg.303 The majority (98%) is excreted in the feces via bile.302 (Pergolide) bioavailability is about 60%; it is 90% bound to plasma proteins. Approximately 40–50% of a dose is excreted in feces over 7 days as at least 10 metabolites.306 (Pramipexole) bioavailability is about 90%; it is 15% bound to plasma proteins. Vd is 7 L/kg; renal Cl is about 0.4 L/hr/kg and markedly exceeds the GFR. About 90% of a dose is excreted unchanged in urine. (Ropinirole) although completely absorbed, bioavailability is 55% because of first-pass metabolism. It is 36% bound to plasma proteins and undergoes extensive metabolism in the liver to inactive metabolites. The N-despropyl metabolite is the major metabolite; the drug also is hydroxylated and glucuronidated.

Adverse Reactions. Nausea, headache, hallucinations, dyskinesias, somnolence, vomiting, symptomatic hypotension, dizziness, fatigue, constipation, and light-headedness occur frequently. Occasionally, abdominal cramps and diarrhea occur. (Bromocriptine, pergolide) rarely, hypertension, stroke, seizures, rhinorrhea, or erythromelalgia are reported. Pleuropulmonary disease is rare and usually occurs in men, especially in smokers receiving 20–100 mg/day of bromocriptine for 3–6 months; it presents with dyspnea and improves with drug discontinuation.307
Contraindications. (Bromocriptine, pergolide) pregnancy; lactation; uncontrolled hypertension; pre-eclampsia; concurrent use of other ergot alkaloids; hypersensitivity to ergot alkaloids.

Precautions. (Bromocriptine, pergolide) use with caution in patients with symptoms of peptic ulcer disease, history of pulmonary disease, MI, liver disease, severe angina, or psychiatric disease. (Bromocriptine) use a barrier contraceptive during treatment for amenorrhea, galactorrhea, or infertility. If pregnancy is detected, discontinue the drug. (Prampexole, ropinirole) several patients have reported “sleep attacks,” or falling asleep during activities of daily living. These sudden occasions of sleepiness have resulted in motor vehicle accidents. Advise patients of this possibility and assess them regularly for symptoms of drowsiness. Instruct patients to avoid other sedating medications and drugs that can increase blood levels of these agents (eg, cimetidine with prampexole, ciprofloxacin with ropinirole). Sudden episodes of falling asleep might necessitate discontinuation of the dopamine agonist. If prampexole or ropinirole is continued after such an incident, instruct the patient not to drive or use dangerous machinery.

Drug Interactions. When used with carbidopa/levodopa, it might be necessary to reduce the dosage of levodopa by as much as 30% to reduce the potential for developing dyskinesias. Drugs that antagonize dopamine (eg, phenothiazines, butyrophenones, metoclopramide) can reduce the effectiveness of these drugs. (Bromocriptine) erythromycin can increase bromocriptine serum levels. (Bromocriptine, pergolide) other ergot alkaloids can exacerbate cardiotoxic effects. (Prampexole) prampexole can result in an earlier and higher peak serum level of levodopa. Drugs that interfere with renal tubular secretion of cations (eg, cimetidine, ranitidine, verapamil, quinidine) can decrease prampexole renal elimination. (Ropinirole) ropinirole is metabolized by CYP1A2; therefore, it might interact with inhibitors or inducers of this isozyme; ciprofloxacin markedly increases AUC and peak serum concentrations. Ropinirole might require dosage reduction with co-administration of estrogen.

Parameters to Monitor. Monitor blood pressure frequently during the first few days of therapy and periodically thereafter. Periodically evaluate hepatic, hematopoietic, cardiovascular, and renal function during long-term therapy. Monitor symptoms of Parkinson’s disease periodically.

Notes. These drugs can be used as single agents for the treatment of early Parkinson’s disease and as adjunctive agents in moderate- to late-stage disease. As first-line agents, dopamine agonists can offer neuroprotection by regulating dopamine turnover and delaying the introduction of levodopa. However, as single agents, they are less effective than levodopa.

ENTACAPONE

Pharmacology. Entacapone is a peripheral acting, selective, and reversible inhibitor of COMT, similar in mechanism to tolcapone. Entacapone is indicated as an adjunct to levodopa/carbidopa to treat patients with Parkinson’s disease who experience end-of-dose “wearing-off.”

Administration and Adult Dosage. PO for Parkinson’s disease 200 mg, taken with each levodopa/carbidopa dose, to a maximum of 8 times daily (1600 mg/day).
The 200 mg dose is optimal and is more efficacious than higher doses, possibly because of interference with carbidopa absorption at doses ≥400 mg.313

**Special Populations.** *Geriatric Dosage.* Same as adult dosage.

**Dosage Forms.** Tab 200 mg.

**Patient Instructions.** Take one tablet of entacapone with each dose of levodopa/carbidopa. Be aware of the possibility of developing dizziness and hypotension when rising from a sitting or supine position. This effect is more likely to occur when the drug is first started. Nausea is another potential side effect in early therapy. Entacapone can cause a brownish-orange discoloration of the urine that is harmless. Dyskinesias and hallucinations can occur with entacapone, which can necessitate the reduction of the carbidopa/levodopa dose. Do not drive a car or operate machinery until you know how entacapone will affect your mental alertness or motor abilities.

**Missed Doses.** Take this drug at regular intervals. If you miss a dose, take it as soon as you remember. If it is close to the time of the next dose, take that dose only. Do not double the dose or take extra.

**Pharmacokinetics. Onset and Duration.** Onset is rapid and occurs with first dose. Peak effect is 0.7–1.3 hr after oral administration. Entacapone can prolong the effects of a carbidopa/levodopa dose by about 30 min.313

**Fate.** Oral bioavailability 35%. Food does not affect entacapone pharmacokinetics, but bioavailability is doubled with liver cirrhosis. Peak after a single 200 mg dose is approximately 1.2 μg/mL. Plasma binding is 98%, mainly to serum albumin. Entacapone does not distribute widely into tissues or CNS; $V_d$ is 0.4 ± 0.16 L/kg. CI is 0.6 ± 0.1 L/kg/hr. Entacapone is metabolized almost completely before elimination, mainly by isomerization followed by glucuronidation. Metabolites are eliminated primarily by biliary excretion, with 90% of the metabolized dose found in the feces and 10% in urine. Only about 0.2% of the dose is eliminated unchanged in urine.314

$t_{1/2}$, β phase 0.4-0.7 hr; γ phase 2.4 hr, which accounts for about 10% of the total AUC.

**Adverse Reactions.** Orthostatic hypotension, diarrhea, dyskinesias, and hallucinations can occur with entacapone therapy, especially during the initial days of therapy. Dopaminergic side effects, including dyskinesias, nausea, dizziness, hallucinations, and insomnia, can occur. Dyskinesias are the most common side effect, usually early in therapy. Their frequency is reduced with lowering of the levodopa/carbidopa dose. Diarrhea is a frequent side effect that is mild to moderate in 4–10% of patients but severe in 1.3%. Orthostatic hypotension, urine discoloration, abdominal pain, and constipation occur occasionally. Elevation of liver enzymes was the same as that with placebo (0.8%). Rarely, rhabdomyolysis, hyperpyrexia and confusion, resembling neuroleptic malignant syndrome, occur.

**Contraindications.** Concurrent use with nonselective MAOIs but it can be taken with a selective MAO-B inhibitor (eg, selegiline).

**Precautions.** Because 90% of drug elimination is by biliary excretion, use caution in patients with biliary obstruction.
Drug Interactions. Entacapone does not inhibit cytochrome P450 enzymes at doses used for Parkinson’s disease. Despite its extensive protein binding, in vitro studies have not shown binding displacement between entacapone and other highly bound drugs such as warfarin, salicylic acid, and diazepam. Drugs that interfere with biliary excretion, glucuronidation, and intestinal β-glucuronidase, such as probenecid, cholestyramine, erythromycin, ampicillin, rifampin, and chlorphenicol, have the potential to interfere with entacapone elimination. Drugs that are metabolized by COMT, such as methyl dopa, dobutamine, isoproterenol, and epinephrine, can have enhanced effects when given with entacapone.

Parameters to Monitor. Monitor symptoms of Parkinson’s disease and excessive dopaminergic activity. The dose of carbidopa/levodopa might need to be reduced if side effects such as dyskinesias and hallucinations are excessive or intolerable.

Pharmacology. Galantamine is a competitive, reversible acetylcholinesterase inhibitor similar to donepezil and rivastigmine.315,316

Administration and Adult Dosage. PO for Alzheimer’s disease 4 mg bid initially with a meal, increasing in 4 mg bid increments at 4-week intervals to a maximum of 12 mg bid. In moderate hepatic or renal dysfunction the maximum dosage is 8 mg bid. Avoid in severe hepatic (Child-Pugh 10–15) or renal (Clcr <9 mL/min) impairment. If more than a few days of therapy are missed, resume therapy at 4 mg bid.

Dosage Forms. Tab 4, 8, 12 mg.

Pharmacokinetics. Oral bioavailability is ≥90%; food decreases peak concentration and rate, but not extent of absorption. Peak cholinesterase inhibition occurs 1 hr after a dose. It is 18% plasma protein bound and distributed extensively into RBCs. \( V_{\text{dss}} \) is 2.6 L/kg; Cl is 0.34 L/hr/kg. Metabolism is primarily by CYP2D6 and 3A4. About 20–25% is excreted unchanged in urine in 24 hr. Half-life is 5–7 hr.315

Adverse Reactions. GI side effects (eg, nausea, vomiting, diarrhea, anorexia, weight loss), are most prominent during dosage escalation. Dizziness, headache, chest pain, tremor, depression, rhinitis, urinary incontinence, flatulence and bradycardia also occur frequently. Various cardiac arrhythmias, increased alkaline phosphatase, thrombocytopenia, GI bleeding, hyperglycemia, and psychiatric symptoms occur occasionally. Esophageal perforation has been reported.

RILUZOLE Rilutek

Pharmacology. In the treatment of amyotrophic lateral sclerosis, riluzole is hypothesized to protect motor neurons from degeneration and death. Although the exact mechanism of action is unknown, there are three pharmacologic properties of the drug that are thought to be relevant: inhibition of glutamate release, inactivation of voltage-dependent sodium channels, and interference with intracellular events that follow activation of excitatory amino acid receptors.

Administration and Adult Dosage. PO for amyotrophic lateral sclerosis 50 mg bid.

Special Populations. Geriatric Dosage. Same as adult dosage.
**Dosage Forms.** Tab 50 mg.

**Patient Instructions.** Take this medication at the same time each day on an empty stomach. It can cause dizziness, drowsiness, or vertigo. Until the extent of these effects is known, use caution when driving, operating machinery, or performing tasks requiring mental alertness. Do not drink alcohol while taking this medication. Contact your doctor if fever or flulike symptoms occur. Protect the drug from exposure to light.

**Missed Doses.** Take this drug at regular intervals. If you miss a dose, take it as soon as you remember. If it is about time for the next dose, take that dose only. Do not double the dose or take extra.

**Pharmacokinetics.** *Fate.* Riluzole is rapidly absorbed, with about 90% absorbed, but absolute bioavailability is 60 ± 9% because of a first-pass effect. Peak serum concentrations occur within 60–90 min. A high-fat meal decreases the AUC by about 20%. About 96% is bound to serum proteins, mainly albumin and lipoproteins; it is distributed extensively throughout the body, with a Vₐ of about 3.4 L/kg; Cl is 0.7 L/hr/kg in white males. Riluzole is metabolized extensively in the liver to at least six major metabolites, mainly by CYP1A2 (hydroxylated derivatives) and P450-dependent glucuronidation. Its metabolism is slower by 32% in women and by 50% in Japanese subjects native to Japan than in white males. Glucuronides account for about 85% of urine metabolites.

\[ t_{1/2} = 12 \pm 1.8 \text{ hr.} \]

**Adverse Reactions.** Nausea, constipation, vomiting, abdominal pain, elevations in AST and ALT, asthenia, dizziness, diarrhea, vertigo, and circumoral paresthesia occur frequently. Decreased lung function, pneumonia, somnolence and neutropenia occur occasionally (3 cases of neutropenia in 4000 during clinical trials).

**Precautions.** Use with caution in patients with renal or hepatic disease, especially the elderly.

**Drug Interactions.** Riluzole might interact with other drugs that also are metabolized by CYP1A2, including theophylline, caffeine, fluoroquinolones, and amitriptyline. Enzyme inducers and cigarette smoking increase the metabolism of riluzole.

**Parameters to Monitor.** Monitor liver function tests and CBC periodically as well as response to therapy.

**Notes.** In amyotrophic lateral sclerosis, riluzole prolongs the time to tracheostomy or death (a 21% risk reduction) compared with placebo. The difference in rates of muscle deterioration between riluzole and placebo was significant in some studies but not in others.

**RIVASTIGMINE (Exelon)**

**Pharmacology.** Rivastigmine is an intermediate-acting (pseudo-irreversible) AChE inhibitor that binds to AChE, resulting in a carbamated form of AChE that cannot hydrolyze acetylcholine. This action increases CNS acetylcholine activity. It is indicated for treatment of mild to moderate symptoms of dementia of the Alzheimer’s type.
Administration and Adult Dosage. **PO** for Alzheimer’s disease 1.5 mg bid initially. After at least 2 weeks, increase in 1.5 mg bid increments q 2 weeks as tolerated, to a maximum of 6 mg bid. If more than a few days of therapy are missed, resume therapy at the initial dose of 1.5 mg bid.

**Special Populations. Geriatric Dosage.** No specific dosage adjustment is needed because the dose is adjusted to patient tolerance.

**Dosage Forms.** Cap 1.5, 3, 4.5, 6 mg; Soln 2 mg/mL.

**Patient Instructions.** Take this drug with food in the morning and evening. Dosage will be increased about every 2 weeks until the maximum tolerable dose is reached. If you experience adverse effects, such as loss of appetite, nausea, vomiting or abdominal pain, stop treatment for several doses and then resume at the same or next lower dose level. Inform your physician if these symptoms occur.

**Missed Doses.** Take this drug at regular intervals. If you miss a dose, take it as soon as you remember. If it is within a few hours of the next dose, take that dose only. Do not double the dose or take extra. If you miss more than a few doses, do not resume the same dosage. Inform your physician.

**Pharmacokinetics. Onset and Duration.** After 6 mg, anticholinesterase activity is present in the CSF for about 10 hr, with a maximum inhibition of about 60% 5 hr after a dose.

**Fate.** Rivastigmine is rapidly and completely absorbed with bioavailability of about 36% after a 3 mg dose. Peak plasma concentrations occur within 1 hr, with peak CSF concentrations achieved in 1.4–2.6 hr. Food delays plasma peak time by 90 min, lowers the peak by 30%, and increases bioavailability by 30%. Rivastigmine is 40% bound to plasma proteins; \( V_d \) is 1.8–2.7 L/kg. \( CL \) is 108 ± 36 L/hr after 6 mg bid. Pharmacokinetics are nonlinear at doses above 3 mg bid. Metabolism is mainly by cholinesterase-mediated hydrolysis. It is then eliminated renally, with 97% of the dose detected in the urine as metabolites, most commonly as the sulfate conjugate of the decarbamylated metabolite (40%). \( CL \) is reduced in the elderly (by 30%) and in patients with renal (by 64%) or hepatic (by 60%) disease. Only 0.4% is eliminated in the feces.

\( t_{1/2} \). 1.5 hr.

**Adverse Reactions.** Nausea (47%) and vomiting (31%) are frequent occurrences in patients, especially women, treated with 6–12 mg/day and are more likely during the titration phase rather than the maintenance phase. Anorexia and weight loss also occur more frequently in women. Other side effects are dizziness, headache, tremor, abdominal pain, dyspepsia, hypotension, orthostatic hypotension, insomnia, tinnitus, palpitations, confusion, anemia, and rash. \(^{322,324} \) Resumption of a high dose after a few days without taking the drug can result in severe vomiting and esophageal perforation.

**Contraindications.** Sensitivity to carbamate derivatives.

**Precautions.** See Donepezil.

**Drug Interactions.** Excessive cholinergic effect can occur if rivastigmine is given with cholinergic drugs (eg, succinylcholine, bethanechol). Rivastigmine can antagonize the effects of anticholinergic drugs and antiparkinson’s drugs. Based on
in vitro studies, rivastigmine does not interact with digoxin, warfarin, diazepam, or fluoxetine. Rivastigmine pharmacokinetics are not altered by antacids, antihypertensives, calcium channel blockers, antidiabetics, NSAIDs, salicylates, antianginals, antihistamines, estrogens, or β-blockers. Rivastigmine can increase the risk of GI side effects from NSAIDs due to the possible increase in stomach acid production.

Parameters to Monitor. Because of the high frequency of nausea, vomiting, and anorexia, monitor patients for these reactions and for possible weight loss.

Notes. Metrifonate (ProMem—Bayer) is an organophosphate being studied for Alzheimer’s disease.

SELEGILINE HYDROCHLORIDE

Pharmacology. Selegiline (formerly L-deprenyl) is a selective, irreversible MAO-B inhibitor that is used as adjunctive therapy in the management of Parkinson’s disease. MAO-B is found in the brain and plays a role in the catabolism of dopamine.325,326 By preventing the breakdown of dopamine by MAO-B, selegiline increases the net amount of dopamine available in the brain. Selegiline also can exert a protective effect by preventing the accumulation of neurotoxic free radicals generated by dopamine metabolism.327–331 However, the exact mechanism of the beneficial effects of selegiline is unclear and might be symptomatic, neuroprotective, or both.

Administration and Adult Dosage. PO for Parkinson’s disease 5 mg bid taken at breakfast and lunch. Alternatively, give an initial dosage of 2.5 mg/day and slowly increase to 10 mg/day over several weeks to minimize side effects.325 There is no evidence that dosages >10 mg/day increase efficacy, and they can lead to nonspecific inhibition of MAO-A.

Special Populations. Pediatric Dosage. Safety and efficacy not established.

Geriatric Dosage. Same as adult dosage.

Dosage Forms. Cap 5 mg; Tab 5 mg.

Patient Instructions. Take this medication with morning and midday meals to minimize nausea and night-time insomnia. At a dosage of 10 mg/day or less, tyramine-containing foods and medications containing amines are safe to consume. Initiation of selegiline might require a reduction of carbidopa/levodopa dosage. Report immediately any severe headache or other unusual or unexpected symptoms.

Missed Doses. Take this drug at regular intervals. If you miss a dose, take it as soon as you remember. If it is about time for the next dose, take that dose only. Do not double the dose or take extra.

Pharmacokinetics. Onset and Duration. Recovery of platelet MAO-B activity after a single oral dose is 2–4 days; after long-term treatment, >90% of platelet MAO-B remains inhibited after 5 days.332 With continual use, clinical efficacy lasts 6–12 months in most patients, to a maximum of 12–24 months.327–330

Fate. Selegiline is readily absorbed from the GI tract, with a peak at 0.5–2 hr; 94% is bound to plasma proteins.332 It is metabolized by the liver to N-desmethylselegi-
line, L-amphetamine, and L-methamphetamine; these isomers, however, are only 10% as potent as the D-isomers. After long-term therapy with 10 mg/day in 2 divided doses, mean trough serum levels of selegiline and N-desmethylselegiline are undetectable; L-amphetamine is 5.9 ± 2.7 µg/L (22 ± 10 nmol/L), and L-methamphetamine is 14.9 ± 6.8 µg/L (100 ± 45 nmol/L). The concentrations of these metabolites are probably too low to contribute to the drug’s clinical efficacy but can contribute to adverse effects. About 86% is excreted in urine as inactive metabolites.325,326

\[ t_{1/2} \text{ (N-desmethylselegiline)} = 2 ± 1.2 \text{ hr; (L-amphetamine)} = 17.7 ± 16.3 \text{ hr; (L-methamphetamine)} = 20.5 ± 11.4 \text{ hr}. \]

**Adverse Reactions.** Nausea, abdominal pain, dry mouth, confusion, hallucinations, dizziness, insomnia, lightheadedness, and/or fainting occur frequently. Vivid dreams, dyskinesias, and headache occur occasionally. In decreasing order of frequency—nausea, hallucinations, confusion, depression, loss of balance, and insomnia—can lead to discontinuation of the drug. Mild, asymptomatic elevations in liver function tests can occur.

**Precautions.** Pregnancy; lactation. Concurrent use with meperidine. Do not use at dosages exceeding 10 mg/day.

**Drug Interactions.** Concurrent administration of selegiline and serotonin reuptake inhibitors (eg, SSRIs, nefazodone, venlafaxine) can cause serotonin syndrome; do not give them within 1–2 weeks of each other (5 weeks after stopping fluoxetine).

**Parameters to Monitor.** Evaluate cardiovascular status and monitor liver function tests periodically. Monitor Parkinson’s disease symptoms periodically.

**Notes.** Selegiline is indicated as adjunctive treatment with carbidopa/levodopa in Parkinson’s disease. Although the efficacy of selegiline is not superior to that of other adjunctive drugs such as dopamine agonists, it appears to be better tolerated.299,300 Approximately 60% of patients who receive selegiline experience a modest (<10%) reduction in “off” periods and can reduce their levodopa dosage by 20%.329 A role for the drug as an initial agent in patients with mild disease is supported by results of a study comparing selegiline 10 mg/day with placebo in patients with early (<5 yr) untreated Parkinson’s disease; selegiline delayed the onset of disease-related disability by nearly 1 yr.327 Zelep (Athena) is a rapidly dissolving form of selegiline in phase III clinical trials.

**TOLCAPONE**

**Tasmar**

**Pharmacology.** Tolcapone reversibly inhibits at least 80% of the activity of COMT. This action prevents the metabolism of levodopa to 3-O-methyldopa and thus prolongs its duration of action, especially with co-administration of carbidopa.333,334 Tolcapone increases plasma levodopa bioavailability by about 2-fold and variably prolongs the terminal half-life of levodopa (given with carbidopa) in the elderly from 2 hr to as long as 3.5 hr, but has no effect on the peak serum levels of levodopa or the time at which they occur. (See Notes.)

**Administration and Adult Dosage.** PO for Parkinson’s disease 100 mg tid initially, always as an adjunct to levodopa/carbidopa therapy, increasing to a maxi-
mum of 200 mg tid. If the patient shows no substantial clinical benefit after 3
weeks of therapy, discontinue tolcapone. (See Parameters to Monitor.)

Special Populations. Geriatric Dosage. Same as adult dosage.

Dosage Forms. Tab 100, 200 mg.

Patient Instructions. This drug can lower blood pressure and cause unsteadiness,
a nausea, or sweating initially. Do not rise rapidly after sitting or lying down. It also
can cause drowsiness. Until the extent of these effects is known, use caution when
driving, operating machinery, or performing tasks that require mental alertness.
This drug also can worsen dyskinesias or dystonia when it is first started and
might require adjustment of the amount of carbidopa/levodopa you are taking. Be-
cause of the risk of liver damage with this drug, you will require regular liver en-
zyme tests. Notify your physician immediately if signs of liver toxicity develop.

Missed Doses. Take this drug at regular intervals. If you miss a dose, take it as
soon as you remember. If it is about time for the next dose, take that dose only. Do
not double the dose or take extra.

Pharmacokinetics. Onset and Duration. Onset 1–2 hr; duration 12–24 hr.

Fate. About 65% is orally absorbed. A peak serum concentration of 6.3 ±
2.9 mg/L occurs 1.8 ± 1.3 hr after a 200 mg dose. Food increases the peak time
and decreases the peak and AUC of tolcapone. About 99.9% is bound to plasma
proteins, mainly albumin. The predominant metabolic pathway is glucuronidation.
Major metabolites that are a product of oxidative processes include 3-O-methyl-
tolcapone and carboxylic acid derivatives metabolized by CYP2A6 and CYP3A4.
Those that are a product of reductive processes include amines and N-acetyl deriv-
atives. Cl is about 0.1 L/kg/hr. About 40% of an orally administered dose is ex-
creted in the urine and feces in 24 hr and >95% in 7–9 days. Less than 0.5% of tol-
capone is excreted unchanged in urine.333,334

\[ t_{1/2} \] (Tolcapone) 2 ± 0.8 hr; (3-O-methyltolcapone) 32 ± 7 hr.333,334

Adverse Reactions. Tolcapone has caused several cases of severe, fulminant liver
failure that were sometimes fatal. Monitor liver function carefully. Adverse reac-
tions consistent with increased levodopa exposure include worsening dyskinesia,
nausea, sleep disorders, dystonia, somnolence, anorexia, hallucinations, and pos-
tural hypotension. They might be lessened by reducing levodopa dosage. Urine
discoloration also occurs and is attributable to the yellow color of tolcapone and
its metabolites. Other tolcapone-related side effects include headache, abdominal
pain, and diarrhea. The most common reason for drug discontinuation from clin-
ical studies was severe diarrhea in 3% of patients. The onset is often delayed and
usually occurs within 0.5–3 months after initiation of therapy.

Contraindications. Liver disease or in patients who have discontinued tolcapone
therapy because of liver toxicity; history of rhabdomyolysis caused by any med-
ication; history of hyperpyrexia and confusion related to medication use or to
medication discontinuation.

Precautions. Orthostatic hypotension, hallucinations, diarrhea, or dyskinesias
can occur at the initiation of therapy. Use with caution in patients with renal im-
pairment. Follow recommended plan for monitoring liver enzymes. Discontinue
tolcapone if ALT or AST exceeds the upper limit of normal or if the patient develops signs and symptoms of liver failure (jaundice, anorexia, dark urine, pruritus, nausea, and right upper quadrant tenderness). The patient should sign a consent form before therapy is initiated.

**Drug Interactions.** Tolcapone can inhibit the metabolism of other drugs also metabolized by COMT (eg, dobutamine, isoproterenol) and exacerbate the dopaminergic side effects of other antiparkinsonian agents. It does not interact with ephedrine or desipramine (a substrate for CYP2D6) but has in vivo affinity for CYP2C9 (although the clinical relevance is undetermined).

**Parameters to Monitor.** Before starting treatment, conduct tests to exclude the presence of liver disease. Obtain ALT and AST levels at baseline and then q 2 weeks for the first year of therapy, q 4 weeks for the next 6 months, and q 8 weeks thereafter. If the dose is increased, begin liver enzyme monitoring again as when the drug was initiated.

**Notes.** Tolcapone can be administered on a schedule (3 times daily) and does not need to be administered with each dose of levodopa. The increase in “on time” in nonfluctuating and fluctuating patients was 1.7–2.9 hr over baseline (vs 0.7–1 hr for placebo). In clinical trials, the average decrease in daily levodopa dosage was about 30% in about 70% of patients.

**TRIHEXYPHENIDYL HYDROCHLORIDE**

**Pharmacology.** Trihexyphenidyl is a competitive antagonist of acetylcholine at central muscarinic receptors. In Parkinson’s disease, it is an adjunctive treatment that balances cholinergic and dopaminergic activities in cerebral synapses.

**Administration and Adult Dosage.** PO for Parkinson’s disease 1 mg/day initially, increasing in 2 mg/day increments q 3–5 days, to a maximum of 12–15 mg/day. **Usual maintenance dosage** 6–10 mg/day in 3 divided doses or 3–6 mg/day in 3 divided doses concurrent with levodopa. SR caps can be given bid once the maintenance dosage has been determined. **PO for drug-induced extrapyramidal disorders** 1 mg initially, increasing in 1 mg increments every few hours until symptoms are controlled, usually 5–15 mg/day in 3–4 divided doses.

**Dosage Forms.** Elxr 0.4 mg/mL; Tab 2, 5 mg; SR Cap 5 mg.

**Pharmacokinetics.** The onset of action is within 1 hr, and the peak effect lasts 2–3 hr; the duration of action is 6–12 hr. The majority of the drug is excreted in the urine probably unchanged, and the elimination half-life is 10.2 ± 4.7 hr.

**Adverse Reactions.** Adverse reactions, precautions, contraindications, and drug interactions are the same as those for benztropine. When trihexyphenidyl is used concurrently with levodopa, the dosages of both drugs might require reduction.

**Precautions.** Use with great caution in patients older than 65 yr because they are more sensitive to the effects of anticholinergic agents.
### DOPAMINE AGONISTS COMPARISON CHART

<table>
<thead>
<tr>
<th>DRUG</th>
<th>DOSAGE FORMS</th>
<th>ADULT DOSAGE</th>
<th>DOPAMINE RECEPTOR SELECTIVITY</th>
<th>EFFECT ON CYTOCHROME P450 ISOZYMES</th>
<th>HALF-LIFE (HR)</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Bromocriptine</strong>&lt;sup&gt;a&lt;/sup&gt;</td>
<td>Cap 5 mg, Tab 2.5 mg.</td>
<td>PO 1.25 mg bid initially, increasing in 2.5 mg/day increments q 2 weeks to a usual dosage of 10–40 mg/day.</td>
<td>D&lt;sub&gt;1&lt;/sub&gt;: +/+  D&lt;sub&gt;2&lt;/sub&gt;: ++  D&lt;sub&gt;3&lt;/sub&gt;: +</td>
<td>Inhibits CYP3A4</td>
<td>4</td>
</tr>
<tr>
<td><strong>Parlodel</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Pergolide</strong>&lt;sup&gt;a&lt;/sup&gt;</td>
<td>Tab 0.05, 0.25, 1 mg.</td>
<td>PO 0.05 mg/day for 2 days initially, increasing in 0.1–0.15 mg/day increments q 3 days for 12 days, then in 0.25 mg/day increments q 3 days to a usual dosage of 1–4 mg/day.</td>
<td>D&lt;sub&gt;1&lt;/sub&gt;: +/+  D&lt;sub&gt;2&lt;/sub&gt;: ++  D&lt;sub&gt;3&lt;/sub&gt;: ++</td>
<td>Inhibits CYP2D6</td>
<td>27</td>
</tr>
<tr>
<td><strong>Permax</strong></td>
<td></td>
<td></td>
<td></td>
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</tr>
</tbody>
</table>

(continued)
### DOPAMINE AGONISTS COMPARISON CHART

<table>
<thead>
<tr>
<th>DRUG</th>
<th>DOSAGE FORMS</th>
<th>ADULT DOSAGE</th>
<th>DOPAMINE RECEPTOR SELECTIVITY</th>
<th>EFFECT ON CYTOCHROME P450 ISOZYMES</th>
<th>HALF-LIFE (HR)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Pramipexole</td>
<td>Tab 0.125, 0.25,</td>
<td>PO 0.125 mg tid initially, increasing in increments of 0.75 mg/day at weekly</td>
<td>0</td>
<td>No effect.</td>
<td>12</td>
</tr>
<tr>
<td></td>
<td>1, 1.5 mg.</td>
<td>intervals to a usual dosage of 1.5–4.5 mg/day in 3 divided doses.</td>
<td>++</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Mirapex</td>
<td></td>
<td></td>
<td>++++</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Ropinirole</td>
<td>Tab 0.25, 0.5, 1,</td>
<td>PO 0.25 mg tid initially, increasing in increments of 0.75 mg/day at weekly</td>
<td>0</td>
<td>Inhibits CYP2D6</td>
<td>4–6</td>
</tr>
<tr>
<td></td>
<td>2, 5 mg.</td>
<td>intervals for 4 weeks, then in increments of 1.5–3 mg/day at weekly intervals</td>
<td>++</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>to a usual dosage of 3–12 mg/day in 3 divided doses.</td>
<td>+++</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Requip</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

0 = none; 0/+ = minimal; + = mild; ++ = moderate; +++ = potent; ++++ = very potent.

*Ergot alkaloid.

From references 302, 303, 305, 306, and 308 and product information.
Ophthalmic Drugs for Glaucoma

Class Instructions. Ophthalmic Solutions. Proper instillation of eye drops improves absorption of the drug into the eye and minimizes systemic absorption and adverse effects. If you wear contact lenses, remove them. Wash your hands before instilling eye drops. Tilt the head back and pull down the lower lid. Place 1 drop into the lower lid. Once medication has been placed in the eye(s), close the eyes and press lightly on the inside corner of each eye. Keep the eyes closed and continue pressure to the inside corner of the eyes for 2–5 minutes. Wash your hands to remove medication. If you miss a dose, apply it as soon as possible. If it is almost time for your next dose, skip the missed dose and go back to your regular schedule. Do not double doses. Ophthalmic Ointments. If you wear contact lenses, remove them. Wash your hands. Tilt the head back and pull down the lower lid. Unless told to use a different amount, squeeze a thin strip (about 0.5 cm) of ointment into the lower lid. Let go of the eyelid and close the eyes for 1–2 minutes. Wash your hands to remove any medication. To keep the medication as germ free as possible, do not touch the tip to any surface. Wipe the tip with a clean tissue before closing. If you miss a dose, take it as soon as possible. If it is almost time for your next dose, skip the missed dose and go back to your regular dosage schedule. Do not double doses.

Pharmacology. The only medical treatment for primary open-angle glaucoma is to decrease intraocular pressure (IOP), the only treatable risk factor. Glaucoma drugs lower IOP by reducing production of aqueous humor, decreasing the resistance to outflow of aqueous humor through the trabecular meshwork, and improving flow through uveoscleral pathway.

Administration and Dosage. The ocular cul-de-sac has a capacity of only about 7 μL. After instillation of an eye drop, this capacity temporarily increases to 30 μL. Although manufacturers’ package inserts often instruct an individual to instill a dose of 1 or 2 drops, the drop size of ophthalmic solutions, from about 26 μL for timolol to about 69 μL for carbachol, exceeds the capacity of the cul-de-sac. Control dosage by changing the concentration of the solution rather than instilling multiple drops. Ophthalmic solutions are generally administered at a frequency that is determined by their duration of action. Gels and ocular inserts provide a sustained-release of active drug from the vehicle, allowing some products to be administered less frequently than solutions of the same drug. Because they are effective and have relatively fewer adverse effects, begin treatment with a β-adrenergic blocker with a goal of decreasing IOP by 30%. To slow progression of visual field loss, patients with more severe glaucoma require greater reductions, possibly to as low as 7–12 mm Hg. If the goal IOP cannot be reached, substitute a carbonic anhydrase inhibitor (CAI), latanoprost, or α₂-adrenergic agonist. If monotherapy is not successful, use a rational combination of drugs.

Patient Instructions. One study found that, due to noncompliance, patients were without treatment for 30% of a 12-month follow-up period. Because noncompli-
ance is a major reason for treatment failure, persist in patient counseling. (See Class Instructions.)

**Pharmacokinetics.** In ocular therapeutics, the eye is considered a separate entity outside the body, with the aqueous humor considered the central compartment. Absorption is the process by which a drug enters the aqueous humor, and bioavailability refers to the rate and extent of absorption into the aqueous humor. Distribution refers to the flow dynamics of partitioning and binding of the drug from the aqueous to surrounding tissues, such as the ciliary body.

**Fate.** In general, ophthalmic solutions must have lipid and aqueous solubility to penetrate the cornea and reach their sites of action in the ciliary body. The epithelium and endothelium of the cornea are lipophilic. The inner layer, the stroma, is hydrophilic. The lipophilic epithelium is penetrated by the undissociated drug. Then the stroma is penetrated by the dissociated, hydrophilic drug. Corneal penetration is enhanced when the epithelium is injured or otherwise compromised. Drug that does not penetrate the cornea can be systemically absorbed through the conjunctival vessels or through nasolacrimal drainage. Most of an eye drop is drained within 15–30 sec of application and 80–85% of the drainage is through the nasolacrimal canal. Drugs that are systemically absorbed after ophthalmic administration do not pass through the liver; therefore, a relatively small amount of absorbed drug can result in adverse systemic effects. Nasolacrimal occlusion increases drug–corneal contact time, thereby enhancing ocular absorption and decreasing systemic absorption. Drugs that pass through the cornea and reach their sites of action can be metabolized by esterases but mostly are eliminated from the eye by aqueous humor turnover, which is 1.5% of the anterior chamber volume per minute. Normally, very little drug reaches the vitreous or crosses the blood–ocular barrier. Because sampling the aqueous humor or ocular tissues would cause severe pain or injury, pharmacokinetic studies are not usually conducted in the eye.

$t_\frac{1}{2}$. Half-life for ophthalmic solutions is determined primarily by tissue binding. For drugs that are not strongly bound to pigments in the iris or other tissues, half-life is determined by the aqueous humor turnover rate of 1.5%/min, which is consistent with a half-life of 46 min.

**Parameters to Monitor.** (See specific drug class.) An ophthalmologist or optometrist should monitor IOP q 2 weeks during initial treatment and stabilization. Once target IOP has been reached, IOP, cup/disc ratios, and visual fields tests should be monitored by an ophthalmologist or optometrist q 3–12 months, depending on the severity of glaucoma and the progression of visual loss. Pharmacokinetic monitoring is limited to noncompliance and detection of adverse effects. When a patient presents with new systemic problems, always consider the ophthalmic drug as a potential cause.

**α2-ADRENERGIC AGONISTS:**

<table>
<thead>
<tr>
<th>APRACLONIDINE HYDROCHLORIDE</th>
<th>1% ophthalmic solution</th>
</tr>
</thead>
<tbody>
<tr>
<td>BRIMONIDINE HYDROCHLORIDE</td>
<td>0.2% ophthalmic solution</td>
</tr>
</tbody>
</table>

**Pharmacology.** Apraclonidine and brimonidine act at α2-adrenergic sites in the ciliary body to inhibit norepinephrine release, causing a decrease in aqueous humor production. Brimonidine also increases uveoscleral outflow.
Clonidine is more polar than clonidine, resulting in less permeability of the blood–brain barrier. Apaclonidine has a high rate of tachyphylaxis, limiting it to short-term use. Brimonidine is more $\alpha_2$-selective and more lipophilic than apraclonidine, allowing the use of lower concentrations.

**Administration and Adult Dosage.** (Apraclonidine) **Ophth** in laser surgery 1 drop of 1% soln in the affected eye 1 hr before surgery and then 1 drop immediately after surgery to prevent the IOP spikes that occur. **Ophth in open-angle glaucoma as a short-term adjunct** 0.5% tid. (Brimonidine) **Ophth** for primary open-angle glaucoma 1 drop of 0.2% soln tid, about 8 hr apart. (See Notes.)

**Special Populations. Pediatric Dosage.** (Apraclonidine) same as adult dosage; (Brimonidine) (<12 yr) not recommended.

**Geriatric Dosage.** Same as adult dosage.

**Dosage Forms.** (Apraclonidine) **Ophth Soln** 0.5, 1%. (Brimonidine) **Ophth Soln** 0.15, 0.2%.

**Patient Instructions.** (See Ophthalmic Solutions Class Instructions.)

**Pharmacokinetics. Onset and Duration** (Apraclonidine) onset 1 hr, peak 3 hr. (Brimonidine) onset 1 hr, peak 2 hr, duration about 6 hr.

**Fate.** (See Ophthalmic Solutions Fate.) Brimonidine that is systemically absorbed is metabolized primarily in the liver; 74% is eliminated in the kidney within 120 hr. $t_{1/2}$ (Apraclonidine) plasma half-life 8 hr. (Brimonidine) plasma half-life 3 hr.

**Adverse Reactions.** (Apraclonidine) causes adverse ocular effects in 15–48% of patients, especially allergic reactions and rarely upper eyelid retraction. Frequent systemic effects are dry mouth and dry nose. Cardiovascular effects have not been reported. (Brimonidine) has similar but less frequent ocular adverse effects. Frequent ocular effects are blepharitis, blepharoconjunctivitis, conjunctival follicles, blurred vision, and headache. Like apraclonidine, it frequently causes dry mouth and dry nose. Brimonidine does not decrease heart rate. It can mildly decrease blood pressure in some patients, although it frequently causes lethargy. Because it crosses the blood–brain barrier, it can cause mild hypotension in adults occasionally. Several severe adverse systemic effects have been reported in children between 28 days and 3 months of age, including bradycardia, hypotension, hypothermia, hypotonia, apnea, dyspnea, hyperventilation, cyanosis, and lethargy. This is believed to be caused by immaturity of the blood–brain barrier and higher systemic concentrations because of low body weight.

**Precautions.** (See Ophthalmic Solutions Precautions.)

**Parameters to Monitor.** Monitor for conjunctivitis and lethargy. (See Ophthalmic Solutions Parameters to Monitor.)

**Notes.** Brimonidine 0.15% preserved with an oxychloro complex (Alphagan P) is equivalent to the 0.2% solution preserved with benzalkonium chloride (Alphagan). Brimonidine bid is equivalent to timolol 0.5% in lowering IOP at peak, 6.5 vs 6.1 mm Hg, but much less effective in lowering trough IOP, 4.3 vs 6.3 mm Hg. Brimonidine bid is more effective than betaxolol 0.25% suspen-
sion at peak and trough. As adjunct therapy in patients who fail to reach target IOP with other therapy, brimonidine bid decreases IOP an additional 4.7 ± 5.3 mm Hg, or 20%. (See Glaucoma Drugs Comparison Chart.)

### β-ADRENERGIC BLOCKING DRUGS:

<table>
<thead>
<tr>
<th>Drug Name</th>
<th>Trade Names</th>
</tr>
</thead>
<tbody>
<tr>
<td>Betaxolol Hydrochloride</td>
<td>Betoptic, Betoptic-S</td>
</tr>
<tr>
<td>Carteolol Hydrochloride</td>
<td>Ocupress</td>
</tr>
<tr>
<td>Levobunolol Hydrochloride</td>
<td>Betagan, Various</td>
</tr>
<tr>
<td>Metipranolol</td>
<td>Optipranolol</td>
</tr>
<tr>
<td>Timolol Maleate</td>
<td>Timoptic, Timoptic-XE</td>
</tr>
</tbody>
</table>

**Pharmacology.** β-Adrenergic blocking drugs downregulate adenylate cyclase by blocking β2-adrenergic receptors in the ciliary body, resulting in a decrease in aqueous production and intraocular pressure. Although betaxolol is a β1-selective adrenergic blocker, it is effective in treating glaucoma. Betaxolol might have more β2 activity than previously thought; small concentrations of β2-blockade might be sufficient to curb aqueous production; β2-receptors in the eye might be different from those in other tissues; or betaxolol’s IOP-lowering effect might be caused by a calcium antagonistic effect. Carteolol has intrinsic sympathomimetic activity (ISA) that theoretically makes it less likely to cause adverse pulmonary or cardiovascular effects and possibly provide increased blood flow to the retina. ISA does not seem to make a difference in cardiac effects in most studies; however, in one study night-time bradycardia was 4-fold greater in patients treated with timolol than in those treated with carteolol. Retinal and optic nerve head circulations are improved by β-adrenergic blocking agents without ISA.

**Administration and Adult Dosage.** (Betaxolol) Ophthalmic 1 drop bid. (Carteolol) Ophthalmic 1 drop bid. (Levobunolol) Ophthalmic initiate treatment with 1 drop/day. (Metipranolol) Ophthalmic 1 drop bid. (Timolol soln) Ophthalmic 1 drop bid of 0.25% soln initially; if target IOP is not reached in 4 weeks, increase to 0.5%. (Timolol gel-forming soln) 1 drop/day of 0.25% soln initially; if target IOP is not reached in 4 weeks, increase to 0.5%. (See Notes.)

**Special Populations.** Pediatric Dosage. Same as adult dosage. Geriatric Dosage. Same as adult dosage.

**Dosage Forms.** (Betaxolol) Ophthalmic 0.5%; Ophthalmic Susp 0.25%. (Carteolol) Ophthalmic Soln 1%. (Levobunolol) Ophthalmic Soln 0.25, 0.5%. (Metipranolol) Ophthalmic Soln 0.3%. (Timolol) Ophthalmic Gel-Forming Soln 0.25, 0.5%; Ophthalmic Soln 0.25, 0.5%.

**Patient Instructions.** (See Class Instructions.)

**Pharmacokinetics.** Onset and Duration. (Betaxolol) onset 30 min, peak 2 hr, duration 12 hr. (Carteolol) onset 1 hr, peak 2 hr, duration 12 hr. (Levobunolol) onset 1 hr, peak 2–6 hr, duration 24 hr. (Metipranolol) onset 30 min, peak 2 hr, duration 24 hr. (Timolol drops) onset 30 min, peak 1–2 hr, duration 24 hr.
**Fate.** (See Ophthalmic Solutions Fate.)

$t_{1/2}$. (Betaxolol) 12–20 hr; (carteolol) 3–7 hr; (levobunolol) 6 hr; (metipranolol) 3–4 hr; (timolol) 3–5 hr. (See Ophthalmic Solutions $t_{1/2}$.)

**Adverse Reactions.** Frequent, but mild, ocular adverse effects include burning and stinging at instillation. Betaxolol ophthalmic suspension and timolol gel-forming solution frequently cause temporary blurred vision. Occasional, but serious, granulomatous anterior uveitis is caused by metipranolol. Occasional, but serious, systemic reactions include bronchospasm, bradycardia, CHF, heart block, cerebral vascular ischemia, and depression.

**Contraindications.** Sinus bradycardia; greater than first-degree AV block; cardiogenic shock; overt cardiac failure. Nonselective drugs are also contraindicated in patients with histories of bronchial asthma or severe COPD.

**Precautions.** Diabetes mellitus; cerebrovascular insufficiency; myasthenia gravis.

**Drug Interactions.** Oral β-adrenergic blocking agents, calcium-channel blockers, and digoxin can cause additive effects on AV conduction. Quinidine can inhibit the metabolism of β-adrenergic blocking agents by CYP2D6, causing bradycardia.

**Parameters to Monitor.** (See Ophthalmic Solutions Parameters to Monitor.) Monitor for complaints of ocular adverse effects such as burning or stinging. Monitor pulse rate, shortness of breath, browache, nervousness, and depression.

**Notes.** If target IOP is not reached with a β-blocker within 4 weeks, consider switching to a topical ophthalmic CAI, α2-adrenergic agonist, or prostaglandin analogue rather than adding another drug. If monotherapy is not successful, a β-blocker can be combined with one of these drugs or pilocarpine. With the exception of betaxolol, β-blockers are not effective when combined with epinephrine or dipivefrin. (See Glaucoma Drugs Comparison Chart.)

**CARBONIC ANHYDRASE INHIBITORS:**

<table>
<thead>
<tr>
<th>ACETAZOLAMIDE</th>
<th>Diamox</th>
</tr>
</thead>
<tbody>
<tr>
<td>BRINZOLAMIDE</td>
<td>Azopt</td>
</tr>
<tr>
<td>DICHLORPHENAMIDE</td>
<td>Daranide</td>
</tr>
<tr>
<td>DORZOLAMIDE HYDROCHLORIDE</td>
<td>Trusopt</td>
</tr>
<tr>
<td>METHAZOLAMIDE</td>
<td>Naptazane</td>
</tr>
</tbody>
</table>

**Pharmacology.** CAIs inhibit the carbonic anhydrase II isoenzyme in the ciliary epithelium, thereby blocking the formation of bicarbonate. This causes a decrease in sodium and water outflow from the ciliary body. More than 99% of carbonic anhydrase must be inhibited to be effective. The result is a decrease of about 40% in aqueous humor production and a decrease in IOP of up to 30–35%. Orally administered CAIs also inhibit carbonic anhydrase in the kidney, red blood cells, and other tissues, causing diuresis and often acidosis and other serious adverse effects that limit their use.
Administration and Adult Dosage. Ophth for primary open-angle glaucoma (brinzolamide) 1 drop tid; (dorzolamide) 1 drop tid. When used adjunctively, dorzolamide is administered bid.366 PO for primary open-angle glaucoma (acetazolamide) SR cap 500 mg bid has been better tolerated than tablets; Tab 125 mg q 4 hr to 250 mg qid. Dosages >1 g/day are no more effective. (Dichlorphenamide) 100–200 mg priming dose, followed by 100 mg q 12 hr until desired response is obtained, then 25–50 mg daily to tid. (Methazolamide) 50–100 mg bid-tid. PO for prevention of altitude sickness (acetazolamide) 750 mg/day.367 (See Notes.)

Special Populations. Pediatric Dosage. Safety and efficacy not established. However, dorzolamide 2% ophthalmic solution is used in infantile glaucoma, and acetazolamide 5–10 mg/kg qid has been used when an oral CAI was necessary.341,364,368

Geriatric Dosage. Same as adult dosage.

Dosage Forms. (Acetazolamide) Tab 125, 250 mg; SR Cap 500 mg; Inj 500 mg. (Brinzolamide) Ophth Susp 1%. (Dichlorphenamide) Tab 50 mg. (Dorzolamide) Ophth Soln 2%. (Methazolamide) Tab 25, 50 mg.

Patient Instructions. (See Class Instructions.) Dorzolamide Tell your doctor if you experience itching, redness, swelling, or other sign of eye or eyelid irritation. This medication can cause you to have blurred vision for a short period. Make sure you know how to react to this medication before you drive, use a machine, or do anything else that might be dangerous if you cannot see properly. Dorzolamide can cause your eyes to become more sensitive to light. Wearing sunglasses and avoiding exposure to bright light can lessen the discomfort.338

Pharmacokinetics. Onset and Duration. (Acetazolamide) Tab peak IOP reduction 2–6 hr, duration 4–12 hr;341,364 SR cap onset 2–4 hr, peak 4–8 hr, duration 12–24 hr.341,364 (Brinzolamide) onset <2 hr, peak 2 hr, duration >12 hr.369 (Dichlorphenamide) onset 30 min, peak 2 hr, duration 6 hr. (Dorzolamide) onset <2 hr, peak 2–4 hr, duration 6–8 hr.341,370,371 (Methazolamide) onset 1–2 hr, peak 4–6 hr, duration 12–24 hr.341

Fate. For all oral CAIs there is a linear relationship between plasma concentration and dose. (Acetazolamide) virtually completely absorbed with a peak serum level of 30 mg/L occurring at 1 hr after a 500 mg dose of tablet; with SR Cap, serum levels remain >10 mg/L for 10 hr. 90% bound to plasma proteins; elimination is by active renal tubular secretion.364 (Methazolamide) well absorbed and distributed in plasma, CSF, aqueous humor, red blood cells, bile, and extracellular fluid. Peak serum concentrations after 50 and 100 mg bid dosages are 5.1 and 10.7 mg/L, respectively. Vdss is 17–23 L. Renal clearance accounts for 20–25% of the total clearance, with about 25% of the drug eliminated in the urine unchanged. Brinzolamide and dorzolamide are systemically absorbed and bind to carbonic anhydrase in erythrocytes with terminal half-lives of 111 and 147 days, respectively; however, there is only a 21% decrease in baseline carbonic anhydrase activity, far below the 99% inhibition level necessary to induce systemic effects.368 Laboratory values of patients receiving dorzolamide did not indicate metabolic acidosis or electrolyte imbalances such as those with long-term systemic CAIs.370
Adverse Reactions. Topical ophthalmic solutions frequently cause ocular burning, stinging, or allergic ocular reactions. However, fewer patients discontinue dorzolamide than pilocarpine. Frequent systemic effects of topical ophthalmic solutions consist of bitter taste, occasional headache, nausea, fatigue, and, rarely, urolithiasis and iridocyclitis. Oral administration frequently causes paresthesias, GI disturbances, anorexia, drowsiness, and confusion. Occasionally, metabolic acidosis, hypokalemia, or urolithiasis occurs. Attempt to treat acidosis with sodium acetate 90 mEq/day. Rare, but possibly fatal, reactions include aplastic anemia, agranulocytosis, and thrombocytopenia.

Contraindications. (Oral) hypokalemia; hyponatremia; hyperchloremic acidosis; adrenocortical insufficiency; marked renal or hepatic impairment; severe COPD. Long-term use of oral CAIs is contraindicated in angle-closure glaucoma.


Drug Interactions. Do not use topical CAIs with oral CAIs because the combination is no more effective and adverse effects are additive, particularly in causing corneal endothelial dysfunction. Oral CAIs can cause salicylate toxicity in patients taking high doses of aspirin, and salicylates can displace acetazolamide from plasma binding sites, causing acetazolamide toxicity and non–anion-gap hyperchloremic metabolic acidosis. Diflunisal displaces acetazolamide from plasma binding sites. In one study, this resulted in a 5.6-fold increase in acetazolamide plasma levels.

Parameters to Monitor. Malaise or fatigue, Cr, serum potassium, serum carbon dioxide. The value of monitoring CBC is controversial because the hematologic adverse effects can be immune mediated and idiosyncratic rather than dose related. However, manufacturers recommend obtaining a baseline CBC and platelet count, with monitoring at regular intervals.

Notes. Because of their severe adverse effects and poor tolerability, use oral CAIs in primary open-angle glaucoma only as a last resort. Some clinicians consider laser surgery before using oral CAIs long term. Use topical CAIs only if a topical β-blocker, prostaglandin analogue, or α2-adrenergic agonist cannot be used or has failed to reach target IOP. If target IOP is not achieved with monotherapy, a topical CAI can be added to another topical treatment. Dorzolamide 2% tid as monotherapy lowers IOP 23% compared with 25% for timolol and 21% for betaxolol. Added to timolol, dorzolamide 2% bid provides another 13–22% decrease in IOP, similar to that from adding acetazolamide. Topical dorzolamide is as effective as oral acetazolamide. (See Glaucoma Drug Comparison Chart.) For prophylaxis of acute altitude sickness, acetazolamide 500 mg/day is
ineffective, but 750 mg/day is about as effective as **dexamethasone** 8–16 mg/day.\textsuperscript{367}

### CHOLINERGICS AND CHOLINESTERASE INHIBITORS:

<table>
<thead>
<tr>
<th>CHOLINERGICS</th>
<th><strong>CARBACHOL</strong></th>
<th>Isopto Carbachol</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>DEMECARIUM</strong></td>
<td><strong>ECHOThIOpHATE IODIDE</strong></td>
<td><strong>PHOSPHOLine Iodide</strong></td>
</tr>
<tr>
<td><strong>PILOCARPINE SALTS</strong></td>
<td>Various</td>
<td></td>
</tr>
</tbody>
</table>

**Pharmacology.** Carbachol and pilocarpine are direct cholinergic agonists that act at acetylcholine receptors to stimulate the ciliary muscle. Carbachol is also a weak cholinesterase inhibitor. Cholinesterase inhibitors act indirectly by inhibiting AChE. Ciliary body contraction causes pupilary constriction and eases the restriction of outflow of aqueous humor through the trabecular meshwork. Demacarium and echothiophate are irreversible cholinesterase inhibitors with long durations of action.\textsuperscript{380}

**Administration and Adult Dosage.** **Ophthalmic** (carbachol) initiate at 1 drop tid of the 0.75% solution; (demecarium) 1 drop daily–bid of the 0.125–0.25% solution; (echothiophate) 1 drop daily–bid; (pilocarpine ophthalmic solution) initiate with 1 drop of 1–2% solution q 6–8 hr. Most patients eventually require qid administration. Because pilocarpine is bound to pigments in the iris and the ciliary body, patients with dark eyes sometimes require 4% and occasionally 6% solutions; (pilocarpine gel) apply a thin strip hs; (pilocarpine inserts) place in conjunctival sac once weekly at hs. When switching to pilocarpine inserts, initiate therapy with Ocusert Pilo-20 because there is no correlation between dosage of solution and that of inserts. **Ophthalmic** (demecarium) 1 drop daily for 2–3 weeks and then q 2–3 days for 3–4 weeks; (echothiophate) 1 drop daily–q 2 days.

**Special Populations.** **Pediatric Dosage.** Same as adult dosage.

**Geriatric Dosage.** Same as adult dosage.

**Dosage Forms.** (Carbachol) **Ophthalmic Solution** 0.75, 1.5, 2.25, 3%. (Demecarium) **Ophthalmic Solution** 0.125, 0.25%. (Echothiophate Iodide) **Powder for reconstitution** 0.03, 0.06, 0.125, 0.25%. (Pilocarpine) **Ophthalmic Gel** 4%; **Ocular therapeutic system** (Ocusert Pilo) 20, 40 \(\mu\)g/hr. (Pilocarpine Hydrochloride) **Ophthalmic Solution** 0.25, 0.5, 1, 2, 3, 4, 5, 6, 8, 10%. (Pilocarpine Nitrate) **Ophthalmic Solution** 1, 2, 4%.

**Patient Instructions.** (See Glaucoma Drugs Class Instructions.) Isoflurophate ophthalmic ointment is inactivated by moisture. Do not rinse the tip of the tube.

**Pharmacokinetics.** **Onset and Duration.** (Carbachol Ophthalmic Solution) onset 13 ± 2.2 min, peak 4 hr, duration 8 hr. (Demecarium Ophthalmic Solution) onset 2–4 hr, peak 24 hr, duration 5–9 days. (Echothiophate Solution) onset within minutes, peak 2–7 weeks, duration several weeks. (Isoflurophate Ophthalmic Gel) onset 15 min, peak within 24 hr, duration 1–4 weeks. (Physostigmine Ophthalmic Solution) onset 8 min, peak 1–2 hr, duration 4–6 hr. (Pilocarpine Ophthalmic Solution) onset within min, peak 2 hr, duration...
8 hr, (Pilocarpine Ophth Gel) 4% maintains IOP reductions of 30% or more for 24 hr.\(^\text{381}\) (Pilocarpine Ocuserts) release drug constantly for 1 week.\(^\text{382,383}\)

**Fate.** For absorption characteristics, see Glaucoma Drugs Fate. Cholinergic and cholinesterase inhibitors are hydrolyzed by acetylcholine.

\(t_{\text{1/2}}\). (See Ophthalmic Solutions \(t_{\text{1/2}}\).)

**Adverse Reactions.** Reduced visual acuity in poor lighting occurs frequently. Occasional effects include ciliary spasm, headache, lacrimation, myopia, blurred vision, retinal detachment, and iris cysts. Adverse effects (eg, iris cysts) occur more often in children, especially with use of long-acting cholinesterase inhibitors. Cataracts occur in 30–50% of elderly patients using echothiophate or demecarium for at least 6 months.\(^\text{341}\) Cholinergic syndrome consisting of weakness, nausea, diaphoresis, and dyspnea occurs rarely.\(^\text{384,385}\) Because of their long duration of action, adverse systemic effects are more likely with long-acting cholinesterase inhibitors.\(^\text{383}\) Patients with myopia of \(\geq\) 6 diopters or greater and those with histories of retinal detachment are at greater risk of developing retinal detachment.\(^\text{346}\)

**Contraindications.** Acute iritis and other conditions in which papillary constriction is undesirable.

**Precautions.** Pregnancy, lactation. Night driving or other activities in poor light. Use cholinesterase inhibitors cautiously in patients with histories of retinal detachment, asthma, bradycardia, hypotension, epilepsy, parkinsonism, recent MI, or patients using systemic cholinesterase inhibitors for myasthenia gravis.

**Drug Interactions.** Antihistamine, antidepressants, antipsychotics, and other anticholinergics can decrease the effects of cholinergics and cholinesterase inhibitors.

**Parameters to Monitor.** Intraocular pressure, cup/disc ratios, and visual field loss should be monitored by an ophthalmologist or optometrist. Miosis is an indication that cholinergic activity is present.\(^\text{386}\) Monitor compliance, pulse for bradycardia, and complaints of visual blurring, nausea, vomiting, diarrhea, and headache.

**Notes.** (See Glaucoma Drugs Comparison Chart.) Use long-acting cholinesterase inhibitors in patients who are not controlled with pilocarpine. Longer-acting agents are also used to diagnose and treat accommodative esotropia.

**PROSTAGLANDINS:**

| BIMATOPROST | Latanoprost is an ester prologue analogue of prostaglandin F\(_{2\alpha}\) that decreases IOP by increasing uveoscleral outflow by an unknown mechanism.\(^\text{387}\) Latanoprost usually lowers IOP by 5–8 mm Hg regardless of baseline pressure.\(^\text{388-391}\) This is important for patients with normal-tension glaucoma who do not respond as well to other drugs.\(^\text{392}\) Unoprostone isopropyl is a docosanoid compound related to a metabolite of prostaglandin F\(_{2\alpha}\).\(^\text{393}\) Unoprostone lowers | Lumigan | Xalatan | Travatan | Rescula |
IOP by about 5 mm Hg in patients with higher IOP\(^{394}\) and about 2 mm Hg in patients with low-tension glaucoma.\(^{393}\)

**Administration and Adult Dosage.** Ophth for glaucoma (Bimatoprost) 1 drop daily in the evening. (Latanoprost) 1 drop of 0.005% solution daily in the evening. Higher concentrations are not as effective as the 0.005% soln.\(^{395}\) Once daily administration is more effective than bid and evening administration is more effective than morning administration.\(^{392,395–397}\) (Travaprost) 1 drop daily in the evening. (Unoprostone) 1 drop bid.

**Special Populations.** **Pediatric Dosage.** Safety and efficacy not established.

**Geriatric Dosage.** Same as adult dosage.

**Dosage Forms.** Ophth Soln (Bimatoprost) 0.03%; (Latanoprost) 0.005%; (Travaprost) 0.004%; (Unoprostone) 0.15%.

**Patient Instructions.** (See Class Instructions.) If other eye drops are used with this drug, separate administrations by at least 5 min. Remove contact lenses before instilling drops. Lenses can be reinserted after 15 min.

**Pharmacokinetics.** **Onset and Duration.** (Bimatoprost) onset 4 hr, peak 8–12 hr, duration 24 hr; (Latanoprost) onset 3–4 hr, peak 8–12 hr, duration 24 hr; (Travaprost) onset 2 hr, peak 12 hr, duration 24 hr; (unoprostone) onset 30 min, peak 1–2 hr, duration 12 hr.\(^{392}\)

**Fate.** (Bimatoprost) peak plasma level of 200 pmol/L is unlikely to produce systemic effects. (Latanaprost) is more lipophilic than its active metabolite, allowing excellent penetration of the cornea. Inside the aqueous, it is hydrolyzed to the active drug and reaches a peak concentration of 55 \(\mu\)g/L at 2–3 hr.\(^{398}\) \(V_{\text{dss}}\) is 0.16 ± 0.02 L/kg. The active drug is not metabolized in the aqueous, but 77–88% is systemically absorbed within 3 min and 90% is bound to plasma proteins. A peak plasma level of 64 ng/L (about \(10^{-10}\) mol/L) is reached within 40 min, a level too low to produce systemic effects. The active drug is metabolized by the liver and 88% of the metabolites are eliminated by the kidneys. Systemic Cl is 0.42 L/hr/kg. Systemic levels cannot be detected after 12 hr.\(^{398}\) (Travaprost) peak plasma levels of 25 ng/L are reached in 30 min and rapidly eliminated. (Unoprostone) peak plasma concentration of 760 ng/L of the de-esterified metabolite of unoprostone is reached 15 min after ocular installation.\(^{393}\)

**Adverse Reactions.** Ocular reactions frequently include burning, stinging, conjunctival hyperemia, foreign-body sensation, blurred vision. The major limitation is increased pigmentation of the iris in patients with green-brown, yellow-brown, and blue/gray-brown eyes that occurs after 3–17 months of use with latanoprost.\(^{388,390,391,396,397,399,400}\) Unoprostone has been used primarily in Japanese patients who have dark irises; however, one case has been reported. Difference in the frequency of this reaction between the two prostoglandins might be caused by differences in the selectivity for prostaglandin receptors.\(^{401,402}\) Flu symptoms occur frequently (6%) in patients receiving unoprostone. Diplopia occasionally occurs; retinal artery embolus, retinal detachment, and vitreous hemorrhage with
latanoprost occur rarely. Occasional upper respiratory infection has been reported but cannot definitively be linked to latanoprost.346,398

Precautions. Infections occur from contamination of multiple-dose containers. Instruct patients to avoid touching the tip of the container to the eye. Patients with diabetic retinopathy or complicated ocular surgery have a greater risk of developing cystoid macular edema, anterior uveitis, or vitreous hemorrhage.346,388

Drug Interactions. Precipitate occurs when used with thimerosal-containing eye drops. Separate doses of different ophthalmic solutions by at least 5 min.

Parameters to Monitor. (See also Ophthalmic Solutions Parameters to Monitor.) Darkening of iris, eye pain.

Notes. In comparisons with timolol, patients receiving latanoprost have an equal or greater reduction in IOP.388,391,396,397,399,403 Patients switched from timolol to latanoprost had an additional 1–5.5 mm Hg reduction in IOP.388,404 Latanoprost is additive when added to another glaucoma treatment. Adding latanoprost to timolol results in an additional IOP reduction of 13–37%.346,404 Latanoprost lowers IOP an additional 15% in patients receiving acetazolamide.405 Pilocarpine given 1 hr before latanoprost does not provide further IOP reduction; however, pilocarpine given 10 min to 1 hr after latanoprost results in a further decrease in IOP of about 5 mm Hg.406 (See Glaucoma Drugs Comparison Chart.)

**SYMPATHOMIMETICS:**

<table>
<thead>
<tr>
<th>DIPIVEFRIN HYDROCHLORIDE</th>
<th>AKPro, Propine</th>
</tr>
</thead>
<tbody>
<tr>
<td>EPINEPHRINE AND SALTS</td>
<td>Various</td>
</tr>
</tbody>
</table>

Pharmacology. Epinephrine stimulates α- and β2-adrenergic receptors in the ciliary body, increasing outflow. Dipivefrin is an epinephrine prodrug that is enzymatically converted into epinephrine in the eye. IOP is reduced by 20–25%.346,407

Administration and Adult Dosage. Ophth for glaucoma (epinephrine) 1 drop (usually 2%) bid; (dipivefrin) 1 drop of 0.1% solution bid.

Special Populations. Pediatric Dosage. Same as adult dosage.

Geriatric Dosage. Same as adult dosage.

Dosage Forms. (Dipivefrin) Ophth Soln 0.1%; (epinephrine HCl) Ophth Soln 0.5, 1, 2%.

Patient Instructions. (See Class Instructions.)

Pharmacokinetics. Onset and Duration. Onset <45 min; peak 4–6 hr; duration 24 hr.407

Fate. Dipivefrin is absorbed 17 times more than epinephrine. Upon entry into the cornea, the two pivalic acid groups are removed by esterases, yielding epinephrine. Because of the better absorption, it can be administered as a 0.1% solution, decreasing the amount of epinephrine exposure to the conjunctiva and available for systemic absorption, thereby decreasing adverse effects.342 Epinephrine that is absorbed systemically is metabolized by MAO and COMT.407
Adverse Reactions. Intolerance to ocular adverse effects leads to discontinuation of epinephrine in 80% of patients. Burning, tearing, reactive conjunctival hyperemia, allergic blepharconjunctivitis, and mydriasis resulting in blurring of vision occur frequently. Mydriasis is minimized when epinephrine is combined with pilocarpine and is more pronounced when used with β-adrenergic blockers. Adrenochrome deposits in palpebral conjunctiva and the superficial cornea occur occasionally. Rare systemic adverse effects include tachycardia, hypertension, anxiety, and arrhythmia.

Precautions. Because epinephrine causes mydriasis, avoid use in patients with narrow-chamber angles because the lens prevents epinephrine from reaching the retina. About 30% of aphakic patients develop cystoid macular edema.

Parameters to Monitor. IOP, cup/disk ratios, and visual fields tests should be performed by an ophthalmologist or optometrist q 3–12 mo, depending on the severity and progression of glaucoma. Monitor for blurring of vision, mydriasis, conjunctival irritation, hypertension and rapid pulse.

Notes. Do not use epinephrine solutions that are cloudy or have become pinkish or brownish. Epinephrine provides no extra benefit when combined with β-adrenergic blockers except betaxolol. (See Glaucoma Drugs Comparison Chart.)
<table>
<thead>
<tr>
<th>Drug Class and Drugs</th>
<th>Dosage Forms</th>
<th>Adult Dosage</th>
<th>Onset</th>
<th>Peak</th>
<th>Duration</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>α-Adrenergic Agonists</strong></td>
<td></td>
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<tr>
<td>Apraclonidine HCl</td>
<td>Ophth Soln 0.5, 1%</td>
<td>0.5% tid short-term; 1% 1 hr before and immediately after surgery</td>
<td>1 hr</td>
<td>3 hr</td>
<td>——</td>
</tr>
<tr>
<td>Iopidine</td>
<td>1 hr before and immediately after surgery</td>
<td></td>
<td></td>
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<tr>
<td>Brimonidine Tartrate</td>
<td>Ophth Soln 0.15%</td>
<td>tid</td>
<td>1 hr</td>
<td>2 hr</td>
<td>6 hr</td>
</tr>
<tr>
<td>Alphagan</td>
<td>(Alphagan P); 0.2% Alphagan P</td>
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</tr>
<tr>
<td>Betaxolol HCl</td>
<td>Ophth Soln 0.5%</td>
<td>bid;</td>
<td>30 min</td>
<td>2 hr</td>
<td>12 hr</td>
</tr>
<tr>
<td>Betoptic</td>
<td>Ophth Susp 0.25%</td>
<td>bid</td>
<td>30 min</td>
<td>2 hr</td>
<td>24 hr</td>
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<tr>
<td>Betoptic-S</td>
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<tr>
<td>Carteolol HCl</td>
<td>Ophth Soln 1%</td>
<td>bid</td>
<td>1 hr</td>
<td>2 hr</td>
<td>12 hr</td>
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<tr>
<td>Ocupress</td>
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<tr>
<td>Levobunolol HCl</td>
<td>Ophth Soln 0.25, 0.5%</td>
<td>daily-bid</td>
<td>1 hr</td>
<td>2–6 hr</td>
<td>24 hr</td>
</tr>
<tr>
<td>Betagan</td>
<td>Various</td>
<td></td>
<td></td>
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</tr>
<tr>
<td>Metipranolol</td>
<td>Ophth Soln 0.3%</td>
<td>bid</td>
<td>30 min</td>
<td>2 hr</td>
<td>24 hr</td>
</tr>
<tr>
<td>Optipranolol</td>
<td></td>
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</tr>
<tr>
<td>Timolol Maleate</td>
<td>Ophth Soln 0.25, 0.5%</td>
<td>bid</td>
<td>30 min</td>
<td>1–2 hr</td>
<td>24 hr</td>
</tr>
<tr>
<td>Timoptic</td>
<td>Timoptic-XE</td>
<td>Ophth Gel-Forming Soln 0.25, 0.5%</td>
<td>daily</td>
<td>30 min</td>
<td>1–2 hr</td>
</tr>
</tbody>
</table>

(continued)
## GLAUCOMA DRUGS COMPARISON CHART

### (continued)

<table>
<thead>
<tr>
<th>DRUG CLASS AND DRUGS</th>
<th>DOSAGE FORMS</th>
<th>ADULT DOSAGE</th>
<th>ONSET</th>
<th>PEAK</th>
<th>DURATION</th>
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<tr>
<td><strong>CAI TOPICAL</strong></td>
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</tr>
<tr>
<td><em>Brinzolamide</em></td>
<td>Ophth Susp 1%.</td>
<td>tid</td>
<td>&lt;2 hr</td>
<td>2 hr</td>
<td>&gt;12 hr</td>
</tr>
<tr>
<td>Azopt</td>
<td></td>
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<tr>
<td><em>Dorzolamide</em></td>
<td>Ophth Soln 2%.</td>
<td>tid</td>
<td>&lt;2 hr</td>
<td>2–4 hr</td>
<td>6–8 hr</td>
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<tr>
<td>Trusopt</td>
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<tr>
<td><strong>CAI ORAL</strong></td>
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<tr>
<td><em>Acetazolamide</em></td>
<td>Tab 125, 250 mg; to 250 mg qid</td>
<td>125–250 mg q 4 hr to 250 mg qid</td>
<td>1–2 hr</td>
<td>2–4 hr</td>
<td>4–12 hr</td>
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<tr>
<td>Diamox</td>
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<tr>
<td>Diamox Sequels</td>
<td>SR Cap 500 mg.</td>
<td>500 mg bid</td>
<td>2–4 hr</td>
<td>8 hr</td>
<td>12–24 hr</td>
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<tr>
<td><em>Dichlorphenamide</em></td>
<td>Tab 50 mg.</td>
<td>25–50 mg daily-tid</td>
<td>30 min</td>
<td>2–4 hr</td>
<td>6–12 hr</td>
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<tr>
<td>Daranide</td>
<td></td>
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<tr>
<td><em>Methazolamide</em></td>
<td>Tab 25, 50 mg.</td>
<td>50–100 mg bid-tid</td>
<td>1–2 hr</td>
<td>4–6 hr</td>
<td>12–24 hr</td>
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<td>Naptazane</td>
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<td><strong>CHOLINERGICS</strong></td>
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<tr>
<td><em>Carbachol</em></td>
<td>Ophth Soln 0.75, 1.5, 2.25, 3%.</td>
<td>tid</td>
<td>13 min</td>
<td>4 hr</td>
<td>8 hr</td>
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<tr>
<td>Isopto Carbachol</td>
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<tr>
<td><em>Pilocarpine HCl</em></td>
<td>Ophth Soln (HCl) 0.25, 0.5, 1, 2, 3, 4, 5, 6, 8, 10%</td>
<td>qid</td>
<td>minutes</td>
<td>2 hr</td>
<td>8 hr</td>
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<tr>
<td>Pilocar</td>
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<tr>
<td>Various</td>
<td>Ophth Gel 4%, daily</td>
<td>minutes</td>
<td>2 hr</td>
<td>24 hr</td>
<td></td>
</tr>
<tr>
<td><em>Pilocarpine Nitrate</em></td>
<td>Ophth Soln (Nitrate) 1, 2, 4%.</td>
<td>qid</td>
<td>minutes</td>
<td>2 hr</td>
<td>8 hr</td>
</tr>
<tr>
<td>Pilagan</td>
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<tr>
<td>DRUG CLASS AND DRUGS</td>
<td>DOSAGE FORMS</td>
<td>ADULT DOSAGE</td>
<td>ONSET</td>
<td>PEAK</td>
<td>DURATION</td>
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<tr>
<td>Pilocarpine Ocular</td>
<td>20, 40 μg/hr.</td>
<td>weekly</td>
<td>minutes</td>
<td>2 hr</td>
<td>7 days</td>
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<td>Therapeutic System</td>
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<td>Ocusert Pilo-20, 40</td>
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<td>CHOLINESTERASE INHIBITORS</td>
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<tr>
<td>Demecarium</td>
<td>Ophth Soln 0.125, 0.25%.</td>
<td>daily-bid</td>
<td>2–4 hr</td>
<td>24 hr</td>
<td>5–9 d</td>
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<tr>
<td>Humorsol</td>
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<tr>
<td>Echothiophate Iodide</td>
<td>Pwdr 0.03, 0.06, 0.125, 0.25%.</td>
<td>daily-bid</td>
<td>minutes</td>
<td>2–7 weeks</td>
<td>several weeks</td>
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<tr>
<td>Phospholine Iodide</td>
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<td>PROSTAGLANDIN ANALOGUES</td>
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<tr>
<td>Bimatoprost</td>
<td>Ophth Soln 0.03%.</td>
<td>p.m.</td>
<td>4 hr</td>
<td>8–12 hr</td>
<td>24 hr</td>
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<td>Lumigan</td>
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<tr>
<td>Latanoprost</td>
<td>Ophth Soln 0.005%.</td>
<td>p.m.</td>
<td>3–4 hr</td>
<td>8–12 hr</td>
<td>24 hr</td>
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<td>Xalatan</td>
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<tr>
<td>Travaprost</td>
<td>Ophth Soln 0.004%.</td>
<td>p.m.</td>
<td>2 hr</td>
<td>12 hr</td>
<td>24 hr</td>
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<tr>
<td>Travatan</td>
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<tr>
<td>Unoprostone</td>
<td>Ophth Soln 0.15%.</td>
<td>bid</td>
<td>30 min</td>
<td>1–2 hr</td>
<td>12 hr</td>
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<td>Rescula</td>
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<td>SYMPATHOMIMETICS</td>
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<tr>
<td>Epinephrine</td>
<td>Ophth Soln 0.5, 1, 2%.</td>
<td>bid</td>
<td>&lt;45 min</td>
<td>4–6 hr</td>
<td>24 hr</td>
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<td>Various</td>
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<td>Dipivefrin</td>
<td>Ophth Soln 0.1%</td>
<td>bid</td>
<td>&lt;45 min</td>
<td>4–6 hr</td>
<td>24 hr</td>
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<td>Propine</td>
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</table>

CAI = carbonic anhydrase inhibitor.
*Dosages in this chart are for primary open-angle glaucoma.
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108. Cooper BR et al. Evidence that the acute behavioral and electrophysiological effects of bupropion (Wellbutrin) are mediated by a noradrenergic mechanism. *Neuropsychopharmacology* 1994;11:333–41.


CENTRAL NERVOUS SYSTEM DRUGS


CENTRAL NERVOUS SYSTEM DRUGS


Acid-Peptic Therapy

**ANTACIDS**

**Pharmacology.** Antacids are weakly basic inorganic salts whose primary action is to neutralize gastric acid; pH >4 inhibits the proteolytic activity of pepsin. Aluminum-containing antacids suppress, but do not eradicate, *Helicobacter pylori* and can promote ulcer healing in peptic ulcer disease (PUD) by enhancing mucosal defense mechanisms.\(^1\,^2\) Aluminum salts also bind phosphate and bile salts in the GI tract, decreasing serum phosphate and serum bile salt levels. Antacids can increase urine pH.

**Administration and Adult Dosage.** PO for symptomatic relief of indigestion, nonulcer dyspepsia, epigastric pain in PUD, or heartburn in gastroesophageal reflux disease (GERD) 10–30 mL prn or 1 and 3 hr after meals and hs.\(^1\,^2\) PO for treatment of PUD 100–160 mEq of acid-neutralizing capacity per dose, given 1 and 3 hr after meals and hs for 4–8 weeks or until healing is complete. Additional doses may be taken if epigastric pain persists. There is evidence that lower dosages can heal peptic ulcers.\(^1\,^2\) PO or NG for prevention or treatment of upper GI bleeding in critically ill patients titrate to maintain intragastric pH >4.0.\(^3\) PO for phosphate binding in renal failure (aluminum hydroxide) 1.9–4.8 g tid or qid or (calcium carbonate) 8–12 g/day; titrate dosage based on serum phosphate.\(^1\)

**Special Populations.** *Pediatric Dosage.* PO for treatment of PUD or GERD (≤12 yr) at least 5–15 mL up to q 1 hr; (>12 yr) same as adult dosage.

*Geriatric Dosage.* Avoid using magnesium-containing antacids in renal impairment.

*Other Conditions.* Avoid using magnesium-containing antacids in patients with Clcr <30 mL/min.

**Dosage Forms.** (See Antacid Products Comparison Chart.)

**Patient Instructions.** If antacids do not relieve symptoms of indigestion, upset stomach, or heartburn within 2 weeks, contact your health care practitioner. Diarrhea can occur with magnesium-containing antacids; decrease the daily dosage, alternate doses with, or switch to, an aluminum- or calcium-containing antacid. Constipation can occur with aluminum-containing antacids; decrease the daily dosage, alternate doses with, or switch to, a magnesium-containing antacid. Refrigerating liquid antacids or flavored antacids can improve their palatability. Antacids can interfere with other medications; take other medications 1 to 2 hours...
before or after antacids unless otherwise directed. If tablets are used, chew thoroughly before swallowing and follow with a glass of water.

**Missed Doses.** If your health care practitioner has told you to take this medicine on a regular schedule and you miss a dose, take it as soon as possible. If it is almost time for your next dose, skip the missed dose and return to your usual dosage schedule. Do not double doses.

**Pharmacokinetics. Onset and Duration.** Onset of acid neutralizing is immediate; duration is 30 ± 10 min in the fasted state and 1–3 hr if ingested with or within 1 hr after meals.¹

**Fate.** Antacid cations are absorbed to different degrees. Sodium is highly soluble and readily absorbed; calcium absorption is generally less than 30% but can decrease with advancing age, intake, achlorhydria, and estrogen loss at menopause; magnesium is generally about 30% absorbed, but percentage of absorption changes inversely with intake; aluminum is slightly absorbed. Calcium, magnesium, and aluminum are excreted renally with normal renal function.¹ The unab sorbed portion is excreted in the feces.

**Adverse Reactions.** Long-term use of sodium- or calcium-containing antacids can cause systemic alkalosis. Hypercalcemia can occur with ingestion of large amounts of calcium; soluble antacids plus a diet high in milk products can result in milk-alkali syndrome, which can lead to nephrolithiasis and, in severe cases, neurologic abnormalities.¹ Magnesium-containing antacids cause dose-related laxative effects; hypermagnesemia occurs in patients with renal impairment.¹ Aluminum-containing antacids cause dose-related constipation, especially in the elderly. Prolonged administration or large dosages of aluminum hydroxide or carbonate can result in hypophosphatemia, particularly in the elderly and alcoholics; encephalopathy has been reported in dialysis patients receiving aluminum-containing antacids alone or with sucralfate.¹ ² ⁴

**Precautions.** Use caution with aluminum and calcium salts and avoid magnesium-containing products in patients with renal insufficiency. Use caution when using sodium bicarbonate in patients with chronic renal failure, edema, hypertension, or CHF. Because antacids are particulate and elevate intragastric pH, they can predispose critically ill patients to nosocomial pneumonia.³

**Drug Interactions.** Antacids reduce the absorption of numerous drugs by three different mechanisms: altering GI pH, altering urinary pH, and binding to drugs in the GI tract. Factors that affect the likelihood of drug interactions are the drug’s dose, valence of cations (eg, tetracycline is polyvalent), and timing of the doses of antacid and drug. Some clinically important interactions include digoxin, oral iron, isoniazid, ketoconazole, oral quinolones, and oral tetracyclines. Antacids can reduce salicylate levels and increase quinidine levels because of urinary pH changes. Large dosages of calcium antacids can produce hypercalcemia in the presence of thiazides. Sodium polystyrene sulfonate resin can bind magnesium and calcium ions from the antacid in the gut, resulting in systemic alkalosis.

**Parameters to Monitor.** Monitor for relief of dyspepsia, epigastric pain or heartburn, and diarrhea or constipation. Monitor serum phosphate during long-term use

Notes. Aggressive antacid therapy is at least as effective as the H$_2$-receptor antagonists or sucralfate when treating PUD or preventing stress-related mucosal bleeding; however, do not use antacids as first-line agents because large, frequent doses are inconvenient and associated with an increased risk of adverse effects.$^{1,2,4}$ Mag-aldrate is a chemical mixture of magnesium and aluminum hydroxides. Alginic acid has foaming and floating properties that can be beneficial in GERD. Most antacid products have been reformulated to contain low amounts of sodium; some antacid products contain considerable amounts of sugar or artificial sweetener. Antacid tablets, if chewed and swallowed, can be as effective as equivalent doses of liquid formulations. Although gastrin is stimulated by calcium, gastric acid rebound with calcium-containing antacids is of questionable clinical importance.$^1$
## ANTACID PRODUCTS COMPARISON CHART

<table>
<thead>
<tr>
<th>ANTACID</th>
<th>ORAL SUSPENSION</th>
<th>TABLETS</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Acid Neutralizing Capacity</td>
<td>Sodium Content</td>
</tr>
<tr>
<td></td>
<td>mEq/5 mL</td>
<td>mg/5 mL</td>
</tr>
<tr>
<td>Aluminum Carbonate, Basic</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Basaljel</td>
<td>11.5</td>
<td>3</td>
</tr>
<tr>
<td>Aluminum Hydroxide</td>
<td></td>
<td></td>
</tr>
<tr>
<td>AlternaGEL</td>
<td>16</td>
<td>&lt;2.5</td>
</tr>
<tr>
<td>Amphojel 300 mg</td>
<td>10</td>
<td>&lt;2.3</td>
</tr>
<tr>
<td>Amphojel 600 mg</td>
<td>—</td>
<td>—</td>
</tr>
<tr>
<td>Aluminum Hydroxide with Magnesium Carbonate</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Gaviscon</td>
<td>44</td>
<td>13</td>
</tr>
<tr>
<td>Gaviscon Extra Strength Relief Formula</td>
<td>14.3d</td>
<td>20.7</td>
</tr>
<tr>
<td>Aluminum Hydroxide with Magnesium Hydroxide</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Maalox</td>
<td>13.3</td>
<td>&lt;1.5</td>
</tr>
<tr>
<td>Maalox High Potency</td>
<td>27.2</td>
<td>&lt;1</td>
</tr>
<tr>
<td>Aluminum Hydroxide with Magnesium Hydroxide and Simethicone</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Fast-Acting Mylanta</td>
<td>12.7</td>
<td>0.6</td>
</tr>
<tr>
<td>Fast-Acting Mylanta, Maximum Strength</td>
<td>25.4</td>
<td>0.05</td>
</tr>
<tr>
<td>Gelusil</td>
<td>—</td>
<td>—</td>
</tr>
<tr>
<td>Maalox Plus</td>
<td>—</td>
<td>—</td>
</tr>
<tr>
<td>Maalox, Maximum Strength, Anti-Gas</td>
<td>29.8</td>
<td>&lt;2.5</td>
</tr>
<tr>
<td>Mylanta</td>
<td>12.7</td>
<td>0.68</td>
</tr>
<tr>
<td>Mylanta, Maximum Strength</td>
<td>25.4</td>
<td>1.14</td>
</tr>
</tbody>
</table>

(continued)
<table>
<thead>
<tr>
<th>ANTACID PRODUCT</th>
<th>ORAL SUSPENSION</th>
<th>TABLETS</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Acid Neutralizing Capacity</td>
<td>Sodium Content</td>
</tr>
<tr>
<td></td>
<td>mEq/5 mL</td>
<td>mg/5 mL</td>
</tr>
<tr>
<td><strong>Aluminum Hydroxide with Magnesium Trisilicate and Sodium Bicarbonate</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Gaviscon (Chewable)</td>
<td>—</td>
<td>—</td>
</tr>
<tr>
<td>Gaviscon-2 (Chewable)</td>
<td>—</td>
<td>—</td>
</tr>
<tr>
<td><strong>Calcium Carbonate</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Children's Mylanta Upset Stomach Relief</td>
<td>8</td>
<td>—</td>
</tr>
<tr>
<td>Maalox, Quick Dissolve</td>
<td>—</td>
<td>—</td>
</tr>
<tr>
<td>Maalox, Quick Dissolve Maximum Strength</td>
<td>—</td>
<td>—</td>
</tr>
<tr>
<td>Titralac</td>
<td>—</td>
<td>—</td>
</tr>
<tr>
<td>Tums</td>
<td>—</td>
<td>—</td>
</tr>
<tr>
<td>Tums E-X</td>
<td>—</td>
<td>—</td>
</tr>
<tr>
<td>Tums Ultra</td>
<td>—</td>
<td>—</td>
</tr>
<tr>
<td><strong>Calcium Carbonate with Magnesium Hydroxide</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Di-GeF</td>
<td>≥9</td>
<td>≤5</td>
</tr>
<tr>
<td>Fast-Acting Mylanta</td>
<td>24</td>
<td>0.3</td>
</tr>
<tr>
<td>Fast-Acting Mylanta, Maximum Strength</td>
<td>48</td>
<td>0.6</td>
</tr>
<tr>
<td>Fast-Acting Mylanta Supreme</td>
<td>12.6</td>
<td>0.7</td>
</tr>
<tr>
<td>Rolaids</td>
<td>—</td>
<td>—</td>
</tr>
</tbody>
</table>

(continued)
### ANTACID PRODUCTS COMPARISON CHART

<table>
<thead>
<tr>
<th>ANTACID</th>
<th>ORAL SUSPENSION</th>
<th>TABLETS</th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Acid Neutralizing Capacity mEq/5 mL</td>
<td>Sodium Content mg/5 mL</td>
<td>Acid-Neutralizing Capacity mEq/Tablet</td>
</tr>
<tr>
<td>Magaldrate</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Riopan</td>
<td>15</td>
<td>&lt;0.3</td>
<td>—</td>
</tr>
<tr>
<td>Riopan Plus&lt;sup&gt;c&lt;/sup&gt;</td>
<td>15</td>
<td>&lt;0.3</td>
<td>13.5</td>
</tr>
<tr>
<td>Riopan Plus Double Strength&lt;sup&gt;c&lt;/sup&gt;</td>
<td>—</td>
<td>—</td>
<td>30</td>
</tr>
</tbody>
</table>

<sup>a</sup>Products listed are representative of numerous brand and generic products on the market. Product formulations and, hence, neutralizing capacity and sodium content, are subject to change by the manufacturer.

<sup>b</sup>To determine the sodium content in mEq, multiply sodium content (mg) by 0.043.

<sup>c</sup>Contains simethicone.

<sup>d</sup>Contains alginate.

<sup>e</sup>Contains sodium bicarbonate.
Pharmacology. Bismuth salts are used to treat nausea, indigestion, diarrhea, gastritis, and peptic ulcers. The precise method by which bismuth heals gastritis and ulcers is uncertain, but possible mechanisms are local gastroprotective effect, stimulation of endogenous prostaglandins, and antimicrobial activity against Helicobacter pylori. Given alone, bismuth salts suppress H. pylori, but long-term eradication requires combination therapy with antibiotics. Bismuth subsalicylate (BSS; Pepto-Bismol, various) is the bismuth salt used most frequently in the United States. Ranitidine bismuth citrate (RBC) is a complex of ranitidine, trivalent bismuth, and citrate. Administration and Adult Dosage. PO for the control of nausea, abdominal cramps, and diarrhea (BSS) 525 mg. Administer dosage q 30–60 min, if needed, to a maximum of 4.2 g/day. When given with antibiotics to eradicate H. pylori, treatment is usually limited to 1–2 weeks. (See also Eradication of Helicobacter pylori Infection.) Special Populations. Pediatric Dosage. PO for the control of nausea, abdominal cramps, and diarrhea (BSS) (<3 yr) not recommended; (3–6 yr) 87 mg; (6–9 yr) 175 mg; (9–12 yr) 262 mg. Administer dosage q 30–60 min, if needed, to a maximum of 8 doses/day. Geriatric Dosage. Same as adult dosage. Dosage Forms. Susp (BSS) 17.5, 35 mg/mL; Chew Tab 262; Tab 262 mg; Tab (RBC) 400 mg (containing ranitidine 160 mg and bismuth citrate 240 mg) (Tritec). Pharmacokinetics. After oral administration, BSS (58% bismuth, 42% salicylate) is converted in the GI tract to bismuth oxide and salicylic acid. Bismuth is less than 0.2% absorbed, with more than 99% of an oral dose excreted in the feces. Over 90% of the salicylate dose is absorbed and excreted in urine. RBC dissociates in intragastric fluid to ranitidine and soluble and insoluble forms of bismuth. Oral absorption of bismuth from RBC is variable. Adverse Reactions. BSS and bismuth derived from RBC can temporarily darken the tongue and stool. Use BSS with caution in children; in the elderly; in patients with renal impairment, salicylate sensitivity, or bleeding disorders; in those receiving high-dosage salicylate therapy; or when potentially interacting medications are taken. Salicylic acid is less likely than aspirin to cause gastric mucosal damage and blood loss. Prolonged use, dosages higher than those recommended, and the use of other salts (subgallate and subnitrates) have been associated with neurotoxicity. Bismuth concentrations can be elevated in the elderly and patients with renal impairment because of decreased renal elimination. RBC should not be used as a single agent for the treatment of active duodenal or gastric ulcers. Use caution in children and teenagers who are experiencing or recovering from nausea and vomiting symptoms because these might be early signs of Reye’s syndrome. Avoid BSS in patients who are hypersensitive to aspirin or nonaspirin salicylates.
Pharmacology. Histamine H₂-receptor antagonists competitively inhibit the actions of histamine at the H₂ receptors of the parietal cell and reduce basal, nocturnal, pentagastrin-, and food-stimulated gastric acid.

**HISTAMINE H₂-RECEPTOR ANTAGONISTS:**

<table>
<thead>
<tr>
<th>NAME</th>
<th>TRADE NAME</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cimetidine</td>
<td>Tagamet, Various</td>
</tr>
<tr>
<td>Famotidine</td>
<td>Pepcid</td>
</tr>
<tr>
<td>Nizatidine</td>
<td>Axid</td>
</tr>
<tr>
<td>Ranitidine</td>
<td>Zantac, Various</td>
</tr>
</tbody>
</table>

**ACID-PEPTIC THERAPY**

<table>
<thead>
<tr>
<th>INDICATION</th>
<th>CIMETIDINE</th>
<th>FAMOTIDINE</th>
<th>NIZATIDINE</th>
<th>RANITIDINE</th>
</tr>
</thead>
<tbody>
<tr>
<td>PO for prevention or symptomatic relief of heartburn or indigestion (OTC)</td>
<td>200 mg/day or 200 mg bid.</td>
<td>10 mg/day or 10 mg bid.</td>
<td>75 mg/day or 75 mg bid.</td>
<td>75 mg/day or 75 mg bid.</td>
</tr>
<tr>
<td>PO for short-term treatment of active duodenal ulcer (4–8 weeks)</td>
<td>300 mg qid, 400 mg bid, 800 mg hs, or 1600 mg hs.</td>
<td>20 mg bid or 40 mg hs.</td>
<td>150 mg bid or 300 mg hs.</td>
<td>150 mg bid or 300 mg hs.</td>
</tr>
<tr>
<td>PO for maintenance of healed duodenal ulcer</td>
<td>400 mg hs.</td>
<td>20 mg hs.</td>
<td>150 mg hs.</td>
<td>150 mg hs.</td>
</tr>
<tr>
<td>PO for short-term treatment of active benign gastric ulcer (6–8 weeks)</td>
<td>300 mg qid, 400 mg bid or 800 mg hs</td>
<td>20 mg bid or 40 mg hs.</td>
<td>150 mg bid or 300 mg hs.</td>
<td>150 mg bid or 300 mg hs.</td>
</tr>
</tbody>
</table>

(continued)
<table>
<thead>
<tr>
<th>INDICATION</th>
<th>CIMETIDINE</th>
<th>FAMOTIDINE</th>
<th>NIZATIDINE</th>
<th>RANITIDINE</th>
</tr>
</thead>
<tbody>
<tr>
<td>PO for main-</td>
<td>400 mg hs&lt;sup&gt;b&lt;/sup&gt;</td>
<td>20 mg hs&lt;sup&gt;b&lt;/sup&gt;</td>
<td>150 mg hs&lt;sup&gt;b&lt;/sup&gt;</td>
<td>150 mg hs. or 300 mg hs.</td>
</tr>
<tr>
<td>tenance of healed gastric ulcer</td>
<td>or 800 mg hs.</td>
<td>or 300 mg hs.</td>
<td>or 300 mg hs.</td>
<td></td>
</tr>
<tr>
<td>PO for symptomatic gastroesophageal reflux disease (6–12 weeks)</td>
<td>300 mg qid or 400 mg bid.</td>
<td>20 mg bid.</td>
<td>150 mg bid.</td>
<td>150 mg bid.</td>
</tr>
<tr>
<td>PO for healing of erosive esophagitis (6–12 weeks)</td>
<td>400 mg qid or 800 mg bid.</td>
<td>20 or 40 mg bid.</td>
<td>150 mg bid or 300 mg bid&lt;sup&gt;b&lt;/sup&gt;</td>
<td>150 mg qid or 300 mg bid&lt;sup&gt;b&lt;/sup&gt;</td>
</tr>
<tr>
<td>PO for maintenance of healed erosive esophagitis</td>
<td>300 mg qid&lt;sup&gt;b&lt;/sup&gt;, 400 mg qid&lt;sup&gt;b&lt;/sup&gt;, or 800 mg bid&lt;sup&gt;b&lt;/sup&gt;</td>
<td>20 mg bid&lt;sup&gt;b&lt;/sup&gt; or 40 mg bid&lt;sup&gt;b&lt;/sup&gt;</td>
<td>150 mg bid&lt;sup&gt;b&lt;/sup&gt; or 300 mg bid&lt;sup&gt;b&lt;/sup&gt;</td>
<td>150 mg bid or 300 mg bid&lt;sup&gt;b&lt;/sup&gt;</td>
</tr>
<tr>
<td>PO for pathological hypersecretory conditions</td>
<td>300 mg q 6–8 hr&lt;sup&gt;c&lt;/sup&gt;</td>
<td>20 mg q 6 hr, up to 160 mg q 6 hr, or adjust to patient needs.</td>
<td>150 mg bid&lt;sup&gt;b&lt;/sup&gt;</td>
<td>150 mg bid, up to 6 g/day; or adjust to patient needs.</td>
</tr>
<tr>
<td>IM</td>
<td>300 mg q 6–8 hr&lt;sup&gt;c&lt;/sup&gt;</td>
<td>20 mg q 12 hr&lt;sup&gt;c&lt;/sup&gt;</td>
<td>50 mg q 6–8 hr&lt;sup&gt;c&lt;/sup&gt;</td>
<td>50 mg q 6–8 hr&lt;sup&gt;c&lt;/sup&gt;</td>
</tr>
<tr>
<td>IV intermittent</td>
<td>300 q 6–8 hr, up to 2.4 g/day&lt;sup&gt;c&lt;/sup&gt;</td>
<td>20 mg q 12 hr&lt;sup&gt;c&lt;/sup&gt;</td>
<td>50 mg q 6–8 hr, up to 400 mg/day&lt;sup&gt;c&lt;/sup&gt;</td>
<td>50 mg q 6–8 hr, up to 400 mg/day&lt;sup&gt;c&lt;/sup&gt;</td>
</tr>
<tr>
<td>IV intermittent bolus</td>
<td>Dilute to 20 mL; inject over not less than 5 min&lt;sup&gt;c&lt;/sup&gt;</td>
<td>Dilute to 5–10 mL; inject over not less than 2 min&lt;sup&gt;c&lt;/sup&gt;</td>
<td>D 20 mL; inject over not less than 5 min&lt;sup&gt;c&lt;/sup&gt;</td>
<td>D 20 mL; inject over not less than 5 min&lt;sup&gt;c&lt;/sup&gt;</td>
</tr>
<tr>
<td>IV intermittent infusion</td>
<td>Dilute to 50 mL; infuse over 15–20 min&lt;sup&gt;c&lt;/sup&gt;</td>
<td>Dilute to 100 mL; infuse over 15–30 min&lt;sup&gt;c&lt;/sup&gt;</td>
<td>D 100 mL; infuse over 15–20 min&lt;sup&gt;c&lt;/sup&gt;</td>
<td>D 100 mL; infuse over 15–20 min&lt;sup&gt;c&lt;/sup&gt;</td>
</tr>
</tbody>
</table>
| IV continuous infusion | 37.5 mg/hr (900 mg/day); adjust to patient needs; up to 600 mg/hr has been given<sup>c</sup> | 1.67 mg/hr<sup>b</sup> (40 mg/day); adjust to patient needs<sup>c</sup> | D 6.25 mg/hr (150 mg/day); adjust to patient needs; up to 220 mg/hr has been given<sup>c</sup> | (continued)
**IV for prevention of upper GI bleeding in critically ill patients** (cimetidine) 50 mg/hr by continuous infusion. In high-risk surgical patients, adjust the dose and/or frequency of intermittent IV therapy or the rate of continuous infusion to maintain the intragastric pH above 4.0.

- Heavy smokers with ulcers larger than 1 cm in diameter.
- Nonlabeled indication and dosage.
- Pathologic hypersecretory states, intractable ulcers, or patients unable to take oral medication.
- Nonlabeled route of administration.
- Loading dose can be given but appears to offer little advantage.

*From references 2, 3, 8, and 11–15.*

**Special Populations. Pediatric Dosage.** Except for ranitidine, the safety and efficacy are not well established.

<table>
<thead>
<tr>
<th></th>
<th>CIMETIDINE</th>
<th>FAMOTIDINE</th>
<th>NIZATIDINE</th>
<th>RANITIDINE</th>
</tr>
</thead>
<tbody>
<tr>
<td>Neonates</td>
<td>5–10 mg/ kg/day.</td>
<td>1–1.2 mg/kg/day.</td>
<td>Unknown.</td>
<td>0.5–3 mg/kg/day.</td>
</tr>
<tr>
<td>Children</td>
<td>20–40 mg/ kg/day.</td>
<td>0.5–2 mg/kg/day.</td>
<td>6–10 mg/kg/day.</td>
<td>2–4 mg/kg/day.</td>
</tr>
</tbody>
</table>

- Duodenal ulcer and gastric ulcer.
- Gastroesophageal reflux disease and esophagitis.

*From references 3 and 16–19.*

**Geriatric Dosage.** Reduce dosage based on renal function.

*Other Conditions.*

<table>
<thead>
<tr>
<th></th>
<th>CIMETIDINE</th>
<th>FAMOTIDINE</th>
<th>NIZATIDINE</th>
<th>RANITIDINE</th>
</tr>
</thead>
<tbody>
<tr>
<td>Renal impairment</td>
<td>$Cl_r$ 15–30 mL/min: 600 mg/day; $&lt;15$ mL/min: 300–400 mg/day.</td>
<td>$Cl_f$ $&lt;10$ mL/min; 20 mg/day or 20 mg every other day.</td>
<td>$Cl_f$ 20–50 mL/min; 150 mg/day; $&lt;20$ mL/min: 150 mg every other day.</td>
<td>$Cl_f$ $&lt;50$ mL/min: PO 150 mg/day or IM/IV 50 mg q 12–24 hr.</td>
</tr>
</tbody>
</table>

*Use the lowest dosage that permits an adequate response; further dosage reduction of cimetidine, famotidine, or ranitidine may be necessary with concomitant severe liver disease. Because only small amounts of H₂-receptor antagonists are removed by hemodialysis and peritoneal dialysis, additional doses may not be necessary; adjust dosage schedule so that the time of the scheduled dose coincides with the end of dialysis.

*From references 11 and 13.*
Dosage Forms.

<table>
<thead>
<tr>
<th>CIMETIDINE</th>
<th>FAMOTIDINE</th>
<th>NIZATIDINE</th>
<th>RANITIDINE</th>
</tr>
</thead>
<tbody>
<tr>
<td>Tab 100, 200, 300, 400, 800 mg.</td>
<td>Tab 10, 20, 40 mg.</td>
<td>Cap 150, 300 mg.</td>
<td>Tab 75, 150, 300 mg.</td>
</tr>
<tr>
<td>Chew Tab 10 mg</td>
<td>Chew Tab 10 mg</td>
<td></td>
<td>Tab (effervescent) 150 mg</td>
</tr>
<tr>
<td>(Pepcid Complete)</td>
<td></td>
<td>Granules (effervescent) 150 mg</td>
<td></td>
</tr>
<tr>
<td>Tab (rapid dissolving) 20, 40 mg.</td>
<td></td>
<td>Cap 150, 300 mg.</td>
<td></td>
</tr>
<tr>
<td>Soln 60 mg/mL.</td>
<td>Susp 8 mg/mL.</td>
<td>Syrup 15 mg/mL.</td>
<td></td>
</tr>
<tr>
<td>Inj 6 mg/mL (premixed)</td>
<td>Inj. 0.4 mg/mL (premixed)</td>
<td>Inj 0.5 (premixed), 25 mg/mL.</td>
<td></td>
</tr>
<tr>
<td>Inj 150 mg/mL.</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

aDissolve dose in approximately 180–240 mL (6–8 fl oz) of water before drinking.
bContains calcium carbonate 800 mg and magnesium hydroxide 165 mg.
cDiscard reconstituted suspension after 30 days.
dStore at 2–8°C (36–46°F).

Patient Instructions. The effectiveness of H₂-receptor antagonists in peptic ulcer disease might be decreased by cigarette smoking. Discontinue or decrease smoking or avoid smoking after the last dose of the day. Antacids can be used as needed for relief of epigastric pain. Even though ulcer or gastroesophageal reflux disease symptoms might improve, continue treatment for the duration of therapy unless instructed otherwise. If symptomatic relief is not obtained in 2 weeks with over-the-counter medication, contact your health care practitioner. Report any bleeding, vomiting, or severe esophageal or abdominal pain.

Missed Doses. If you miss a dose, take it as soon as possible. If it is almost time for your next dose, skip the missed dose and return to your usual dosage schedule. Do not double doses.

Pharmacokinetics.

Onset.
All agents have an oral onset of 1 hr and an IV onset of 15 min.

Serum Levels.

<table>
<thead>
<tr>
<th>CIMETIDINE</th>
<th>FAMOTIDINE</th>
<th>NIZATIDINE</th>
<th>RANITIDINE</th>
</tr>
</thead>
<tbody>
<tr>
<td>EC₅₀</td>
<td>625 ± 375 µg/L</td>
<td>11 ± 2 µg/L</td>
<td>167 ± 13 µg/L</td>
</tr>
</tbody>
</table>

Fate.

<table>
<thead>
<tr>
<th>Oral bio-availability</th>
<th>60 ± 20%</th>
<th>41 ± 4%</th>
<th>95 ± 5%</th>
<th>55 ± 25%</th>
</tr>
</thead>
</table>

(continued)
ACID-Peptic Therapy

<table>
<thead>
<tr>
<th></th>
<th>CIMETIDINE</th>
<th>FAMOTIDINE</th>
<th>NIZATIDINE</th>
<th>RANITIDINE</th>
</tr>
</thead>
<tbody>
<tr>
<td>$V_d$</td>
<td>1 ± 0.2 L/kg</td>
<td>1.2 ± 0.3 L/kg</td>
<td>1.4 ± 0.2 L/kg</td>
<td>1.6 ± 0.4 L/kg</td>
</tr>
<tr>
<td>Protein binding</td>
<td>20 ± 6%</td>
<td>16%</td>
<td>30 ± 5%</td>
<td>15%</td>
</tr>
<tr>
<td>Excreted unchanged in urine</td>
<td>75%</td>
<td>65–70%</td>
<td>70 ± 5%</td>
<td>68–79%</td>
</tr>
<tr>
<td>$t_{1/2}$ Normal</td>
<td>1.9 ± 0.4 hr</td>
<td>3 ± 0.5 hr</td>
<td>1.4 ± 0.2 hr</td>
<td>2 ± 0.4 hr</td>
</tr>
<tr>
<td>Anuric</td>
<td>4.5 ± 0.5 hr</td>
<td>20+ hr</td>
<td>7.2 ± 1.3 hr</td>
<td>7 ± 3 hr</td>
</tr>
</tbody>
</table>

*EC$_{50}$ is the serum concentration necessary to inhibit pentagastrin-stimulated secretion of acid by 50%.

From references 2, 11, 12, and 21.

**Adverse Reactions.** Adverse reactions are generally mild. The most frequent adverse events occur in 1–7% of patients and include headache, diarrhea, constipation, dizziness, drowsiness, and fatigue. Reversible confusional states, depression, agitation, and other CNS manifestations can occur occasionally with all H$_2$-receptor antagonists, predominantly in severely ill patients or those with renal and/or hepatic disease or advanced age. Reversible dose-dependent increases in ALT have been reported with IV cimetidine and IV ranitidine; administration over 15–30 min minimizes this effect. Rare cases of fatal hepatic disease with and without jaundice have been reported with cimetidine and ranitidine.

**Precautions.** Pregnancy; lactation. Dosage reduction might be required in severe renal and/or hepatic failure. Dosage reduction might not be required in patients treated with cimetidine because this drug can increase Cr, by competing for renal tubular secretion. This effect should not be interpreted as renal dysfunction. Symptomatic response to therapy does not preclude the possibility of gastric or esophageal malignancy.

**Drug Interactions.** Cimetidine inhibits hepatic CYP1A2, CYP2C8–10, CYP2D6, and CYP3A3–5; ranitidine inhibits CYP2D6 and CYP3A3–5 to a much lesser ex-
tent. Clinically important interactions with drugs metabolized by these iso-enzymes can occur (the most important of which are carbamazepine, chlordiazepoxide, clozapine, diazepam, glipizide, lidocaine, phenytoin, propranolol, theophylline, tolbutamide, tricyclic antidepressants, quinidine, tacrine, and warfarin). Hepatic microsomal enzyme interactions with cimetidine are dose dependent.\textsuperscript{22,23} Controversy remains about interactions with ranitidine, although in general they seem less likely and less severe than with cimetidine.\textsuperscript{11,22,23} Cimetidine can inhibit the elimination of certain drugs secreted by renal tubules (eg, procainamide).\textsuperscript{22,23} Nizatidine has been reported to increase serum salicylate concentrations in patients on high aspirin doses (3.9 g/day). Cimetidine, ranitidine, and nizatidine (but not famotidine) inhibit gastric mucosal alcohol dehydrogenase; the clinical importance of this interaction is uncertain.\textsuperscript{24,25} Elevations in gastric pH can alter the rate or extent of absorption of ketoconazole, itraconazole, and other drugs whose dissolution and absorption are pH dependent.\textsuperscript{22,23}

**Parameters to Monitor.** Improvement in epigastric pain or heartburn. However, pain relief in PUD and GERD does not correlate directly with endoscopic evidence of healing. Monitor Crs, CBC, AST, ALT, and CNS status periodically. In patients receiving IV doses of cimetidine (≥2.4 g/day) or ranitidine (≥400 mg/day), it is advisable to monitor serum transaminases routinely throughout IV therapy. When the drug is used to prevent upper GI bleeding in critically ill patients, measure the intragastric pH periodically. Monitor for potential drug interactions.

**Notes.** In general, the H\textsubscript{2}-receptor antagonists are similar in efficacy when conventional dosages are prescribed for the treatment of gastric and duodenal ulcers and for maintenance of healing of duodenal ulcer. Ulcer healing rates are similar to those with sucralfate or aggressive antacid therapy.\textsuperscript{2,12,13} The H\textsubscript{2}-receptor antagonists are often used in combination with a number of antibiotics to eradicate *Helicobacter pylori* in peptic ulcer disease. (See Eradication of *Helicobacter pylori* Infection.) Usual dosages of H\textsubscript{2}-receptor antagonists are less effective than misoprostol or proton pump inhibitors in preventing NSAID-induced gastric ulcer.\textsuperscript{26,27} However, high-dose famotidine can be effective in preventing and healing NSAID-induced gastric and duodenal ulcers.\textsuperscript{28–30} Although all H\textsubscript{2}-receptor antagonists provide symptomatic relief and esophageal healing, higher dosages are required in patients with moderate to severe esophagitis than those used in patients with mild GERD symptoms.\textsuperscript{8,12,15} Intermittent administration or continuous infusion of IV cimetidine, ranitidine, or famotidine is more effective than placebo in preventing upper GI bleeding in critically ill patients.\textsuperscript{3} The maintenance of intragastric pH above 4.0 does not conclusively prevent upper GI bleeding.\textsuperscript{3} Although it is easier to maintain the intragastric pH above 4.0 by continuous infusion, the superiority of continuous infusion of the H\textsubscript{2}-receptor antagonists in preventing upper GI bleeding in the critically ill patient has not been established; intermittent administration and continuous infusion are at least as effective as sucralfate or aggressive antacid therapy.\textsuperscript{3} Combination of an H\textsubscript{2}-receptor antagonist with sucralfate provides two different mechanisms of drug action and might be beneficial. However, enhanced efficacy of two drugs compared with single-drug therapy has not been substantiated in controlled trials in patients with duodenal
ulcer, gastric ulcer, or GERD or when used to prevent or treat upper GI bleeding. Coadministration of an H2-receptor antagonist with a proton pump inhibitor is without established benefit and might compromise the action of the proton pump inhibitor.2 Controlled trials have not demonstrated that H2-receptor antagonists are of benefit in patients with active upper GI bleeding.11 The relation of H2-receptor antagonist therapy to the development of nosocomial pneumonia in critically ill patients is inconclusive.3 (See Sucralfate.) Cimetidine can augment cell-mediated immunity by blockade of H2-receptors on suppressor T lymphocytes; it remains difficult to determine whether this effect is clinically useful or potentially dangerous, especially after organ transplantation and in autoimmune disorders.22 Cimetidine can reduce pain and hasten resolution of herpes zoster lesions.31 Although it is not certain whether this immune system action is class specific or drug specific, it appears to be related to the cimetidine molecule. All H2-receptor antagonists are stable in D5W, D10W, NS, LR, parenteral nutrition, or 5% sodium bicarbonate for 48 hr at room temperature.

MISOPROSTOL Cytotec

Pharmacology. Misoprostol is a synthetic prostaglandin E1 analogue that inhibits gastric acid secretion and enhances gastric mucosal defense. Antisecretory effects are dose dependent over the range of 50–200 μg; cytoprotective effects occur at doses of 200 μg or more. The gastric ulcer protective effect of misoprostol appears to plateau between 200 μg bid–tid, but no dose–response effect is apparent in preventing duodenal ulcers.30,32 Cotherapy with misoprostol reduces the frequency of NSAID-induced complications, including GI perforation, obstruction, and bleeding, but its cost effectiveness remains controversial.30,33 Misoprostol also causes uterine contraction.

Administration and Adult Dosage. PO for GI protection during NSAID therapy 200 μg qid with food; if this dosage is not tolerated, 100 μg qid can be used. Lower-dosage regimens of misoprostol 200 μg tid or bid appear similar in efficacy and better tolerated for protection against NSAID-induced gastric and duodenal ulcers than the 200 mg qid dosage. Dosage reduction is not required in renal impairment, hepatic failure, or the elderly. PO or Vag for use with mifepristone for pregnancy termination 400 μg 2 days after mifepristone dose. Vag for cervical ripening at term 25–100 μg.34,35 PO for cervical ripening at term 100–200 μg.34,36 (See Notes.)

Dosage Forms. Tab 100, 200 μg. Misoprostol is also available in combination with diclofenac in Arthrotec. (See Diclofenac.)

Pharmacokinetics. After oral administration, misoprostol is extensively absorbed and rapidly de-esterified to the active drug, misoprostol acid. Peak serum concentrations of misoprostol acid are reduced when the drug is taken with food. Plasma protein binding of misoprostol acid is <90%. Misoprostol acid undergoes further metabolism, but approximately 80% is excreted unchanged in urine.

Adverse Reactions. Diarrhea is reported to occur within 2 weeks of initiating therapy in 14–40% of patients on NSAIDs receiving 800 μg/day and less frequently with 400–600 μg/day. Diarrhea usually resolves in about 1 week with
continued treatment; rarely, profound diarrhea occurs in patients with inflammatory bowel disease. Abdominal pain occurs in 13–20% of patients on NSAIDs receiving misoprostol 800 µg/day, but there is no consistent difference from placebo. Antacids (except those containing magnesium) can be used for abdominal pain relief. Nausea, flatulence, headache, dyspepsia, vomiting, and constipation occur occasionally. Women who receive misoprostol occasionally develop gynecologic disorders including cramps or vaginal bleeding.

**Precautions.** Advise patients (especially those receiving concurrent corticosteroids or anticoagulants) to report bleeding, vomiting, severe abdominal pain, and diarrhea. For GI protective uses, misoprostol is contraindicated in pregnancy because of the risk of abortion and women of childbearing potential should have a negative serum pregnancy test within 2 weeks of beginning therapy, should begin treatment on the second or third day of the next menstrual period, should comply with effective contraceptive measures, and should receive oral and written warnings of the hazards of misoprostol therapy and the risk of contraceptive failure. Warn patients not to give misoprostol to others.

**Drug Interactions.** Misoprostol does not affect the hepatic cytochrome P450 microsomal enzyme system and does not interfere with the beneficial effects of NSAIDs in rheumatoid arthritis.

**Notes.** In most trials of patients receiving long-term NSAID therapy for rheumatoid arthritis, misoprostol 200 µg qid was superior to the H2-receptor antagonists or sucralfate in preventing NSAID-induced gastric ulcer; however, misoprostol did not relieve GI pain or discomfort associated with NSAID use.26,30,33 Omeprazole 20 mg/day is associated with a lower relapse rate and is better tolerated than misoprostol 200 µg bid for prophylactic treatment. Misoprostol appears to be more effective than conventional methods of cervical ripening at term;34,35 oral and vaginal administrations appear to be equivalent in efficacy.34

### PROTON PUMP INHIBITORS:

<table>
<thead>
<tr>
<th><strong>LAN SOPRA ZOLE</strong></th>
<th>Prevacid</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>OME PRAZOLE</strong></td>
<td>Prilosec</td>
</tr>
<tr>
<td><strong>PAN TOPRA ZOLE</strong></td>
<td>Protonix</td>
</tr>
<tr>
<td><strong>RABE PRA ZOLE</strong></td>
<td>Aciphex</td>
</tr>
</tbody>
</table>

**Pharmacology.** Proton pump inhibitors (PPIs) are inactive substituted benzimidazoles that, when protonated in the secretory canaliculi of the parietal cells, covalently bind to H+/K⁺-ATPase (proton pump), which is the final pathway for acid secretion. PPIs produce a profound and prolonged antisecretory effect and inhibit basal, nocturnal, pentagastrin-, and food-stimulated gastric acid secretion. Serum gastric levels increase during treatment but return to pretreatment levels within 1–2 weeks of discontinuing therapy.

**Administration and Adult Dosage.** Administer the PPI at least 30–60 min before meals, preferable in the morning, because these agents inhibit only those proton pumps that are actively secreting acid. Infuse IV pantoprazole doses over 15 min via a dedicated line and the in-line filter provided. IV dosage is the same as PO dosage.
<table>
<thead>
<tr>
<th>INDICATION</th>
<th>ESOMEPRAZOLE</th>
<th>LANSOPRAZOLE</th>
<th>OMEPRAZOLE</th>
<th>PANTOPRAZOLE</th>
<th>RABEPRAZOLE</th>
</tr>
</thead>
<tbody>
<tr>
<td>Treatment of active</td>
<td>—</td>
<td>15 mg/day. a</td>
<td>20 mg/day. a</td>
<td>40 mg/day. b</td>
<td>20 mg/day. a</td>
</tr>
<tr>
<td>duodenal ulcer (4 weeks)</td>
<td>—</td>
<td>15 mg/day. a</td>
<td>20 mg/day. b</td>
<td>20 mg/day. b</td>
<td>20 mg/day. b</td>
</tr>
<tr>
<td>Maintenance of duodenal</td>
<td>—</td>
<td>30 mg/day. a</td>
<td>40 mg/day. a</td>
<td>40 mg/day. b</td>
<td>20–40 mg/day. b</td>
</tr>
<tr>
<td>ulcer healing (1 yr)</td>
<td>—</td>
<td>15–30 mg/day. b</td>
<td>20–40 mg/day. b</td>
<td>40 mg/day. b</td>
<td>20–40 mg/day. b</td>
</tr>
<tr>
<td>Treatment of active</td>
<td>20 mg/day. a</td>
<td>15 mg/day. a</td>
<td>20 mg/day. a</td>
<td>20 mg/day. b</td>
<td>20 mg/day. a</td>
</tr>
<tr>
<td>gastric ulcer (4–8 weeks)</td>
<td>20–40 mg/day. a</td>
<td>30 mg/day. a</td>
<td>20 mg/day. a</td>
<td>40 mg/day. a</td>
<td>20 mg/day. a</td>
</tr>
<tr>
<td>Maintenance of erosive esopha</td>
<td>20 mg/day. a</td>
<td>15 mg/day. a</td>
<td>20 mg/day. a</td>
<td>20 mg/day. b</td>
<td>20 mg/day. a</td>
</tr>
<tr>
<td>gitis (4–8 weeks)</td>
<td>20–40 mg/day. a</td>
<td>60 mg/day. a</td>
<td>20 mg/day. a</td>
<td>20–40 mg/day. b</td>
<td>20 mg/day. b</td>
</tr>
<tr>
<td>PO for treatment of pathologic</td>
<td>40 mg/day. a</td>
<td>—</td>
<td>up to 90 mg bid. c</td>
<td>up to 120 mg tid. c</td>
<td>up to 120 mg tid. c</td>
</tr>
<tr>
<td>hypersecretory conditions</td>
<td>40 mg/day. a</td>
<td>30 mg bid a</td>
<td>20 mg bid a</td>
<td>20–40 mg bid b</td>
<td>20 mg bid b</td>
</tr>
<tr>
<td>Helicobacter pylori eradication</td>
<td>—</td>
<td>—</td>
<td>—</td>
<td>—</td>
<td>—</td>
</tr>
<tr>
<td>for reduction of the risk of</td>
<td>—</td>
<td>—</td>
<td>—</td>
<td>—</td>
<td>—</td>
</tr>
<tr>
<td>duodenal ulcer recurrence</td>
<td>—</td>
<td>15 mg/day. a</td>
<td>20 mg/day. b</td>
<td>40 mg/day. b</td>
<td>—</td>
</tr>
<tr>
<td>Risk reduction of NSAID-induced</td>
<td>—</td>
<td>—</td>
<td>—</td>
<td>—</td>
<td>—</td>
</tr>
<tr>
<td>gastric ulcers</td>
<td>—</td>
<td>—</td>
<td>—</td>
<td>—</td>
<td>—</td>
</tr>
</tbody>
</table>

FDA-approved regimen.

Nonlabeled indication and dosage.

Adjust dosage to patient's needs and continue as long as clinically indicated.

Combined with clarithromycin 500 mg bid and amoxicillin 1 g bid for 10 days.

Combined with clarithromycin 500 mg bid and amoxicillin 1 g bid for 7 days.

Combined with clarithromycin 500 mg bid and metronidazole 400 mg bid for 7 days.

From references 2, 8, 15, and 38–48.
Special Populations. Pediatric Dosage. Safety and efficacy not well established. 

PO for esophagitis (omeprazole) (>10 months) 0.7 mg/kg/day initially, increasing as necessary up to 3.5 mg/kg/day; PO for peptic ulcer disease in combination with antimicrobials for H. pylori (<10 yr) 0.6 mg/kg/day or 20 mg/day; (>10 yr) 20 mg daily–bid. Lansoprazole 0.45 mg/kg/day in 2 divided doses; up to 15 mg bid has been used for treatment of H. pylori in combination with antimicrobial therapy.

Geriatric Dosage. Dosage reduction is not usually necessary; reduce only if the drug is not well tolerated.

Other Conditions. Dosage adjustments of PPIs are unnecessary in renal impairment or mild to moderate liver disease. However, dosage reduction should be considered for chronic or severe hepatic impairment. Certain groups (ie, Asians) tend to be poor metabolizers, so a decrease in dose might be considered. PPIs are not readily dialyzable.

Dosage Forms.

<table>
<thead>
<tr>
<th>LANSOPRAZOLE</th>
<th>OMEPRAZOLE</th>
<th>PANTOPRAZOLE</th>
<th>RABEPRAZOLE</th>
</tr>
</thead>
<tbody>
<tr>
<td>Enteric-coated granules</td>
<td>Enteric-coated granules</td>
<td>Enteric-coated tablet</td>
<td>Delayed-release tablet</td>
</tr>
<tr>
<td>Cap 15, 30a mg.</td>
<td>Cap 10, 20, 40 mg.</td>
<td>Tab 40 mg</td>
<td>Tab 20 mg.</td>
</tr>
</tbody>
</table>

aPrevac for Helicobacter pylori therapy consists of 2 lansoprazole 30 mg capsules, 4 amoxicillin 500 mg capsules, and 2 clarithromycin 500 mg tablets in an individual daily administration pack.

Patient Instructions. Swallow capsule (lansoprazole, omeprazole) or tablet (pantoprazole, rabeprazole) whole; do not crush or chew. Take medication 30–60 minutes before meals, preferably in the morning. Capsules can be opened and the granules sprinkled on applesauce, yogurt, or apple or orange juice if you have difficulty swallowing. Do not chew, and do swallow the preparation immediately after sprinkling the content onto food. (See Notes.) The effectiveness of PPIs in peptic ulcer disease can be decreased by cigarette smoking. Even though symptoms can improve quickly, continue treatment for the duration of therapy unless instructed otherwise.

Missed Doses. If you miss a dose, take it as soon as possible. If it is almost time for the next dose, skip the missed dose and return to your usual dosage schedule. Do not double doses.

Pharmacokinetics. Onset and Duration. PO onset of antisecretory activity is 1–3 hr, with rabeprazole having the quickest onset and pantoprazole the slowest. Duration is dose dependent. Gastric acid inhibition increases with repeated daily doses, reaching a plateau after several days. Upon discontinuing the PPI, gastric secretory activity gradually returns to pretreatment level within 2–7 days. There is no indication that rebound gastric acidity occurs.
Pharmacokinetics.

<table>
<thead>
<tr>
<th></th>
<th>Lansoprazole</th>
<th>Omeprazole</th>
<th>Pantoprazole</th>
<th>Rabeprazole</th>
</tr>
</thead>
<tbody>
<tr>
<td>Oral Bioavailability (%)</td>
<td>80–85</td>
<td>30–40</td>
<td>77</td>
<td>52</td>
</tr>
<tr>
<td>Protein Binding (%)</td>
<td>97</td>
<td>95</td>
<td>98</td>
<td>96</td>
</tr>
<tr>
<td>Peak (hr)</td>
<td>1.7</td>
<td>0.5–3.5</td>
<td>2.4</td>
<td>2.9–3.8</td>
</tr>
<tr>
<td>(V_d) (L/kg)</td>
<td>0.4</td>
<td>0.13–0.35</td>
<td>0.15</td>
<td>0.34</td>
</tr>
<tr>
<td>(t_{1/2}) (hr)</td>
<td>1.5–1.7</td>
<td>0.5–1</td>
<td>1–1.9</td>
<td>1–2</td>
</tr>
<tr>
<td>Urinary Excretion (%)</td>
<td>33</td>
<td>77</td>
<td>71</td>
<td>90</td>
</tr>
</tbody>
</table>

From references 2, 21, and 38–44.

Adverse Effects. All have similar short-term (<12 weeks) and long-term (>12 weeks) side effect profiles. The most frequent short-term adverse effects are headache, diarrhea, nausea, and abdominal pain. Flu-like symptoms, constipation, fatigue, malaise, muscle cramps, joint pain, myalgia, anxiety, skin rash, confusion, sleep disturbances, and taste perversion occur occasionally. Anaphylactic reactions, gynecomastia, hemolytic anemia, thrombocytopenia, and psychic disturbances occur rarely. Rare cases of severe skin reactions (eg, Stevens–Johnson syndrome, toxic epidermal necrolysis), hepatic failure, cholestatic jaundice, pancreatitis, interstitial nephritis, and agranulocytosis have occurred. Long-term use of PPIs have been thought to cause gastric cancer, gastric enterochromaffin cell hyperplasia, carcinoid tumors, bacterial overgrowth, atrophic gastritis, and decreased absorption of certain nutrients; however, recent studies have shown that the risk of such adverse effects is not increased. Patients infected with *H. pylori* are at greater risk for atrophic gastritis. Patients older than 65 yr have similar side effect profiles as younger individuals.

Precautions. Pregnancy; there are sporadic reports of congenital abnormalities in infants born to women who took omeprazole during pregnancy. Symptomatic response to PPI therapy does not preclude the possibility of gastric malignancy.

Drug Interactions. Elevations in gastric pH can increase the extent of absorption of ampicillin and pancreatic enzyme supplements and decrease the rate or extent of absorption of digoxin, itraconazole, iron salts, ketoconazole, and other drugs or dosage forms that are pH dependent. PPIs are metabolized to different degrees via the CYP450 isoenzymes CYP3A4, CYP2C19, CYP1A2, and CYP2C9. Lansoprazole, pantoprazole, and rabeprazole do not increase diazepam, (R)-warfarin or phenytoin concentrations, but these medications are affected by omeprazole because of its extensive metabolism via the hepatic CYP2C19 isoenzyme. Lansoprazole can increase the clearance of theophylline by 10%. Rabeprazole does not interact with phenytoin, warfarin, or theophylline; however, rabeprazole causes a 20% increase in serum digoxin trough levels. Pantoprazole has no interaction with warfarin, phenytoin, diazepam, or theophylline, even though it is metabolized via CYP2C19. Absorption of PPIs is not affected by antacids.
Parameters to Monitor. Improvement in epigastric pain or heartburn; however, pain relief in PUD and GERD does not correlate directly with endoscopic evidence of healing. Monitor for potential drug interactions and adverse effects. Monitor laboratory values, including liver function tests, CBC, and SMA-7. Assess the indication, dosage, and duration of PPI therapy, especially as it relates to the need for treatment beyond 16 weeks. Monitor serum vitamin B₁₂ concentrations every few years in patients on long-term PPI therapy, especially the elderly.55

Notes. PPIs are the drugs of choice for erosive esophagitis and Zollinger–Ellison syndrome. Standard dosages provide more rapid relief of symptoms and ulcer or esophageal healing than standard dosages of H₂-receptor antagonists.8,15,38 Patients with gastric or duodenal ulcers or esophagitis refractory to H₂-receptor antagonists are likely to respond to PPIs, but the rate of recurrence after discontinuation is similar to that of H₂-receptor antagonists.8,15,38 NSAID-induced gastric and duodenal ulcers can be prevented or treated by PPIs and are superior to and have a better side effect profile than misoprostol 200 μg bid or ranitidine 150 mg bid.27,37,38,56 IV pantoprazole is no more effective than oral PPIs.

Coadministration of a PPI with an H₂-receptor antagonist or sucralfate is without established benefit. Lansoprazole and omeprazole are available as gelatin capsules that are formulated as enteric-coated granules from which the drug is released when pH rises above 6. It is important for patients not to chew or crush the capsules or enteric-coated granules because the protective enteric coating might be destroyed and thus decrease the drug’s bioavailability.57 Patients who have difficulty swallowing capsules or have feeding tubes can open the capsule and mix the granules with apple or orange juice. Granules also can be sprinkled onto applesauce for oral administration. Lansoprazole granules can be sprinkled onto Ensure pudding, cottage cheese, yogurt, or strained pears. Omeprazole and lansoprazole can be mixed with sodium bicarbonate to make a simplified suspension administered through feeding tubes.57

Esomeprazole magnesium (Nexium) is the (S)-isomer of omeprazole that is as effective as omeprazole in controlling pH and produces longer-lasting gastric acid suppression. Doses of 20–40 mg are effective for symptomatic relief of GERD, H. pylori infections, and erosive esophagitis healing and maintenance.58–60 Side effects are similar to other PPIs and include diarrhea, abdominal pain, flatulence, headaches, and nausea. It is available as 20 and 40 mg delayed-release tablets.

Pharmacology. Sucralfate is an aluminum hydroxide salt of a sulfated disaccharide. Its exact mechanism of action is not known. It forms an ulcer-adherent complex with proteinaceous exudates at the ulcer site, thereby protecting against further attack by acid, pepsin, and bile salts. Adherence to the ulcer crater is enhanced at pH <3.5. Sucralfate inhibits pepsin activity; a dose of 1 g has about 14–16 mEq of acid-neutralizing capacity. The aluminum moiety stimulates endogenous prostaglandins and binds bile salts and phosphate in the GI tract.2,61,62

Administration and Adult Dosage. PO for short-term treatment of duodenal ulcer 1 g qid on an empty stomach, 1 hr before meals and at bedtime, or 2 g bid for 4–8 weeks.2,61,62 PO for maintenance of healed duodenal ulcer 1 g bid. PO
for short-term treatment of active benign gastric ulcer 1 g qid.\textsuperscript{2,61,62} PO for treatment of symptomatic GERD or erosive esophagitis 1 g qid.\textsuperscript{61,62} PO or NG for prevention of upper GI bleeding in critically ill patients 1 g q 4–6 hr.\textsuperscript{3,61,62} PO or NG for phosphate binding in renal failure titrate dosage based on serum phosphate.\textsuperscript{2,61}

**Special Populations.** **Pediatric Dosage.** Safety and efficacy not well established. PO 40–80 mg/kg/day in 4 divided doses. Alternatively, (<10 kg) 0.5 g q 6 hr; (>10 kg) 1 g q 6 hr.\textsuperscript{63}

**Geriatric Dosage.** Dosage reduction usually not necessary.

**Dosage Forms.** Tab 1 g; Susp 100 mg/mL.

**Patient Instructions.** Take this drug with water on an empty stomach 1 hr before each meal and at bedtime. Antacids can be used as needed for pain relief but do not take them within 30 minutes before or after taking sucralfate. Take potentially interacting drugs 2 hours before taking sucralfate to avoid or minimize drug interactions. Even though symptoms can decrease, continue treatment for the duration of therapy unless instructed otherwise.

**Missed Doses.** If you miss a dose, take it as soon as possible. If it is almost time for your next dose, skip the missed dose and return to your usual dosage schedule. Do not double doses.

**Pharmacokinetics.** **Onset and Duration.** Onset (attachment of sucralfate to ulcer site) is within 1 hr; duration is about 6 hr.

**Fate.** Sucralfate is only minimally absorbed from the GI tract and is excreted primarily in the feces. About 3–5% (primarily aluminum) is absorbed and excreted in urine.\textsuperscript{2,61} Aluminum excretion is decreased in uremia.

**Adverse Reactions.** Adverse reactions are usually minor and occur in about 5% of patients. Constipation occurs in about 2% of patients. Other effects, including diarrhea, nausea, gastric discomfort, indigestion, dry mouth, rash, pruritus, backache, dizziness, drowsiness, vertigo, and a metallic taste, occur occasionally. Aluminum accumulation and toxicity, including osteodystrophy, osteomalacia, encephalopathy, and seizures, have been reported in patients with chronic renal failure. Hypophosphatemia can develop in critically ill patients and those on prolonged sucralfate therapy. Bezoar formation in the esophagus and GI tract and intestinal obstruction and perforation have been reported.\textsuperscript{3,61}

**Precautions.** Use with caution in patients receiving other aluminum-containing drugs or in chronic renal failure and dialysis. Avoid administration through feeding tubes because the drug can occlude the tube.

**Drug Interactions.** Sucralfate can inhibit the absorption of drugs including digoxin, ketoconazole, levothyroxine, phenytoin, quinidine, oral fluoroquinolones, tetracyclines, theophylline, and warfarin. In most cases, drug interactions can be avoided if the drug is given 2 hr before sucralfate administration, especially in patients receiving tube feedings.

**Parameters to Monitor.** Improvement in epigastric pain or heartburn; however, pain relief in PUD and GERD does not correlate directly with endoscopic evidence of healing. Monitor for constipation and signs of aluminum toxicity in the
elderly, in chronic renal failure, or in patients receiving other aluminum-containing drugs. Obtain serum phosphate levels periodically in patients receiving concurrent aluminum-containing drugs or with prolonged use. Monitor for potential drug interactions.

Notes. Sucralfate is as effective as standard doses of the H₂-receptor antagonists in the short-term treatment and maintenance of healed duodenal ulcer.²⁵,²⁶ Sucralfate can overcome the negative effect of cigarette smoking on duodenal ulcer healing and recurrence.⁶¹ Its efficacy in healing erosive esophagitis and maintaining esophageal healing is inferior to the H₂-receptor antagonists or proton pump inhibitors. Sucralfate is effective in preventing stress-related mucosal bleeding and can be more cost effective than H₂-receptor antagonists.³,⁶²,⁶⁴ Its efficacy as a single agent in preventing NSAID-induced gastric and duodenal ulcers, chemotherapy-induced stomatitis, and stress-related bleeding in high-risk critically ill surgical patients is unsubstantiated. Whether sucralfate is associated with a lower frequency of nosocomial pneumonia in critically ill patients than H₂-receptor antagonists or antacids remains controversial.³,⁶⁵ Although therapy with sucralfate and an H₂-receptor antagonist or PPI provides two different mechanisms of drug action, enhanced efficacy of two drugs has not been substantiated for any indication.

ERADICATION OF HELICOBACTER PYLORI INFECTION

One cause of peptic ulcer disease (PUD) is associated with H. pylori infections. Patients infected or colonized with H. pylori are at increased risk for developing PUD, gastric carcinoma, atrophic gastric and gastric mucosa-associated lymphoid tissue.⁶⁶,⁶⁷ Thus, all patients who test positive for H. pylori should be treated.⁶⁷,⁶⁸ The value of H. pylori eradication in patients with dyspepsia or nonulcer dyspepsia remains controversial.⁶⁹,⁷⁰ The goal of therapy is to promote rapid ulcer healing and prevent relapse by eradicating the infection. Combination therapy that includes an antisecretory agent (PPI or ranitidine bismuth citrate) and two antimicrobial agents (amoxicillin and clarithromycin or metronidazole) for 10–14 days is effective in resolving the infection.³⁸,⁴⁴,⁶⁷ The use of any PPI combined with at least two antibiotics have achieved similar eradication rates against H. pylori infections.³⁸,⁴⁴,⁴⁷,⁴⁸

Factors to consider when choosing a H. pylori regimen include eradication rates, patient compliance, and minimizing drug resistance and adverse effects associated with the drug therapy. Dual therapy (PPI and one antibiotic) is rarely used because eradication rates are often poor (<70%), but triple therapy is used because it obtains an eradication rate of at least 80–90%. The eradication rate for quadruple therapy is >90%.⁵⁸ Even though quadruple therapy is effective in eliminating the infection, it is not ideal because of the complicated dosage regimen that can lead to decreased patient compliance. Adverse effects can decrease patient compliance, especially those treated with metronidazole.

When choosing antibiotics, resistance becomes an issue. Metronidazole resistance to H. pylori infections is most common, with a range of 7–80%, and is more frequent in women. Macrolide resistance is less common (1–10%) and is even less frequent with tetracycline and amoxicillin.⁷¹ Quadruple therapy should be considered for patients who failed initial treatment.
Antibiotics should not be used for longer than 2 weeks. If treatment fails, a different antibiotic regimen should be considered. Although any PPI can be used in the various regimens, there are some substitutions that should not be done (e.g., ampicillin for amoxicillin, doxycycline for tetracycline, azithromycin for clarithromycin, or an H2-receptor antagonist for a PPI).38,44,47,48
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<thead>
<tr>
<th>DRUGS</th>
<th>DOSE</th>
<th>FREQUENCY</th>
<th>DURATION</th>
<th>EFFICACY</th>
<th>ADVERSE EFFECTS</th>
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**DRUG TREATMENT REGIMENS USED TO ERADICATE *HELICOBACTER PYLORI* (continued)**

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<th>DOSE</th>
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<th>COMPLIANCE&lt;sup&gt;c&lt;/sup&gt;</th>
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BSS = bismuth subsalicylate; H<sub>2</sub>RA = H<sub>2</sub>-receptor antagonist; PPI = proton pump inhibitor; RBC = ranitidine bismuth citrate.

<sup>a</sup>Efficacy (eradication rate): excellent >90%; good >80–90%; fair >70–80%; poor <70%.

<sup>b</sup>Adverse Effects = frequency of clinically important adverse effects.

<sup>c</sup>Compliance = estimate based on total number of tablets/capsules, frequency of administration, and clinically important adverse effects.

<sup>d</sup>Any PPI can be used (esomeprazole 40 mg, lansoprazole 30 mg, omeprazole 20 mg, pantoprazole 20–40 mg, rabeprazole 40 mg).

<sup>e</sup>PPI therapy can be extended to 28 days in patients with active ulcer.

<sup>f</sup>FDA-approved regimen.
Antiemetics

DOLESETRON MESYLATE

**Pharmacology.** Dolasetron and its active metabolite, hydrodolasetron, are selective serotonin3 (5-HT3) antagonists. Dolasetron is approved for the prevention of chemotherapy-induced nausea and vomiting and for the prevention and treatment of postoperative nausea and vomiting. Its use is similar to those of ondansetron and granisetron.72 (See Antiemetic Drugs Comparison Chart.)

**Adult Dosage.** PO or IV for chemotherapy-induced nausea and vomiting 100 mg or 1.8 mg/kg. IV for postoperative nausea and vomiting 12.5 mg. PO for postoperative nausea and vomiting 100 mg. IV doses can be administered over a minimum of 30 sec or further diluted in 50 mL of NS or D5W and infused over a period of up to 15 min. In some dose-finding trials in adults, higher response rates were obtained with 200 mg PO and 2.4 mg IV than with lower doses by the respective routes.72

**Pediatric Dosage.** PO or IV for chemotherapy-induced nausea and vomiting (≤2 yr) safety and efficacy not established; (2–16 yr) 1.8 mg/kg, to a maximum of 100 mg. IV for postoperative nausea and vomiting (2–16 yr) 0.35 mg/kg, to a maximum of 12.5 mg. PO for postoperative nausea and vomiting (2–16 yr) 1.2 mg/kg, to a maximum of 100 mg.

**Dosage Forms.** Inj 20 mg/mL; Tab 50, 100 mg.

**Pharmacokinetics.** Approximately 75% of dolasetron mesylate is dolasetron base. The apparent absolute bioavailability of oral dolasetron is approximately 75%. Little dolasetron is detected in the plasma because of rapid conversion to hydrodolasetron by the ubiquitous enzyme, carbonyl reductase. Hydrodolasetron is partly metabolized in the liver and 61% is excreted unchanged in the urine. Hydrodolasetron has a Vd of 5.8 ± 1.5 L/kg and Cl of 0.56 ± 0.16 L/hr/kg. Its half-life is 7.3 ± 1.8 hr.

**Adverse Reactions.** Acute, reversible ECG changes (PR and QTc prolongation, and QRS widening) have occurred in clinical trials and in healthy volunteers. Other adverse effects are similar to those of ondansetron and granisetron.

**Notes.** Dolasetron can be prepared extemporaneously as an oral solution by mixing the injectable form in apple or apple–grape juice. The diluted product can be kept up to 2 hr at room temperature before use.

**DRONABINOL**

**Pharmacology.** Dronabinol (Δ9-tetrahydrocannabinol) is an active antiemetic component of Cannabis. Its mechanism of action as an antiemetic is complex and poorly understood but it probably inhibits the chemoreceptor trigger zone in the medulla.

**Administration and Adult Dosage.** PO as an antiemetic 5 mg/m² 1–3 hr before chemotherapy and then q 2–4 hr after chemotherapy, for a total of 4–6 doses/day. Dosage can be increased in 2.5 mg/m² increments, to a maximum of 15 mg/m²/dose.73–75 PO for appetite stimulation 2.5 mg before lunch and dinner.
or 2.5 mg at bedtime if unable to tolerate daytime administration. Dosage can be increased to a maximum of 20 mg/day.

**Special Populations.**

**Pediatric Dosage.** PO as an antiemetic during cancer chemotherapy same as adult dosage in mg/m².

**Geriatric Dosage.** Same as adult dosage. (See Adverse Reactions.)

**Dosage Forms.** Cap 2.5, 5, 10 mg.

**Patient Instructions.** This drug can cause drowsiness or changes in mood. Until the extent of this effect is known, use caution when driving, operating machinery, or performing other tasks requiring mental alertness. Avoid excessive concurrent use of alcohol or other drugs that cause drowsiness. Store this medication in the refrigerator.

**Missed Doses.** Take this drug at regular intervals. If you miss a dose, take it as soon as you remember. If it is about time for the next dose, take that dose only. Do not double the dose or take extra.

**Pharmacokinetics.**

**Onset and Duration.** Oral onset 30–60 min; peak 2–4 hr; duration is 4–6 hr but can be longer in those who have not previously used the drug.⁷⁶,⁷⁷ Cannabis smoking onset 15 sec–2 min; peak 8–16 min; duration 3–12 hr.⁷⁸

**Fate.** Bioavailability is 4–12% orally, 2–50% by smoking. About 95% bound to plasma proteins. Vd is 8.9 ± 4.2 L/kg; Cl is 0.21 ± 0.054 L/hr/kg. Primarily metabolized by hydroxylation to active and inactive metabolites. Ultimately, 35% of metabolites are found in feces and 15% in urine, with <1% excreted unchanged in urine.⁷⁹–⁸¹

**t₁/₂.** Terminal phase 32 ± 12 hr, although time course of effects more closely parallels initial distribution phase.⁷⁹

**Adverse Reactions.** Euphoria, dizziness, paranoia, or drowsiness occur frequently and might be accompanied by ataxia, loss of balance, and disorientation to the point of being disabling. Other frequent side effects are dry mouth, orthostatic hypotension, and conjunctival injection.⁸² The "high" experienced by some is not always well tolerated, especially by older patients.⁸³

**Contraindications.** Allergy to dronabinol, marijuana, or sesame oil; mentally ill patients.

**Precautions.** Avoid during lactation. Use with caution in patients with hypertension or heart disease. Use with caution in patients with epilepsy.⁷³

**Drug Interactions.** Not well studied, but some apparent interactions have been reported after Cannabis use: additive or supra-additive sedation with alcohol and other CNS depressants; additive hypertension and/or tachycardia with anticholinergics, antihistamines, sympathomimetics, or tricyclic antidepressants; and hypomania with disulfiram or fluoxetine.

**Parameters to Monitor.** Observe for frequency of emesis, drowsiness, or disorientation.

**Notes.** Dronabinol is at least as effective as phenothiazines for chemotherapy-induced nausea and vomiting⁷⁵ but not as effective as serotonin antagonists or IV metoclopramide.⁸⁴,⁸⁵ It is not particularly effective for cisplatin-induced nausea.
and vomiting. It might not be as effective as smoking *Cannabis*, which is easier to titrate.85 (See Antiemetic Drugs Comparison Chart.)

**Granisetron**

**Pharmacology.** Granisetron is a selective antagonist at the 5-HT3 receptor used for the prevention of nausea and vomiting associated with cancer chemotherapy. Its use in cancer chemotherapy is similar to that of ondansetron, and its efficacy and side effects are comparable to those of ondansetron.72,86 (See Antiemetic Drugs Comparison Chart.)

**Adult Dosage.** IV for cancer chemotherapy-induced nausea and vomiting 10 µg/kg administered in 20–50 mL NS or D5W over 5 min, 30 min before the start of chemotherapy. **PO for cancer chemotherapy-induced nausea and vomiting** 1 mg up to 1 hr before chemotherapy and additional 1 mg doses at 12-hr intervals thereafter while receiving chemotherapy.

**Pediatric Dosage.** IV (>2 yr) same as adult dosage, but this dosage might not be adequate for children.86

**Dosage Forms.** Inj 1 mg/mL; Tab 1 mg.

**Pharmacokinetics.** Oral absorption is approximately 60%. Vd is 30 ± 1.5 L/kg; Cl is 0.060 ± 0.54 L/hr/kg. Elimination is mostly by hepatic metabolism, with 16 ± 14% appearing in the urine as unchanged drug. The half-life is 5.3 ± 3.5 hr (can be longer in cancer patients than in normals). The metabolism of granisetron might be changed by inducers or inhibitors of the cytochrome P450 system, but dosage adjustment is not recommended.

**Adverse Reactions.** (See Ondansetron).

**Ondansetron Hydrochloride**

**Pharmacology.** Ondansetron is a selective antagonist at the 5-HT3 receptor used for the prevention of nausea and vomiting associated with cancer chemotherapy, especially cisplatin, and for postoperative nausea and vomiting. It also is useful for radiotherapy-induced nausea and vomiting. Ondansetron is thought to block these receptors at both peripheral sites in the GI tract and within the area postrema in the CNS.87 It is not a dopamine receptor antagonist, so it has no extrapyramidal side effects. (See Notes.)

**Administration and Adult Dosage.** IV for chemotherapy-induced nausea or vomiting 0.15 mg/kg for 3 doses (30 min before chemotherapy and then 4 and 8 hr after) or 0.45 mg/kg, to a maximum of 32 mg as a single dose or 8 mg IV as a single dose for cisplatin doses <100 mg/m².72 (See Notes.) Infuse slowly over 15 min in 50 mL D5W or NS. **IV bolus for postoperative nausea or vomiting** 4 mg over 2–5 min before induction or postoperatively. **PO for chemotherapy-induced nausea or vomiting** 8 mg bid–tid or 24 mg once daily. **PO for radiotherapy-induced nausea or vomiting** 8 mg daily–tid. **PO for postoperative nausea or vomiting** 8–16 mg 1 hr before surgery.72

**Special Populations.** **Pediatric Dosage.** IV for chemotherapy-induced nausea or vomiting (<2 yr) safety and efficacy not established; (2–18 yr) same as adult dosage or 0.15 mg/kg for 2 doses for moderately emetogenic chemotherapy.88–90
IV for postoperative nausea or vomiting 0.05–0.1 mg/kg, to a maximum of 4 mg as a single dose over 30 sec before the surgical incision.\textsuperscript{72} PO for chemotherapy-induced nausea and vomiting (4–11 yr) 4 mg q 8 hr. PO for postoperative nausea or vomiting 0.15 mg/kg 30–45 min before IV catheter placement.\textsuperscript{91}

**Geriatric Dosage.** Same as adult dosage.

**Other Conditions.** In hepatic function impairment, do not exceed a single oral dose of 8 mg or a total daily IV dosage of 8 mg.

**Dosage Forms.** \textbf{Inj} 0.64, 2 mg/mL; \textbf{Tab} 4, 8, 24 mg; \textbf{Tab} (rapidly dissolving) 4, 8 mg; \textbf{Soln} 0.8 mg/mL.

**Pharmacokinetics.** \textbf{Fate}. Oral absorption is 62 ± 15%. $V_d$ is 1.9 ± 0.5 L/kg; $Cl$ is 0.35 ± 0.16 L/hr/kg in adults and can be higher in children. The drug is extensively metabolized to glucuronide and sulfate conjugates. About 5% appears in urine as unchanged ondansetron.\textsuperscript{79} $t_\text{1/2}$. 3.5 ± 1.9 hr in normal adults, increased in the elderly.\textsuperscript{79} However, the duration of activity is not related to the half-life.\textsuperscript{72}

**Adverse Reactions.** Headache occurs frequently. Transient increased serum levels of hepatic enzymes also occur frequently, but these are probably caused by chemotherapy rather than by ondansetron.\textsuperscript{72,92–94}

**Contraindications.** None known.

**Precautions.** Pregnancy; lactation; suspected ileus. Patients who are hypersensitive to other 5-HT\textsubscript{3} antagonists might cross-react with ondansetron.\textsuperscript{95}

**Drug Interactions.** The metabolism of ondansetron can be changed by inducers or inhibitors of the cytochrome P450 system, but dosage adjustment is not recommended.

**Parameters to Monitor.** Frequency of vomiting.

**Notes.** Protect vials from light; inspect for discoloration and particulate matter before using.

Dexamethasone enhances the antiemetic effect of the 5-HT\textsubscript{3} antagonists;\textsuperscript{72,96} the combination of ondansetron and dexamethasone is more effective than metoclopramide and dexamethasone for the acute component but not for the delayed phase of severely emetogenic chemotherapy.\textsuperscript{72} Several studies have documented the lack of additional efficacy beyond that achieved with a total daily ondansetron dosage of 0.45 mg/kg.\textsuperscript{96,97} (See Antiemetic Drugs Comparison Chart.)

**PROCHLORPERAZINE SALTS** Compazine, Various

**Pharmacology.** Prochlorperazine is a phenothiazine tranquilizer with anticholinergic and weak antiemetic activities. It suppresses the chemoreceptor trigger zone in the CNS and is used mainly for its antiemetic properties. It is not effective for the treatment of motion sickness or vertigo.

**Administration and Adult Dosage.** PO as an antiemetic 5–10 mg tid or qid; PR as an antiemetic 25 mg bid; IM as an antiemetic (deep in upper outer quadrant of buttock) 5–10 mg q 4–6 hr, to a maximum of 40 mg/day. IM presurgically (deep in
upper outer quadrant of buttock) 5–10 mg 1–2 hr before induction, can repeat once before or after surgery; IV presurgically 5–10 mg 15–30 min before induction or as infusion (20 mg/L) started 15–30 min before induction. SC not recommended.

**Special Populations. Pediatric Dosage.** Not to be used in surgery or in patients <9 kg or <2 yr. PO or PR as an antiemetic (9–13 kg) 2.5 mg daily–bid; (14–18 kg) 2.5 mg bid–tid; (19–39 kg) 2.5 mg tid–5 mg bid. (See Notes.) IM as an antiemetic (deep in upper outer quadrant of buttock) 0.13 mg/kg. SC not recommended.

**Geriatric Dosage.** Use the lower end of the recommended dosage range in elderly patients.

**Dosage Forms.** Inj 5 mg/mL; Supp 2.5, 5, 25 mg; Syrup 1 mg/mL; Tab 5, 10, 25 mg. Larger-dose tablets are available for psychiatric use. Sustained-release products have no demonstrated advantage over rapid-release products.

**Patient Instructions.** This drug can cause drowsiness. Until the extent of this effect is known, use caution when driving, operating machinery, or performing other tasks requiring mental alertness. Avoid excessive concurrent use of alcohol or other drugs that cause drowsiness.

**Missed Doses.** Take this drug as prescribed. If you miss a dose, take it as soon as you remember. If it is about time for the next dose, take that dose only. Do not double the dose or take extra.

**Pharmacokinetics. Onset and Duration.** PO onset 30–40 min; PR onset 60 min; IM onset 10–20 min. Duration for all routes 3–4 hr.

**Fate.** The drug is well absorbed, but extensive and variable presystemic metabolism in the gut wall and liver limits bioavailability. Eliminated primarily by hepatic metabolism and biliary excretion.

\[ t_{1/2} = 23 \text{ hr.} \]

**Adverse Reactions.** Extrapyramidal reactions, especially dystonias and dyskinesias, occur occasionally in adults and frequently in children (other extrapyramidal reactions are less likely because of the short duration of therapy when used as an antiemetic). Anticholinergic effects such as dry mouth, mydriasis, cycloplegia, urinary retention, decreased GI motility, and tachycardia occur occasionally. SC administration can cause local reactions at injection site.

**Contraindications.** Pediatric surgery; children <9 kg or <2 yr; coma or greatly depressed state caused by CNS depressants.

**Precautions.** Antiemetic action can mask signs and symptoms of overdose with other drugs and can mask the diagnosis and treatment of other conditions such as intestinal obstruction, brain tumor, or Reye’s syndrome. Use with caution in conditions in which the drug’s anticholinergic effects might be detrimental, in children with acute illnesses or dehydration, or in patients with histories of allergy to phenothiazine derivatives (eg, blood dyscrasias, jaundice). Avoid exposure to concentrate on hands or clothing because of the possibility of contact dermatitis.
Drug Interactions. Phenothiazines can decrease the efficacy of guanethidine or guanadrel or have additive hypotensive effects with hypotensive drugs. Phenothiazines can inhibit the antiparkinson activity of levodopa.

Parameters to Monitor. Monitor for extrapyramidal side effects and drug efficacy.

Notes. Protect the solution from light; a slight yellowish discoloration does not indicate altered potency, but markedly discolored solution should be discarded. Protect suppositories from heat. Prochlorperazine does not predictably reduce chemotherapy-induced nausea and vomiting in children and might be associated with an increase in symptoms.99 (See Antiemetic Drugs Comparison Chart.)
### ANTIEMETIC DRUGS COMPARISON CHART

<table>
<thead>
<tr>
<th>DRUG</th>
<th>DOSAGE FORMS</th>
<th>INITIAL DOSE&lt;sup&gt;a,b&lt;/sup&gt;</th>
<th>INDICATIONS</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>Adult</td>
<td>Pediatric</td>
</tr>
<tr>
<td><strong>ANTIHISTAMINES</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Buclizine</td>
<td>Chew Tab 50 mg.</td>
<td>PO 50 mg.</td>
<td>—</td>
</tr>
<tr>
<td>Bucladin-S</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Cyclizine</td>
<td>Tab 50 mg.</td>
<td>PO 50 mg.</td>
<td>PO (6–12 yr) 25 mg.</td>
</tr>
<tr>
<td>Marezine</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Dimenhydrinate</td>
<td>Tab 50 mg.</td>
<td>PO 50–100 mg</td>
<td>PO (2–6 yr) 12.5–25 mg, (6–12 yr) 25–50 mg, (6–12 yr) 25–50 mg, (6–12 yr) 25–50 mg</td>
</tr>
<tr>
<td>Dramamine</td>
<td>Chew Tab 50 mg</td>
<td>IM, IV 50 mg.</td>
<td>IM, IV &gt;2 yr 1.25 mg/kg.</td>
</tr>
<tr>
<td>Various</td>
<td>Liquid 2.5, 3.1 mg/mL, Inj 50 mg/mL</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Diphenhydramine</td>
<td>Cap 25, 50 mg</td>
<td>PO 50 mg.</td>
<td>PO (2–6 yr) 6.25 mg or 1.25 mg/kg, (6–12 yr) 12.5 mg, (6–12 yr) 12.5 mg</td>
</tr>
<tr>
<td>Benadryl</td>
<td>Chew Tab 12.5 mg</td>
<td>IM, IV 10–50 mg.</td>
<td>IM, IV &gt;9 kg 1.25 mg/kg.</td>
</tr>
<tr>
<td>Various</td>
<td>Elixir 2.5 mg/mL, Soln 1.25, 2.5 mg/mL, Syrup 2.5 mg/mL, Inj 50 mg/mL.</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Meclizine</td>
<td>Cap 25, 30 mg</td>
<td>PO 25–50 mg.</td>
<td>—</td>
</tr>
<tr>
<td>Antivert</td>
<td>Tab 12.5, 25, 50 mg</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Bonine</td>
<td>Chew Tab 25 mg.</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Various</td>
<td></td>
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</tr>
</tbody>
</table>

<sup>a</sup> For children, doses should be adjusted based on age and body weight.

<sup>b</sup> Doses may vary depending on the specific formulation and manufacturer's recommendations.
<table>
<thead>
<tr>
<th>DRUG</th>
<th>DOSAGE FORMS</th>
<th>INITIAL DOSE&lt;sup&gt;a,b&lt;/sup&gt;</th>
<th>INDICATIONS</th>
<th>Nausea and Vomiting</th>
<th>Motion Sickness</th>
<th>Vertigo</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>CANNABINOIDS</strong></td>
<td></td>
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</tr>
<tr>
<td><strong>Dronabinol (C-III)&lt;sup&gt;d&lt;/sup&gt;</strong></td>
<td>Cap 2.5, 5, 10 mg.</td>
<td>PO 5 mg/m&lt;sup&gt;2&lt;/sup&gt;.</td>
<td>PO 5 mg/m&lt;sup&gt;2&lt;/sup&gt;.</td>
<td>X</td>
<td></td>
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<tr>
<td><strong>Marinol</strong></td>
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<tr>
<td><strong>PHENOTHIAZINES</strong></td>
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<td></td>
</tr>
<tr>
<td><strong>Chlorpromazine</strong></td>
<td>Tab 10, 25, 50 mg</td>
<td>PO 10–25 mg</td>
<td>PO, IM (&gt;6 months) 0.55 mg/kg</td>
<td>X</td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Thorazine</strong></td>
<td>Liquid 30, 100 mg/mL</td>
<td>PR 50–100 mg</td>
<td>PR (&gt;6 months) 1 mg/kg.</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Various</strong></td>
<td>Syrup 2 mg/mL</td>
<td>IM 25 mg.</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Supp 25, 100 mg</strong></td>
<td>Inj 25 mg/mL.</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Perphenazine</strong></td>
<td>Tab 2, 4, 8, 16 mg</td>
<td>PO 2–4 mg</td>
<td>—</td>
<td>X</td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Trilafon</strong></td>
<td>Liquid 3.2 mg/mL</td>
<td>IM 5 mg.</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Inj 5 mg/mL.</strong></td>
<td></td>
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<td></td>
<td></td>
</tr>
<tr>
<td><strong>Prochlorperazine</strong></td>
<td>Tab 5, 10, mg</td>
<td>PO, IM 5–10 mg</td>
<td>PO, PR (&gt;9 kg or &gt;2 yr) 2.5 mg</td>
<td>X</td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Compazine</strong></td>
<td>Syrup 1 mg/mL</td>
<td>IV 2.5–10 mg</td>
<td>IM (&gt;9 kg or &gt;2 yr) 0.13 mg/kg.</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Various</strong></td>
<td>Supp 2.5, 5, 25 mg</td>
<td>PR 25 mg.</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Inj 5 mg/mL.</strong></td>
<td></td>
<td></td>
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<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Promethazine</strong></td>
<td>Syrup 1.25, 5 mg/mL</td>
<td>PO, PR 25 mg</td>
<td>PO, PR, IM (&gt;2 yr) 0.25–0.5 mg/kg.</td>
<td>X</td>
<td>X</td>
<td></td>
</tr>
<tr>
<td><strong>Phenergan</strong></td>
<td>Tab 12.5, 25, 50 mg</td>
<td>IM, IV 12.5–25 mg.</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Various</strong></td>
<td>Supp 12.5, 25, 50 mg</td>
<td>Inj 25, 50 mg/mL.</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

(continued)
<table>
<thead>
<tr>
<th>DRUG</th>
<th>DOSAGE FORMS</th>
<th>Initial Dosea,b</th>
<th>INDICATIONS</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>Adult</td>
<td>Pediatric</td>
</tr>
<tr>
<td>Thiethylperazine</td>
<td>Tab 10 mg</td>
<td>PO, IM 10 mg.</td>
<td>—</td>
</tr>
<tr>
<td>Torecan</td>
<td>Inj 5 mg/mL.</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Triflupromazine</td>
<td>Inj 10, 20 mg/mL.</td>
<td>IM 5–15 mg</td>
<td>IM (&gt;2.5 yr)</td>
</tr>
<tr>
<td>Vesprin</td>
<td></td>
<td>IM (elderly) 2.5 mg</td>
<td>0.2–0.25 mg/kg.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>IV 1 mg.</td>
<td></td>
</tr>
<tr>
<td>SEROTONIN 5-HT₃ ANTAGONISTS</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Dolasetron</td>
<td>Tab 50, 100 mg</td>
<td>PO 100 mg</td>
<td>PO 1.8 mg/kg</td>
</tr>
<tr>
<td>Anzemet</td>
<td>Inj 20 mg/mL.</td>
<td>IV 1.8 mg/kg.</td>
<td>IV 1.8 mg/kg.</td>
</tr>
<tr>
<td>Granisetron</td>
<td>Tab 1 mg</td>
<td>PO 1 mg</td>
<td>IV (&gt;2 yr) 10 µg/kg.</td>
</tr>
<tr>
<td>Kytril</td>
<td>Inj 1 mg/mL.</td>
<td>IV 10 µg/kg.</td>
<td></td>
</tr>
<tr>
<td>Ondansetron</td>
<td>Tab 4, 8, 24 mg</td>
<td>PO 8 or 16 mg</td>
<td>PO (&gt;4 yr) 4 mg</td>
</tr>
<tr>
<td>Zofran</td>
<td>Tab (rapidly dissolving) 4, 8 mg</td>
<td>IV 0.15–0.45 mg/kg (max 32 mg).</td>
<td>IV (&gt;2 yr) 0.15 mg/kg.</td>
</tr>
<tr>
<td></td>
<td>Soln 6.8 mg/mL.</td>
<td></td>
<td></td>
</tr>
<tr>
<td>MiscellaneouS</td>
<td>Elixir 0.1 mg/mL.</td>
<td>PO, IV 10–20 mg.</td>
<td>PO, IV 10 mg/m².</td>
</tr>
<tr>
<td>Dexamethasone</td>
<td>Soln 0.1, 1 mg/mL.</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Decadron</td>
<td>Tab 0.25, 0.5, 0.75, 1, 1.5, 2, 4.6 mg</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Various</td>
<td>Inj 4, 10, 20, 24 mg/mL.</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

(continued)
# Antiemetic Drugs Comparison Chart (continued)

<table>
<thead>
<tr>
<th>Drug</th>
<th>Dosage Forms</th>
<th>Initial Dose[^a]</th>
<th>Indications</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>Adult</td>
<td>Pediatric</td>
</tr>
<tr>
<td><strong>Droperidol</strong></td>
<td>Inj 2.5 mg/mL, IM, IV 0.625–1.25 mg, IV 0.015–0.075 mg/kg</td>
<td>f</td>
<td></td>
</tr>
<tr>
<td>Inapsine</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Lorazepam (C-IV)</strong></td>
<td>Tab 0.5, 1, 2 mg, PO 1–2 mg, PO 0.05 mg/kg</td>
<td>e</td>
<td></td>
</tr>
<tr>
<td>Ativan</td>
<td>Soln 2 mg/mL, IV 2–4 mg/kg</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Various</td>
<td>Inj 2, 4 mg/mL</td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Methylprednisolone</strong></td>
<td>Inj 40, 125, 500 mg, IV up to 100 mg, IV 2–4 mg/kg</td>
<td>e</td>
<td></td>
</tr>
<tr>
<td>Solu-Medrol</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Various</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Metoclopramide</strong></td>
<td>Inj 5 mg/mL, IV 1–2 mg/kg, IV 1–2 mg/kg</td>
<td>X</td>
<td></td>
</tr>
<tr>
<td>Reglan</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Scopolamine</strong></td>
<td>Transdermal Patch, 1 disk behind</td>
<td>—</td>
<td>X</td>
</tr>
<tr>
<td>Scopace</td>
<td>1.5 mg (delivers ear over 3 days)</td>
<td>—</td>
<td>X</td>
</tr>
<tr>
<td>Transderm Scop</td>
<td>1 mg over 3 days, PO 0.4 mg</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Tab 0.4 mg</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Trimethobenzamide</strong></td>
<td>Cap 100, 250 mg, PO 250 mg, PO (14–40 kg) 100–200 mg</td>
<td>X</td>
<td></td>
</tr>
<tr>
<td>Tigan</td>
<td>Supp 100, 200 mg, PR 200 mg, PR (&lt;14 kg) 100 mg, (14–40 kg) 100–200 mg</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Various</td>
<td>Inj 100 mg/mL, IM 200 mg, 100–200 mg</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

[^a]: Initial dose only; check prescribing information for subsequent dosage.

[^b]: Doses of serotonin antagonists are for chemotherapy-induced nausea and vomiting. (See monograph for doses in postoperative nausea and vomiting.)

[^c]: Possibly effective.

[^d]: Controlled substance schedule designated in parentheses.

[^e]: Not labeled for this use; used as adjunctive for cancer chemotherapy-induced nausea and vomiting.

[^f]: Effective, but not labeled for this use.
BISACODYL

Pharmacology. Bisacodyl is a stimulant cathartic structurally similar to phenolphthalein that produces its effect by direct contact with colonic mucosa. It can inhibit water absorption in the small bowel and colon.\textsuperscript{100,101}

Administration and Adult Dosage. PO as a laxative/cathartic 10–30 mg; PR 10 mg. Adjust dosage based on response.

Special Populations. Pediatric Dosage. PO (≤ 6 yr) safety and efficacy not established; (>6 yr) 5–10 mg or 0.3 mg/kg once daily; PR (≤ 6 yr) safety and efficacy not established; (6–12 yr) 5 mg; (>12 yr) 10 mg.

Geriatric Dosage. Same as adult dosage.

Dosage Forms. EC Tab 5 mg; Enema (adult) 10 mg; Supp 10 mg.

Patient Instructions. Swallow tablets whole (not chewed or crushed) and do not take within 1 hour of antacids or dairy products. Do not use oral products in children 6 years of age or less.

Pharmacokinetics. Onset and Duration. Onset PO 6–12 hr; PR 15 min–1 hr.\textsuperscript{101,102}

Fate. Absorption is less than 5% by oral or rectal route, with subsequent conversion to the glucuronide salt and excretion in urine and bile. Rapidly converted in the gut by intestinal and bacterial enzymes to its active, but nonabsorbed, desacetyl metabolite.\textsuperscript{103}

Adverse Reactions. Abdominal cramps occur frequently; with long-term use, metabolic acidosis or alkalosis, hypocalcemia, tetany, loss of enteric protein, and malabsorption occur occasionally; suppositories can cause proctitis and rectal inflammation and are not recommended for long-term use.

Drug Interactions. Antacids or milk can dissolve the enteric coating of oral bisacodyl tablets, causing drug release in the stomach and gastric irritation.

Contraindications. Acute surgical abdomen; nausea, vomiting, or other symptoms of appendicitis; fecal impaction; intestinal or biliary tract obstruction; abdominal pain of unknown origin.

Notes. Useful for preoperative or preradiographic bowel preparation. Bisacodyl has been used in combination with polyethylene glycol (PEG) electrolyte lavage solution to decrease the amount of solution required.\textsuperscript{100,101,104} (See PEG Electrolyte Lavage Solution.)

DIPHENOXYLATE HYDROCHLORIDE AND ATROPINE SULFATE Lomotil, Various

Pharmacology. Diphenoxylate is a synthetic meperidine congener without analgesic activity that slows GI motility. Because high doses of diphenoxylate (40–60 mg) cause systemic opioid activity, atropine is added in subtherapeutic amounts to decrease abuse potential.

Administration and Adult Dosage. PO for diarrhea 2 tablets or 10 mL qid initially and then, if control is achieved (usually within 48 hr), decrease to a maintenance dosage as low as 2 tablets or 10 mL daily prn. If chronic diarrhea is not
controlled in 10 days at the full dosage, then symptoms are unlikely to be controlled by further administration.

**Special Populations. Pediatric Dosage.** Use liquid only. (<2 yr) Not recommended. PO for diarrhea 0.3–0.4 mg/kg/day of diphenoxylate in 4 divided doses initially, not to exceed adult dosage. Reduce dosage once diarrhea is controlled.

**Geriatric Dosage.** Same as adult dosage.

**Dosage Forms.** Syrup 500 μg diphenoxylate and 5 μg atropine/mL; Tab 2.5 mg diphenoxylate and 25 μg atropine.

**Patient Instructions.** This drug can cause dry mouth, blurred vision, drowsiness, or dizziness; use caution while driving or performing other tasks requiring alertness, coordination, or physical dexterity. Avoid alcohol and other CNS depressants. Seek medical attention if diarrhea persists or if fever, palpitations, or abdominal distention occurs.

**Missed Doses.** If you miss a dose, take it as soon as possible. If it is almost time for your next dose, skip the missed dose and return to your usual dosage schedule. Do not double doses.

**Pharmacokinetics. Onset and Duration.** Onset 45–60 min; duration 3–4 hr.

**Fate.** Diphenoxylate is well absorbed from the GI tract and metabolized to an active metabolite, diphenoxyllic acid. Drug and metabolite attain peak serum levels in 2 hr. Diphenoxylate Vₐ is 3.8 ± 1.1 L/kg; Cl is 1.04 ± 0.14 L/hr/kg. Conjugates of the drug and metabolite are excreted primarily in the urine.

**t₁/₂.** (Diphenoxylate) 2.5 ± 0.6 hr; (diphenoxyllic acid) 7.2 ± 0.7 hr.

**Adverse Reactions.** Anticholinergic symptoms such as dry mouth, urinary retention, blurred vision, fever, or tachycardia occur frequently with high daily dosages and occasionally with usual dosages in children. Drowsiness, dizziness, and headache occur occasionally.

**Contraindications.** Children <2 yr; obstructive jaundice; diarrhea associated with pseudomembranous enterocolitis or enterotoxin-producing bacteria. (See Notes.)

**Precautions.** Use with caution in children because of variable response and potential for toxicity (atropinism) with recommended dosages (particularly in children with Down’s syndrome) and in patients with acute ulcerative colitis, hepatic dysfunction, or cirrhosis. (See Notes.)

**Drug Interactions.** Because of its chemical similarity to meperidine, avoid diphenoxylate use with MAOIs. Use with caution in combination with CNS depressants.

**Parameters to Monitor.** Frequency and volume of bowel movements; body temperature; blood in stool. Watch for signs of atropine toxicity. Monitor for abdominal distention.

**Notes.** In chronic diarrhea, diphenoxylate 5 mg is about equipotent with loperamide 2 mg or codeine 30–45 mg. It might provide temporary symptomatic relief of infectious traveler’s diarrhea (although loperamide is preferred) if used cautiously with an antibiotic, but discontinue if fever occurs, symptoms persist beyond 48 hr, or blood or mucus appears in the stool.
Docusate is an anionic surfactant that lowers the surface tension of the oil–water interface of the stool, allowing fecal material to be penetrated by water and fat, thereby softening the stool. The emulsifying action also enhances the absorption of many fat-soluble drugs and mineral oil. These agents also can cause subtle effects on fluid absorption and secretion in the GI tract.\textsuperscript{100,101}

**Administration and Adult Dosage.** PO as a stool softener (sodium salt) 50–500 mg/day in single or divided doses (give solution/syrup in milk or fruit juice to mask taste); begin therapy with up to 500 mg/day and adjust after maximal effects occur (about 3 days);\textsuperscript{103} (calcium salt) 240 mg/day; (potassium salt) 100–300 mg/day. Use of these agents in elderly, bedridden patients might be ineffective in altering the prevalence of constipation.\textsuperscript{101} PR as enema 50–100 mg in water.

**Special Populations.** Pediatric Dosage. PO (sodium salt) (<3 yr) 10–40 mg/day; (3–6 yr) 20–60 mg/day; (6–12 yr) 40–120 mg/day; (>12 yr) same as adult dosage; give solution/syrup in milk, fruit juice, or formula to mask taste; (calcium salt) (≥6 yr) 50–150 mg/day; (potassium salt) (≥6 yr) 100 mg/day.

**Geriatric Dosage.** Same as adult dosage. (See Notes.)

**Dosage Forms.** (Sodium salt: Colace, various) Cap 50, 100, 240, 250 mg; Soln 10, 50 mg/mL; Syrup 3.3, 4 mg/mL; Tab 100 mg. (Calcium salt: Surfak, various) Cap 50, 240 mg. (Potassium salt: Dialose, various) Cap 240 mg; Tab 100 mg.

**Patient Instructions.** Take this with a full glass of fluid; take the liquid or solution forms in milk, fruit juice, or infant formula to mask the bitter taste.

**Missed Doses.** If you miss a dose, take it as soon as possible. If it is almost time for your next dose, skip the missed dose and return to your usual dosage schedule.

**Pharmacokinetics.** **Onset and Duration.** Onset of effect on stools is 2–3 days after first dose with continuous use.

**Fate.** Drug action is local to the gut, but docusate can be partially absorbed in the duodenum and jejunum and secreted in the bile.\textsuperscript{107}

**Adverse Reactions.** Bitter taste, throat irritation, and nausea (more common with syrup and liquid) occur frequently, abdominal cramps occasionally. Docusate can change intestinal morphology and cellular function and cause fluid and electrolyte accumulation in the colon.\textsuperscript{107}

**Contraindications.** Undiagnosed abdominal pain; intestinal obstruction; concomitant use with mineral oil.

**Precautions.** Rectal bleeding or failure to respond to therapy might indicate a serious condition and the need for medical attention.

**Drug Interactions.** Concomitant use with mineral oil can enhance mineral oil absorption.\textsuperscript{100}

**Parameters to Monitor.** Frequency and consistency of stools; ease of defecation.

**Notes.** Surfactant stool softeners are useful for softening hard, dry stools, in painful anorectal conditions, and in cardiac and other conditions to lessen the strain of defecation. They are more useful in preventing than in treating constipa-
tion but they might not be effective for long-term prevention of constipation in institutionalized, elderly patients.100,101,110

**LACTULOSE**

**Pharmacology.** Lactulose is a synthetic disaccharide analogue of lactose that contains galactose and fructose and is metabolized by colonic bacteria to lactic and small amounts of acetic and formic acids. These acids result in acidification of colonic contents, decreased ammonia production and absorption, and an osmotic catharsis.111

**Administration and Adult Dosage.** PO as a cathartic 15–30 mL (10–20 g), to a maximum of 60 mL; PO for hepatic encephalopathy 30–45 mL (20–30 g) q 1 hr until laxation, then 30–45 mL tid or qid, titrated to produce about 2 or 3 soft stools/day. PR for hepatic encephalopathy as an enema 300 mL with 700 mL water or NS retained for 30–60 min, can repeat q 4–6 hr. Repeat immediately if evacuated too promptly.

**Special Populations.** Pediatric Dosage. PO for hepatic encephalopathy (infants) 2.5–10 mL/day in divided doses; (older children and adolescents) 40–90 mL/day in divided doses, titrated to produce 2 or 3 soft stools daily. If initial dose causes diarrhea, reduce dose immediately; if diarrhea persists, discontinue.

**Geriatric Dosage.** Same as adult dosage. (See Notes.)

**Dosage Forms.** Syrup 667 mg/mL.

**Patient Instructions.** This syrup can be mixed with fruit juice, water, or milk to improve its palatability. In the treatment of hepatic encephalopathy, 2–3 loose stools per day are common, but report any worsening of diarrhea. Report belching, flatulence, or abdominal cramps if they are bothersome.

**Missed Doses.** If you miss a dose, take it as soon as possible. If it is almost time for your next dose, skip the missed dose and return to your usual dosage schedule.

**Pharmacokinetics.** Onset and Duration. (Catharsis) onset 24–48 hr; duration 24–48 hr. (Hepatic encephalopathy) onset and duration variable; however, reversal of coma can occur within 2 hr of the first enema.

**Fate.** After oral administration, less than 3% is absorbed and most reaches the colon unabsorbed and unchanged. Unabsorbed drug is metabolized in the colon by bacteria to low-molecular-weight acids and carbon dioxide. The small amount of absorbed drug is excreted in the urine unchanged.112

**Adverse Reactions.** Flatulence, belching, and abdominal discomfort are frequent initially. Colonic dilation occurs occasionally. Excessive diarrhea and fecal water loss can result in hypernatremia.113

**Contraindications.** Patients who require a low-galactose diet.

**Precautions.** Use with caution in diabetics because the drug contains small amounts of free lactose and galactose. Rectal bleeding or failure to respond to therapy might indicate a serious condition and the need for medical attention.

**Drug Interactions.** Do not use other laxatives concomitantly because their induction of loose stools might confound proper lactulose dosage titration for hepatic...
encephalopathy. Nonabsorbable antacids can interfere with the colonic acidification of lactulose. Theoretically, some antibacterials might interfere with the intestinal bacteria that metabolize lactulose; however, oral neomycin has been used concurrently in hepatic encephalopathy.  

**Parameters to Monitor.** (Hepatic encephalopathy) observe for changes in hepatic encephalopathy and number of stools per day. Periodically obtain serum sodium, chloride, potassium, and bicarbonate levels during prolonged use, especially in elderly or debilitated patients.

**Notes.** Lactulose is effective in hepatic encephalopathy, but as a general laxative it offers no advantage over less expensive drugs. One study of constipation in the elderly found that up to 60 mL/day of 70% sorbitol was equivalent in laxative effects and caused less nausea than the same dosage of lactulose syrup.

**Pharmacology.** Loperamide is a synthetic antidiarrheal structurally similar to haloperidol and without appreciable opiate activity that causes a dose-related inhibition of colonic motility and affects water and electrolyte movement through the bowel. Tolerance has not been observed.

**Administration and Adult Dosage.** PO for acute diarrhea (B) or traveler’s diarrhea (over-the-counter [OTC]) 4 mg initially and then 2 mg after each unformed stool (often with an antibiotic for traveler’s diarrhea), to a maximum of 16 mg/day (8 mg/day for no more than 2 days with OTC product). PO for chronic diarrhea (B) initiate therapy as above and then individualize dosage; usual maintenance dosage is 4–8 mg/day in single or divided doses. If clinical improvement does not occur after treatment with 16 mg/day for at least 10 days, symptoms are unlikely to be controlled by further use.

**Special Populations. Pediatric Dosage.** (<2 yr) safety and efficacy not established. PO for acute diarrhea (B) (2–5 yr) up to 1 mg tid as liquid; (6–8 yr) 2 mg bid; (8–12 yr) 2 mg tid. After the first day of therapy, give 1 mg/10 kg after each loose stool, to a maximum daily dosage equal to the initial daily dosage. PO for acute or traveler’s diarrhea (OTC) (2–5 yr) not recommended; (6–8 yr) 1 mg initially and then 1 mg after each loose stool, to a maximum of 4 mg/day for 2 days; (9–11 yr) 2 mg initially and then 1 mg after each loose stool, to a maximum of 6 mg/day for 2 days. PO for chronic diarrhea dosage not established.

**Geriatric Dosage.** Same as adult dosage.

**Dosage Forms.** Cap 2 mg; Chew Tab 2 mg; Tab 2 mg; Liquid 0.2, 1 mg/mL; Chew Tab 2 mg plus simethicone (Imodium Advanced).

**Patient Instructions.** This drug can cause drowsiness or dizziness. Until the severity of these reactions is known, use caution when performing tasks that require mental alertness. It can cause dry mouth. Drink plenty of clear fluids to prevent the dehydration that can accompany diarrhea. If diarrhea does not stop after a few days, or if abdominal pain, distention, or fever occurs, seek medical attention.

**Missed Doses.** If you miss a dose, take it as soon as possible. If it is almost time for your next dose, skip the missed dose and return to your usual dosage schedule.
Pharmacokinetics. Onset and Duration. Onset 45–60 min; duration 4–6 hr.

Fate. GI absorption is approximately 40%; ≥25% is excreted in the stool unchanged; <2% of a dose is recovered in the urine. \( t_{1/2} \) 10.8 ± 1.7 hr.

Adverse Reactions. Abdominal cramping, constipation, distention, headache, rash, tiredness, drowsiness, dizziness, and dry mouth occur frequently.

Contraindications. (Rx, OTC) patients who must avoid constipation; children <2 yr. (OTC) bloody diarrhea; body temperature above 38°C (101°F); diarrhea associated with pseudomembranous colitis; or enterotoxin-producing bacteria. (See Notes.)

Precautions. Use with caution in patients with ulcerative colitis. Discontinue if improvement is not observed within 48 hr. Use cautiously in patients with hepatic dysfunction.

Drug Interactions. Absorption of loperamide can be decreased by cholesterol-binding resins.


Notes. Adverse reactions might be less frequent and efficacy might be greater than with diphenoxylate with atropine. Loperamide can provide temporary symptomatic relief of infectious traveler’s diarrhea if used cautiously with an antibiotic, but discontinue if fever occurs or other symptoms persist beyond 48 hr, or blood or mucus in stool develops.

Pharmacology. Magnesium salts act as saline cathartics that inhibit fluid and electrolyte absorption by increasing osmotic forces in the gut lumen. Part of the action might be caused by cholecystokinin release, which stimulates small bowel motility and inhibits fluid and electrolyte absorption from the small intestine.

Administration and Adult Dosage. PO as a laxative/cathartic (citrate) 240 mL; (sulfate) 20–30 mL of 50% solution (10–15 g) in a full glass of water; (hydroxide; milk of magnesia) 30–60 mL with liquid; (concentrate) 10–30 mL. (See Notes.)

Special Populations. Pediatric Dosage. PO (citrate) one-half the adult dosage; (sulfate) (2–5 yr) 2.5–5 g, (≥6 yr) 5–10 g in one-half glass or more of water; (hydroxide; milk of magnesia) 0.5 mL/kg.

Geriatric Dosage. Same as adult dosage.

Other Conditions. Avoid use in patients with impaired renal function.

Dosage Forms. Soln (citrate) 77 mEq/dL magnesium, 300 mL; (sulfate); Susp (hydroxide; milk of magnesia) 7–8.5%, many sizes, (see Notes) (also available as concentrates with 10 mL equivalent to 20 or 30 mL of susp); Tab (hydroxide; milk of magnesia) 311 mg; Pwdr (sulfate) 150, 240, 454 g.
Patient Instructions. Take milk of magnesia or magnesium sulfate with at least one full glass of liquid. You can take magnesium sulfate with fruit juice to partly mask its bitter taste. Refrigerating magnesium citrate improves its palatability.

Pharmacokinetics. Onset and Duration. Onset is dose dependent: (high end of dosage range) 0.5–3 hr; (low end of dosage range) 6–8 hr.\textsuperscript{103}

Fate. Slow absorption of about 10\% of a dose from the GI tract. Absorbed magnesium is rapidly excreted in the urine in normal renal function.\textsuperscript{100,101}

Adverse Reactions. Abdominal cramping, excessive diuresis, nausea, vomiting, and diarrhea occur frequently. Excessive use can lead to electrolyte abnormalities; dehydration can occur if taken with insufficient fluids. Use in patients with renal impairment can lead to hypermagnesemia, CNS depression, and hypotension.\textsuperscript{100,101,107,118}

Contraindications. Acute surgical abdomen; fecal impaction; intestinal obstruction; abdominal pain of unknown origin; nausea; vomiting.

Precautions. Rectal bleeding or failure to respond to therapy might indicate a serious condition and the need for medical attention. Avoid use in patients with impaired renal function.\textsuperscript{100,101,107,118}

Drug Interactions. None known.

Parameters to Monitor. Periodically check serum magnesium levels in patients with impaired renal function who are receiving long-term daily administration.

Notes. Magnesium salts are useful for preparing the bowel for radiologic examination and surgical procedures. One regimen used magnesium citrate solution 300 mL 2 hr before PEG electrolyte lavage solution that was continued until the stool return was clear.\textsuperscript{119} The following amounts of various magnesium salts are approximately equivalent to 80 mEq of magnesium: 100 mL citrate; 2.4 g (30 mL) milk of magnesia; and 10 g sulfate. The sulfate salt is the most potent but the least palatable cathartic. (See also Magnesium Salts in the Renal and Electrolyte sections.)

METOCLOPRAMIDE

Pharmacology. Metoclopramide stimulates the release of acetylcholine from the gastric myenteric plexus by antagonizing peripheral and central dopamine (D) receptors, specifically the D\textsubscript{2} subtype receptors. Metoclopramide also acts as a partial agonist at the 5-HT\textsubscript{4} receptors, thereby facilitating the release of acetylcholine in the GI tract; however, it acts as an antagonist at the 5-HT\textsubscript{3} receptor site.\textsuperscript{120} It increases peristalsis of the gastric antrum, duodenum, and jejunum, relaxes the pyloric sphincter and duodenal bulb, and has little effect on the colon or gallbladder. In patients with GERD, metoclopramide produces a dose-dependent increase and duration of action in lower esophageal sphincter pressure. Its antiemetic action results from a direct antidopaminergic effect on the chemoreceptor trigger zone and vomiting center and from 5-HT\textsubscript{3}–receptor blocking effects. Metoclopramide increases prolactin secretion and serum prolactin. It also produces a transient increase in aldosterone secretion, thought to be related to direct stimulation of the adrenal gland via stimulation of the 5-HT\textsubscript{3} receptor.\textsuperscript{121}
Administration and Adult Dosage. PO for short-term treatment of symptomatic GERD in patients who fail to respond to conventional therapy up to 15 mg qid 30 min before each meal and hs for 4–12 weeks or intermittent single doses of up to 20 mg; PO for symptomatic diabetic gastroparesis 10 mg qid 30 min before each meal and hs for 2–8 weeks; IM or IV for severe symptoms associated with gastroparesis 10 mg qid for up to 10 days; IV to facilitate small bowel intubation or to aid in radiologic examination 10 mg over 1–2 min, 10-30 min before tube placement.\textsuperscript{122,123} PO to increase maternal milk supply 10 mg tid for 10–14 days.\textsuperscript{124} PO, IM, or IV for the treatment of hiccups, PO 10 mg q 6 hr for 10 days, or IM, IV 5–10 mg q 8 hr for 24–48 hr and then switch to PO.\textsuperscript{125} IV for prevention of chemotherapy-induced emesis 2 mg/kg q 2–4 hr for 2–5 doses; IV for delayed nausea and vomiting 0.5 mg/kg or 30 mg IV q 4–6 hr for 3–5 days. PO 2 mg/kg q 2–4 hr for 2–5 doses; PO for delayed nausea and vomiting 0.5 mg/kg or 30 mg PO q 4–6 hr for 3–5 days.\textsuperscript{72} IM for prevention of postoperative nausea and vomiting 10–20 mg near the end of surgery. IV for treatment of postoperative nausea and vomiting 10 mg q 4–6 hr pn postoperatively.\textsuperscript{72} Administer undiluted IV metoclopramide slowly (at least 1–2 min for a 10-mg dose); infuse diluted IV doses over at least 15 min (See Notes.)

Special Populations. Pediatric Dosage. IV to facilitate small bowel intubation or aid radiologic examination (<6 yr) 0.1 mg/kg; (6–14 yr) 2.5–5 mg; (>14 yr) same as adult dosage. IV for postoperative nausea and vomiting 0.1–0.2 mg/kg.\textsuperscript{72}

Geriatric Dosage. Begin at one-half the initial dose (usually 5 mg) and increase or decrease based on efficacy and side effects.

Other Conditions. With Clcr <40 mL/min, begin at one-half the initial dose (usually 5 mg) and increase or decrease based on efficacy and side effects.

Dosage Forms. Tab 5, 10 mg; Soln 10 mg/mL; Syrup 1 mg/mL; Inj 5 mg/mL.

Patient Instructions. Take each dose 30 minutes before meals and at bedtime. This drug can cause drowsiness. Until the degree of drowsiness is known, use caution when driving, operating machinery, or performing other tasks requiring mental alertness. Avoid excessive concurrent use of alcohol or other drugs that cause drowsiness. Report any involuntary movements (eg, muscle spasms and jerky movements of the head and face) that occur, especially in children and the elderly.

Missed Doses. If you miss a dose, take it as soon as possible. If it is almost time for your next dose, skip the missed dose and return to your usual dosage schedule. Do not double doses.

Pharmacokinetics. Onset and Duration. (GI effects) PO onset 45 ± 15 min, IM 12.5 ± 2.5 min, IV 2 ± 1 min; duration 1–2 hr.

Fate. Bioavailabilities are PO 80 ± 15.5% and IM 85 ± 11%. Peak serum concentration after a PO dose occurs in 1–2 hr but can be delayed with impaired gastric emptying. The drug is about 30% bound to plasma proteins. $V_d$ is 3.4 ± 1.3 L/kg, increased in uremia and in cirrhosis; $Cl$ is 0.37 ± 0.08 L/hr/kg, decreased in uremia and in cirrhosis. About 85% of orally administered drug is recovered in the urine after 72 hr as unchanged drug; 20% of an IV dose is excreted unchanged in urine.\textsuperscript{79}
Adverse Reactions. Most side effects are related to dosage and duration of use. Drowsiness, restlessness, fatigue, and lassitude occur in 10% of patients with a dosage of 10 mg qid and in 70% with IV doses of 1–2 mg/kg. Acute dystonic reactions occur in 0.2% of patients receiving 30–40 mg/day, 2% in cancer chemotherapy-treated patients >35 yr receiving doses of 1–2 mg/kg, and 25% in cancer chemotherapy-treated children without prior diphenhydramine treatment. Parkinsonian symptoms, tardive dyskinesia, and akathisia occur less frequently. Rapid IV push produces transient, intense anxiety, and restlessness followed by drowsiness. Transient flushing of the face and/or diarrhea occur frequently after large IV doses. Hyperprolactinemia can occur, resulting in gynecomastia and impotence in males and galactorrhea and amenorrhea in females. Fluid retention can result from transient elevation of aldosterone secretion that occurs after parenteral, but not oral, administration. Diarrhea, hypertension, and mental depression have been reported. Neuroleptic malignant syndrome is a rare, but potentially fatal, adverse effect reported to occur with metoclopramide.

Contraindications. GI hemorrhage; mechanical obstruction or perforation; pheochromocytoma; epilepsy; concurrent use of drugs that cause extrapyramidal effects.

Precautions. Pregnancy; lactation. Use with caution in the elderly and in patients with hypertension, renal failure, or Parkinson’s disease, history of depression or attempted suicide, and after gut anastomosis. In patients with diabetic gastroparesis, insulin dosage or timing might require adjustment.

Drug Interactions. Absorption of drugs from the stomach or small bowel can be altered by metoclopramide (eg, digoxin and cimetidine absorption is decreased; cyclosporine absorption is increased). Anticholinergics and narcotics may antagonize GI effects of metoclopramide. Use with an MAOI can result in hypertension, and the combination should be avoided. Additive sedation can occur with alcohol or other CNS depressants.

Parameters to Monitor. Monitor periodically for CNS effects, extrapyramidal reactions, and changes in Crs, blood glucose, or blood pressure. (GERD or diabetic gastroparesis) observe for symptomatic relief.

Notes. Tolerance to the drug’s gastrokinetic effect can develop with long-term therapy. Metoclopramide has been used in the treatment of neurogenic bladder, orthostatic hypotension, Tourette’s syndrome, adynamic or chemotherapy-induced ileus, anorexia, and complications of scleroderma. If extrapyramidal symptoms occur, administer diphenhydramine 50 mg IM or benztropine 1–2 mg IM.

Cisapride (Propulsid—Janssen Pharmaceutica) was available for the symptomatic treatment of adults with nighttime heartburn due to GERD. Cisapride is no longer marketed in the United States, but will be available only through an Investigational Limited-Access Program because of serious cardiovascular effects (eg, prolonged QT interval, torsades de pointes) in patients taking interacting medications or with certain underlying health conditions. For patients to be considered for the Propulsid Investigational Limited-Access Program, they must have
failed all standard therapies and have baseline laboratory tests and ECG, and undergone an appropriate diagnostic evaluation including radiologic examinations or endoscopy. Contact Janssen Pharmaceutica at 1-800-JANSSEN to determine whether a patient qualifies for the program.

**Domperidone** (Motilium—Janssen) is a prokinetic agent available outside the U.S. for the treatment of diabetic gastroparesis. It selectively blocks peripheral dopamine-D₂ receptors in the GI tract; it has antiemetic effects related to its action at the chemoreceptor trigger zone; and it stimulates pituitary prolactin release in humans but has no cholinergic activity. The drug does not cross the blood–brain barrier and thus does not produce CNS and extrapyramidal effects. Domperidone improves delayed gastric emptying and enhances antral and duodenal peristalsis but does not affect esophageal or colonic motility. PPIs, H₂-receptor antagonists, and antacids should not be coadministered with domperidone because the drug requires an acidic environment for activity. Dosages of 10–20 mg tid have been studied for dyspepsia and 20 mg qid is being studied for the treatment of diabetic gastroparesis. The most frequent side effects of domperidone are headache, dry mouth, anxiety, and elevation in serum prolactin concentrations.⁵²⁸

**Erythromycin** is a macrolide antibiotic that has prokinetic activity by acting as a motilin receptor agonist in the GI tract to stimulate GI contractility.¹²⁹ In gastroparesis, doses of 200–250 mg IV given over 15–30 min of the lactobionate salt, 250 mg PO tid of the ethylsuccinate salt, or 500 mg PO of the stearate salt 15–120 min before meals and at hs appears to be effective.¹²⁹,¹³⁰ Erythromycin ethylsuccinate suspension formulation has a faster prokinetic action than erythromycin stearate tablets.¹³⁰

**PEG ELECTROLYTE LAVAGE SOLUTION**

**Pharmacology.** Polyethylene glycol (PEG) electrolyte lavage solution is an isosmotic solution containing approximately 5.69 g/L sodium sulfate, 1.68 g/L sodium bicarbonate, 1.46 g/L sodium chloride, 745 mg/L potassium chloride, and 60 g/L PEG 3350; it is used for total bowel cleansing before GI examination. A solution lacking sodium sulfate, with a slight variation in other salts and PEG (NULYTELY), and flavored solutions are available with improved palatability. PEG acts as an osmotic cathartic, and the electrolyte concentrations are such that there is little net fluid or electrolyte movement into or out of the bowel.¹⁰¹,¹⁰⁴,¹³¹

**Adult Dosage.** PO or NG 200–300 mL orally q 10 min or by NG tube at a rate of 20–30 mL/min until about 4 L are consumed or the rectal effluent is clear. Use a 1-L trial before the full dosage in patients suspected of having bowel obstruction. Use the solution at least 4 hr before the examination, allowing the patient 3 hr for drinking and a 1-hr period to complete bowel evacuation. Another method is to give the solution the evening before the examination. Chilling the solution might improve its palatability but do not add other ingredients. Withhold solid food for 2 hr and medication for 1 hr before the solution is administered.

**Pediatric Dosage.** PO or NG 25–40 mL/kg/hr for 4–10 hr appear safe and useful for bowel evacuation.

**Dosage Forms.** Available as powder for reconstitution and oral solution.
Pharmacokinetics. The first bowel movement usually occurs after 1 hr, with total bowel cleansing 3–4 hr after starting.

Adverse Reactions. Frequent side effects are nausea, abdominal fullness, bloating (in up to 50% of patients), cramps, anal irritation, and vomiting. Urticaria, rhinorrhea, and dermatitis occur occasionally. Do not use PEG electrolyte lavage solution in patients with GI obstruction, gastric retention, toxic colitis, toxic megacolon, ileus, or bowel perforation; the solution seems to be safe for patients with liver, kidney, or heart disease.

Notes. This product is well suited for bowel cleansing before colonoscopy, but, because of some residual lavage fluid retained in the colon, other cleansing methods might be preferred before barium enema. Colonic cleaning with bisacodyl 15 mg orally followed by 2 L of PEG lavage solution 8 hr later has been found to be equally effective and more acceptable to patients than 4 L of solution used alone. Similar results were obtained using 300 mL of magnesium citrate solution 2 hr before PEG lavage solution that was continued until stool return was clear. The drug might be useful as a GI evacuant in ingestions and overdoses with iron and some EC and SR drug products.

Psyllium is a bulk-forming cathartic that absorbs water and provides an emollient mass.

Administration and Adult Dosage. PO for constipation 2.5–12 g daily–tid, stirred in a full glass of fluid, followed by an additional glass of liquid. PO for mild diarrhea usual doses titrated to effect can be used to “firm up” effluent. PO to lower cholesterol 10–30 g/day in divided doses in combination with diet can decrease cholesterol in patients with mild to moderate hypercholesterolemia.

Special Populations. Pediatric Dosage. PO for constipation (<6 yr) safety and efficacy not established; (6–12 yr) 2.5–3 g (psyllium) daily–tid, with fluid as above.

Geriatric Dosage. Same as adult dosage.

Dosage Forms. Pwdr (sugar-free) Konsyl (containing 100% psyllium) 6 g packet, 200–660 g; Metamucil, Sugar-Free Orange Flavor (containing 65% psyllium); Pwdr (containing sugar) Metamucil Orange Flavor (50% or 65% sucrose), Konsyl-Orange (28% psyllium, 72% sucrose) 7, 11, 12 g packet, 210, 420, 538, 630, 960 g; Wafer Metamucil (containing 5 g fat) 3.4 g of psyllium per wafer.

Patient Instructions. Mix powder with a full glass of fluid before taking and follow with another glass of liquid.

Pharmacokinetics. Onset and Duration. Onset 12–24 hr, but 2–3 days might be required for full effect.

Fate. Not absorbed from GI tract; eliminated unchanged in feces.

Adverse Reactions. Flatulence occurs frequently. Serious side effects are rare, but esophageal, gastric, intestinal, and rectal obstructions have been reported. Allergic reactions and bronchospasm have occurred after inhalation of dry powder.
Contraindications. Acute surgical abdomen; fecal impaction; intestinal obstruction; abdominal pain of unknown origin; nausea; vomiting.

Precautions. Rectal bleeding or failure to respond to therapy might indicate a serious condition and the need for medical attention. Use with caution in patients who require fluid restriction because constipation can occur unless fluid intake is adequate. Psyllium can be hazardous in patients with intestinal ulcerations, stenosis, or disabling adhesions. Use effervescent Metamucil formulations (packet) with caution in patients who require potassium restriction (7.4 and 7.9 mEq potassium/packet). Use the noneffervescent formulations of Metamucil cautiously in diabetics because they contain 50% or 65% sucrose. Sugar-free preparations include Konsyl and Metamucil Sugar Free.

Drug Interactions. None known.

Notes. Psyllium is useful in lessening the strain of defecation and for inpatients who are on low-residue diets or constipating medications. It is safe to use during pregnancy.\(^{100,101}\)

### Miscellaneous Gastrointestinal Drugs

**ACTIVATED CHARCOAL** Various

**Pharmacology.** Activated charcoal is a nonspecific GI adsorbent with a surface area of 900–2000 m\(^2\)/g that is used primarily in the management of acute poisonings.\(^{136}\)

**Administration and Adult Dosage.** PO or via gastric tube 50–120 g dispersed in liquid as soon as possible after ingestion of poison (the Food and Drug Administration suggests 240 mL diluent/30 g activated charcoal). Repeat administration of activated charcoal after gastric lavage. (See Notes.)

**Special Populations.** Pediatric Dosage. PO or via gastric tube (≤12 yr) 25–50 g or 1–2 g/kg dispersed in liquid; (>12 yr) same as adult dosage.\(^{137}\)

**Geriatric Dosage.** Same as adult dosage.

**Dosage Forms.** Pwdr plain or dispersed in water or sorbitol-water.

**Patient Instructions.** This drug causes stools to turn black.

**Pharmacokinetics.** *Onset and Duration.* Onset is immediate; duration is continual while it remains in the GI tract.

**Fate.** Not orally absorbed; eliminated unchanged in the feces.

**Adverse Reactions.** Black stools; gritty consistency can cause emesis in some patients.

**Precautions.** Insufficient hydration or use in patients with decreased bowel motility can result in intestinal bezoars.

**Drug Interactions.** Activated charcoal can decrease the oral absorption and efficacy of many drugs. (See Notes.)

**Parameters to Monitor.** Passage of activated charcoal in the stools. If sorbitol or other cathartics are administered, limit their dosages to prevent excessive fluid and electrolyte losses.
Notes. A suspension of activated charcoal in 25–35% sorbitol can increase palatability of the drug; total dosage of sorbitol should not exceed 1 g/kg. Substances not adsorbed by activated charcoal are mineral acids, alkalis, iron, cyanide, lithium and other small ions, and alcohols. Repeated oral doses of activated charcoal (eg, 15–30 g q 4–6 hr) have been used to enhance the elimination of some drugs, most notably carbamazepine, phenobarbital, salicylates, and theophylline.

Anti-Irritable Bowel Syndrome Agents

Alosetron (Lotronex) is a selective serotonin 5-HT3 antagonist that was removed from the market after numerous reports of ischemic colitis and several deaths.138–140 Several other drugs are being studied for use in treating irritable syndrome. Tegaserod (Zelnorm—Novartis) is a 5-HT3-receptor partial agonist that appears to decrease abdominal pain and bloating and increase the frequency of bowel movements in patients with constipation-predominant irritable bowel syndrome. It also appears to be effective in alternating irritable bowel syndrome. The most effective dose is 6 mg bid, and the most common adverse effect is diarrhea with initial therapy, which eventually dissipates with continued treatment.141 Prucalopride (Rezolor—Janssen) is being evaluated for patients with delayed small bowel and colonic motility. Patients with chronic constipation might benefit from this drug, which is a benzofurancarboxamide selective 5-HT4-receptor agonist. In healthy subjects, prucalopride stimulates colonic activity; however, it has minimal effects on gastric and small bowel transit times.142 Diarrhea, abdominal pain, headache, flatulence, and nausea are its most common side effects.142,143 Cilansetron (Solvay) is a 5-HT3-receptor antagonist similar to alosetron that is being evaluated for diarrhea-predominant irritable bowel syndrome. Cilansetron might have pharmacologic effects similar to those of alosetron.144

Mesalamine Preparations

Pharmacology. Mesalamine (5-aminosalicylic acid [5-ASA]) is thought to be the active moiety of sulfasalazine. The mechanism of action of mesalamine in inflammatory bowel disease is unknown, but mesalamine seems to inhibit cyclooxygenase and 5-lipoxygenase, thereby downregulating the production of inflammatory prostaglandins in the colon. An immunomodulatory response also might occur because mesalamine inhibits and prevents the secretion of antibodies and lymphocytes during active disease. Mesalamine inhibits macrophage and neutrophil chemotaxis, reduces intestinal mononuclear cell production of immunoglobulin A and G antibodies, and is a scavenger of oxygen-derived free radicals, which are increased during active inflammatory bowel disease.146–150 Balsalazide disodium is a prodrug that is cleaved by bacterial azoreductase in the colon to release mesalamine and the inactive carrier, 4-aminobenzoyl-β-alanine.145 Balsalazide 750 mg is equivalent to 267 mg of mesalamine. Each molecule of olsalazine that reaches the colon is converted to 2 molecules of mesalamine.
<table>
<thead>
<tr>
<th>INDICATION</th>
<th>ASACOL</th>
<th>COLAZIDE</th>
<th>DIPENTUM</th>
<th>PENTASA</th>
<th>AXCAN, ROWASA</th>
</tr>
</thead>
<tbody>
<tr>
<td>Short-term treatment of active mild to moderate ulcerative colitis.</td>
<td>PO 800 mg tid, or 1.6 g tid&lt;sup&gt;a&lt;/sup&gt; for 6 weeks.</td>
<td>PO 2.25 g tid.</td>
<td>PO 500 mg tid&lt;sup&gt;a&lt;/sup&gt; or 1 g tid&lt;sup&gt;b&lt;/sup&gt; for 3–6 weeks.</td>
<td>PO 1 g qid for 6–8 weeks.</td>
<td>PR 2 g hs,&lt;sup&gt;a,b&lt;/sup&gt; or 4 g hs&lt;sup&gt;a&lt;/sup&gt; for 3–6 weeks (enema).</td>
</tr>
<tr>
<td>Maintenance of ulcerative colitis remission.</td>
<td>PO 800 mg bid.</td>
<td>a</td>
<td>PO 500 mg bid.&lt;sup&gt;c&lt;/sup&gt;</td>
<td>PO 1 g bid&lt;sup&gt;a&lt;/sup&gt; or 1 g qid&lt;sup&gt;a&lt;/sup&gt;</td>
<td>PR 1–2 g hs&lt;sup&gt;a,b&lt;/sup&gt; (enema).</td>
</tr>
<tr>
<td>Short-term treatment of active mild to moderate Crohn’s disease.</td>
<td>PO 800 mg tid&lt;sup&gt;d&lt;/sup&gt; or 1.6 g tid&lt;sup&gt;d&lt;/sup&gt; for 8–16 weeks.</td>
<td>a</td>
<td>a</td>
<td>PO 1 g qid&lt;sup&gt;a&lt;/sup&gt;</td>
<td>a</td>
</tr>
<tr>
<td>Maintenance of Crohn’s disease remission.</td>
<td>PO 800 mg–1.6 g tid.&lt;sup&gt;a&lt;/sup&gt;</td>
<td>a</td>
<td>a</td>
<td>PO 1 g bid&lt;sup&gt;a&lt;/sup&gt; or 1 g qid&lt;sup&gt;a&lt;/sup&gt;</td>
<td>a</td>
</tr>
<tr>
<td>Treatment of active proctitis.</td>
<td>PO 800 mg tid.&lt;sup&gt;a&lt;/sup&gt;</td>
<td>a</td>
<td>a</td>
<td>PO 1 g qid&lt;sup&gt;a&lt;/sup&gt;</td>
<td>PR 1–2 g hs.&lt;sup&gt;a,b&lt;/sup&gt; 4 g hs&lt;sup&gt;a&lt;/sup&gt; (enema); 500 mg bid or tid&lt;sup&gt;a&lt;/sup&gt; (Axcan, Rowasa, suppository).</td>
</tr>
</tbody>
</table>

<sup>a</sup>Nonlabeled indication and dosage; optimal dosage regimen has not been determined.

<sup>b</sup>Retain enema for approximately 8 hr.

<sup>c</sup>Patients intolerant to sulfasalazine.

<sup>d</sup>Retain suppository for 1–3 hr or longer.

*From references 146–149.*
Special Populations. Pediatric Dosage. Safety and efficacy not established. Pediatric use has been reported. PO mesalamine (Asacol, Pentasa) 30–50 mg/kg/day in 3–4 divided doses (maximum dosages: Asacol 4.8 g/day, Pentasa 4 g/day); PR mesalamine enema 1–4 g q hs; PR mesalamine suppository 500 mg q hs or bid.172,173

Geriatric Dosage. No dosage reduction is necessary. Older patients are more likely to have renal impairment. (See Precautions.)

Other Conditions. Dosage reduction might be considered in severe renal and/or hepatic impairment.146 (See Precautions.)

Dosage Forms.
## Table: Pharmacology of Mesalamine Prodrugs

<table>
<thead>
<tr>
<th>DRUG</th>
<th>ASACOL</th>
<th>COLAZIDE</th>
<th>DIPENTUM</th>
<th>PENTASA</th>
<th>AXCAN, CANASA, ROWASA</th>
</tr>
</thead>
<tbody>
<tr>
<td>Site of action</td>
<td>Distal ileum to colon.</td>
<td>Colon.</td>
<td>Colon.</td>
<td>Proximal jejunum to colon.</td>
<td>Rectum or splenic flexure (enema); rectum (suppository).</td>
</tr>
<tr>
<td>Dosage forms</td>
<td>EC Tab 400 mg.</td>
<td>Cap 750 mg.</td>
<td>Cap 250 mg.</td>
<td>SR Cap 250 mg.</td>
<td>Enema 4 g/60 mL; (Rowasa) Supp 500 mg. (Axcan, Canasa, Rowasa)</td>
</tr>
</tbody>
</table>

5-ASA = 5-aminosalicylic acid.

*Each molecule of olsalazine that reaches the colon is converted to 2 molecules of mesalamine.*
**Patient Instructions.** (Oral) take mesalamine with food and a full glass of water. Swallow tablets or capsules whole without breaking or chewing. The tablet core (Asacol) or small beads (Pentasa) might appear in the stool after mesalamine is released, but this does not mean there was a lack of effect. Report intact or partly intact tablets in the stool (Asacol) because this might indicate that the expected amount of mesalamine was not released from the tablet. Report nausea, vomiting, abrupt change in character or volume of stools, or skin rashes. (Rectal) empty bowel immediately before insertion of enema or suppository. Use enema at bedtime and retain for 8 hours, if possible. Retain suppository for at least 1 to 3 hours. Report signs of anal or rectal irritation. Rectal formulations can stain materials that come into direct contact with them.

**Missed Doses.** (Oral) if you miss a dose, take it as soon as possible. If it is almost time for your next dose, skip the missed dose and return to your usual dosage schedule. Do not double doses. (Rectal) if you miss a dose, use it as soon as possible if you remember it that same night. If you do not remember it until the next morning, skip the missed dose and return to your usual dosage schedule.

**Pharmacokinetics. Onset and Duration.** The onsets of action of Asacol, Dipentum, and Pentasa is delayed because of the release characteristics of their dosage forms; duration of action depends on intestinal transit time. The onset of symptom relief is sooner with the balsalazide than with delayed-released mesalamine.

**Fate.** About 70 ± 10% of oral mesalamine is absorbed in the proximal small bowel when administered in an uncoated product or unbound to a carrier molecule; some absorption can occur in the distal small bowel, but mesalamine is poorly absorbed from the colon. Various oral dosage forms have been formulated to deliver mesalamine topically to the more distal sites of inflammation. (See Dosage Forms and Notes.) After oral administration, about 50% of mesalamine from Pentasa is released in the small bowel and 50% in the colon, although the amount released is patient specific. About 20–30% of released mesalamine is absorbed after oral administration of Asacol or Pentasa; the remainder is excreted in the feces. About 98% of an oral olsalazine dose reaches the large bowel; less than 2% is absorbed. Mesalamine absorption from the enema is pH dependent; neutral solutions are better absorbed than acidic solutions. Rowasa (at pH 4.5) is less than 15% rectally absorbed. Plasma protein binding: mesalamine (40%); N-acetylmesalamine (80%); olsalazine and olsalazine-O-sulfate (>99%). Absorbed mesalamine is rapidly acetylated to N-acetyl-5-aminosalicylate (N-acetylmesalamine) in the intestinal mucosal wall and the liver. A small amount of olsalazine is metabolized to olsalazine-O-sulfate. N-acetylmesalamine is excreted in urine. Less than 1% of a dose of olsalazine is recovered unchanged in urine.

$t_{1/2}$. (Mesalamine) 1 ± 0.5 hr; (N-acetylmesalamine) 7.5 ± 1.5 hr; (olsalazine-O-sulfate) 7 days.

**Adverse Reactions.** Adverse effects are usually less frequent than with oral sulfasalazine. Headache, flatulence, abdominal pain, diarrhea, dizziness, anorexia, and dyspepsia are the most frequent side effects reported with oral formulations and, to a lesser extent, rectal formulation. An acute intolerance syndrome associated with mesalamine occurs in approximately 3% of patients. About 17%
of patients taking olsalazine 1 g/day experience secretory diarrhea. Dermatologic reactions include rash (1%), acne, pruritus, urticaria, alopecia, and photosensitivity. Renal insufficiency occurs in 0.2% of patients; renally impaired patients are at increased risk.149 Rare adverse effects are oral, esophageal, and duodenal ulcerations; hepatotoxicity; jaundice; cholestasis; cirrhosis; liver failure; pancytopenia; leukopenia; agranulocytosis; and anemia.149 Pericarditis, fatal myocarditis, hypersensitivity pneumonitis, pancreatitis, nephrotic syndrome, and interstitial nephritis occur rarely.149,152 Allergic cross-reactions can occur in sulfasalazine-allergic patients.

Contraindications. Pyloric stenosis; intestinal obstruction; salicylate hypersensitivity.

Precautions. Mesalamine is considered safe in pregnancy; however, higher-than-normal doses have resulted in renal insufficiency in the fetus.153,154 Monitor Cr, periodically, especially in those with pre-existing renal impairments.149,152 Use caution in impaired hepatic function. Patients who experience rash or fever with sulfasalazine might have the same reaction to mesalamine or olsalazine; oral desensitization is an option for those who are allergic to mesalamine.155,156 Avoid Rowasa rectal suspension enemas in those with sulfite allergy.

Drug Interactions. In patients on warfarin, olsalazine can increase and mesalamine can decrease INR.157 Omeprazole has no effect on mesalamine absorption.149

Parameters to Monitor. Improvement in abdominal cramping, diarrhea, and rectal bleeding. Monitor for adverse effects, including diarrhea (olsalazine), acute intolerance syndrome, and hypersensitivity reaction. Monitor BUN, Cr, and urinalysis before and periodically during therapy. Monitor INR in patients taking concurrent warfarin.

Notes. The release characteristics of Pentasa are primarily time dependent, whereas those of Asacol are pH dependent; consequently, Asacol might not provide reliable site-specific release of 5-ASA if intestinal pH is inadequate. Balsalazide appears to more consistently distribute and liberate mesalamine in the colonic area, thus having greater effectiveness and less frequent side effects than sulfasalazine or olsalazine.145,158 In 2–3% of patients taking Asacol intact or partly intact, tablets were found in the stools.

There appears to be no clinically important advantage of one oral mesalamine product over another, or over sulfasalazine, in treating or maintaining remission of mild to moderate ulcerative colitis.146,149,159,160 A dose–response relationship exists when mesalamine is used to treat and maintain remission of mild to moderate ulcerative colitis.149,161 A mesalamine preparation might be beneficial in the sulfasalazine-sensitive patient and men who wish to have children because mesalamine does not alter sperm count, morphology, or motility.154 The enema is as effective as oral sulfasalazine or hydrocortisone enema in patients with active mild to moderate left-sided ulcerative colitis and proctitis and is associated with a more rapid response and fewer and milder adverse effects.146,149,160 Patients refractory to oral sulfasalazine and oral or rectal hydrocortisone might respond to rectal mesalamine. Rectal mesalamine combined with oral sulfasalazine
or corticosteroids can enhance induction and maintenance of remission in patients with mild to moderate ulcerative colitis, but the risk of adverse effects is increased.\textsuperscript{162}

In Crohn’s disease involving the ileum or proximal large bowel, oral formulations that deliver mesalamine to the small bowel and colon are preferable to sulfasalazine or olsalazine. Oral mesalamine preparations seem to be effective in treating active mild to moderate Crohn’s disease\textsuperscript{161} (including ileal or ileocolonic) and maintaining remission.\textsuperscript{149,160} Preventing recurrence after surgery with mesalamine prophylaxis in Crohn’s disease is not effective.\textsuperscript{163} Rectal mesalamine appears to be less effective in Crohn’s disease, but efficacy depends on disease location and severity.\textsuperscript{149,159,160}

**OCTREOTIDE ACETATE**  
**Sandostatin, Sandostatin LAR Depot**

**Pharmacology.** Octreotide is a synthetic octapeptide with pharmacologic actions similar to those of somatostatin. The actions of somatostatin are regulated by somatostatin receptors (five known subtypes) located in regions of the brain, leptomeninges, anterior pituitary, endocrine and exocrine pancreas, GI mucosa, and cells of the immune system. Octreotide binds primarily to somatostatin-receptor subtype 2, to a lesser extent to subtype 5, and to an even lesser extent to subtype 3; it does not bind to subtypes 1 and 4. It suppresses the secretion of numerous substances including serotonin, gastrin, vasoactive intestinal peptide (VIP), cholecystokinin, insulin, glucagon, secretin, motilin, pancreatic polypeptide, and growth hormone (GH). It suppresses the luteinizing hormone response to gonadotropin-releasing hormone and the secretion of thyroid-stimulating hormone. It also decreases splanchnic and venous blood flow.\textsuperscript{164,165}

**Administration and Adult Dosage.** (\textit{See also Notes.})
<table>
<thead>
<tr>
<th>INDICATION</th>
<th>OCTREOTIDE ACETATE IMMEDIATE</th>
<th>OCTREOTIDE ACETATE DEPOT</th>
</tr>
</thead>
<tbody>
<tr>
<td>Metastatic carcinoid tumor</td>
<td>SC 100–600 µg/day in 2–4 doses × 2 weeks; dosages of 50–1500 µg/day (median 450 µg/day) have been used.</td>
<td>(Patients not currently receiving octreotide injection) initiate octreotide acetate injection for 2–4 weeks (see left for dosage). If tolerated and effective, then continue with depot formulation, as below.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>(Patients currently receiving octreotide injection) IM intraglutely 20 mg initially q 4 weeks × 2 months. If symptoms do not resolve in 2 months, increase to 30 mg q 4 weeks. If symptoms resolve on 20 mg, then decrease to 10 mg q 4 weeks as a trial period. If symptoms worsen, then increase dose back to 20 mg IM q 4 weeks.⁹</td>
</tr>
<tr>
<td>VIP-secreting tumors (VIPomas)</td>
<td>SC 200–300 µg/day in 2–4 doses × 2 weeks. Dosages of 150–750 µg/day have been used (dosages &gt;450 µg/day are usually not required).</td>
<td>Same as for metastatic carcinoid tumor.</td>
</tr>
<tr>
<td>Acromegaly</td>
<td>SC 50 µg tid, increasing q 2 weeks based on serum IGF-I level.⁸,⁹ Most common dosage is 100 µg tid. Some require dosages up to 500 µg tid, but doses &gt;300 µg/day usually do not have any added biochemical benefit.</td>
<td>(Patients not currently receiving octreotide injection) initiate octreotide acetate injection for 2–4 weeks (see left for dosage). If tolerated and effective, continue with depot formulation (see below)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>(Patients currently receiving octreotide injection) IM intraglutely 20 mg of depot formulation q 4 weeks × 3 months, then base dosage on serum GH level.⁷,⁹</td>
</tr>
</tbody>
</table>

GH = growth hormone; IGF-I = insulin-like growth factor-I; VIP = vasoactive intestinal peptide.

⁸If the patient experiences exacerbation of symptoms, consider giving doses of octreotide acetate injection for a few days at the dose used before switching to the depot formulation.

⁹A more rapid titration can be obtained by drawing multiple GH levels during the 8 hr after the octreotide dose. The goal is to achieve GH <5 µg/L (or IGF-I <1.9 units/mL in men and <2.2 units/mL in women).

⁷Individuals who have received irradiation should discontinue octreotide for about 4 weeks each year. If symptoms worsen or laboratory results are abnormal, resume therapy.

⁹If GH ≤2.5 µg/L and symptoms are controlled, maintain dosage at 20 mg q 4 weeks; if GH >2.5 µg/L and symptoms not controlled, increase dosage to 30 mg q 4 weeks; if GH ≤1 µg/L and symptoms controlled, decrease dosage to 10 mg q 4 weeks; patients whose GH levels and symptoms are not controlled can increase dosage to 40 mg q 4 weeks; dosages >40 mg are not recommended.

⁷Individuals who have received pituitary irradiation should discontinue octreotide for about 8 weeks each year. If symptoms worsen or laboratory results are abnormal, resume therapy.
IV (immediate injection only) same dosage as SC, dilute in 50–200 mL of NS or D5W and infuse over 15–30 min or give by IV push over 3 min. In emergency situations (eg, carcinoid crisis), give by rapid IV bolus.

**Special Populations. Pediatric Dosage.** SC (immediate) (≥1 month) 1–10 μg/kg are well tolerated, and studies of various GI disorders have used widely different dosages in children 3 days–16 yr. Octreotide has been studied in the treatment of hyperinsulinemic hypoglycemia in neonates in different dosages. SC for anti-VIP effects 3.5 μg/kg/day divided q 8 hr. SC for chronic GI bleeding 4–8 μg/kg/day.

**Geriatric Dosage.** Dosage reduction is recommended because of decreased renal clearance, but specific guidelines are not established.

**Other Conditions.** The effect of hepatic disease on the disposition of octreotide is unknown. Reduction of maintenance dosages might be required in patients with renal impairment and those undergoing dialysis.

**Dosage Forms.** Inj (immediate) 50, 100, 200, 500, 1000 μg/mL; Inj (depot) 2, 4, 6 mg/mL.

**Patient Instructions.** (Immediate-release) Instruct patient in sterile SC injection technique. Avoid multiple SC injections at the same site within a short period. Systematically rotate injection sites. Do not use solution if particulates and/or discoloration are present. Store medication in refrigerator but do not allow it to freeze; individual ampules can remain at room temperature for up to 24 hours. Octreotide is stable at room temperature for 14 days if protected from light. Pain at injection site can be minimized by using the smallest volume necessary to obtain the desired dose and by bringing the solution to room temperature before injection, but do not warm artificially. Stop medication and report if symptoms worsen or you have abnormal blood sugar levels or abnormal blood pressure. Inspect the vial for particulate matter or discoloration of the solution; do not use if either is present.

**Missed Doses.** (Immediate-release) If you miss a dose, take it as soon as possible. If it is almost time for your next dose, skip the missed dose and return to your usual dosage schedule. Do not double doses. Although you will not be harmed by forgetting a dose, the symptoms that you are trying to control might reappear. To control your symptoms, your doses should be evenly spaced over 24 hours.

**Pharmacokinetics.** **Onset and Duration.** (Immediate-release) SC peak concentrations occur in 0.4 hr (0.7 hr in acromegaly). Duration is up to 12 hr, depending on tumor type. (Depot) IM initial peak occurs at 1 hr and then slowly decreases over 3–5 days; a second peak appears 2–3 weeks postinjection. Duration is up to 2–3 weeks. Steady-state levels are usually attained after about 12 weeks.

**Fate.** Oral absorption is poor; SC and IV routes are bioequivalent. The drug is 65% protein bound (41% in acromegaly), primarily to lipoprotein and, to a lesser extent, albumin. \( V_d = 0.35 \pm 0.22 \text{ L/kg} \); \( Cl = 0.16 \pm 0.08 \text{ L/hr/kg} \). \( V_d \) and \( Cl \) are both increased in acromegaly; \( Cl \) is decreased in the elderly by 26% and in those with renal impairment. Octreotide exhibits nonlinear pharmacokinetics at dosages of 600 μg/day. About 32% is excreted unchanged in urine.
$t_{1/2}$. 1.5 ± 0.4 hr; increased by 46% in the elderly.

**Adverse Reactions.** Single doses of octreotide acetate can inhibit gallbladder contractility and decrease bile secretion. Approximately half of patients treated for at least 12 months experience cholesterol gallstones or sludge unrelated to age, sex, or dosage. About 22% of patients with acromegaly treated with the depot formulation developed new cholelithiasis, 7% of which were microstones. About 24% of patients with malignant carcinoid who received 18 months of depot therapy developed gallstones; 1% might require cholecystectomy. Five to 10 percent of nonacromegalic patients and 34–61% of acromegalics experience diarrhea, loose stools, nausea, and abdominal discomfort. The severity, but not frequency, is dose dependent and usually occurs with the initial dose, with the symptoms spontaneously resolving within 10–14 days. Hypoglycemia (in 3%) and hyperglycemia (in 16%) are more common in acromegals than in nonacromegals. The frequencies of hypoglycemia (4%) and hyperglycemia (27%) are higher in carcinoid patients treated with the depot formulation. Octreotide suppresses secretion of TSH; alters the balance between insulin, glucagon, and GH; and might be responsible for cardiac conduction abnormalities, which are particularly frequent in acromegaly: bradycardia (25%), conduction abnormalities (10%), and arrhythmias (9%). Pain on injection occurs frequently with the immediate-release formulation and can be minimized by warming the solution before injection and using the smallest possible volume of solution to obtain the appropriate dose. Pain on injection is more frequent with the depot injection, from 2–11% in acromegals to 20–50% in carcinoid patients. Flu-like symptoms, vomiting, flatulence, constipation, and headaches occur in 1–10%. Several cases of pancreatitis have been reported. Steatorrhea also can occur while on long-term therapy. Abnormal Schilling’s tests and decreased vitamin B₁₂ levels have been reported.

**Precautions.** Pregnancy; lactation. Never give depot formulation by the IV or SC route. Use with caution in patients with diabetic gastroparesis because octreotide slows GI transit time; insulin-dependent diabetics might require a reduction in insulin dosage.

**Drug Interactions.** In acromegaly, reducing the dosage of medications that cause bradycardia (eg, β-blockers) might be required. In all patients, the dosage of calcium-channel blocking drugs, diuretics, insulin, or oral hypoglycemics might require an adjustment with concurrent octreotide. Octreotide can decrease the absorption of some orally administered nutrients and drugs (eg, fat, cyclosporine).

**Parameters to Monitor.** Perform ultrasound of the gallbladder periodically during extended therapy. Obtain baseline and periodic total and/or free T₄ levels during long-term therapy. Monitor closely for hyper- or hypoglycemia, especially in diabetics. Periodically monitor vitamin B₁₂ during long-term therapy. Evaluate cardiac function at baseline and periodically during therapy, especially in acromegals. Monitor serum concentrations of drugs whose absorption might be affected by octreotide (eg, cyclosporine). To evaluate response, monitor GH or IGF-I concentrations in acromegals; urinary 5-hydroxyindole acetic acid, plasma serotonin, plasma substance P in carcinoid patients; and plasma VIP in VIPoma patients.
Notes. The absorption of dietary fats can decrease while on octreotide therapy. Zinc levels should be monitored periodically in patients receiving parenteral nutrition and octreotide.

Store depot formulation at 2–8°C. Before administration, leave the drug at room temperature for 30–60 min. Octreotide must be administered immediately after mixing and should only be given IM intragluteally and not in the deltoid region to avoid injection site discomfort.

Store the immediate-release formulation at 2–8°C and protected from light. If stored at room temperature (20–30°C) and protected from light, the product is stable for 14 days. Before SC administration, the solution can be kept at room temperature to decrease injection site discomfort, but do not warm artificially. Octreotide 200 μg/mL is stable for up to 60 days in polypropylene syringes under refrigeration and protected from light. Octreotide is not compatible with parenteral nutrition because of the formation of glycosyl octreotide conjugate.

Sulfasalazine is a conjugate of sulfapyridine linked to mesalamine by an azo bond. This bond is cleaved by colonic bacteria to sulfapyridine and mesalamine, the active moiety. (See Mesalamine Preparations.)

Administration and Adult Dosage. PO for short-term treatment of active mild to moderate ulcerative colitis or Crohn’s disease 4–6 g/day in equally divided doses; do not exceed an interval of 8 hr between nighttime and morning doses; administer with or after meals when feasible. A lower initial dosage can decrease adverse GI effects. PO for maintenance of remission of ulcerative colitis 2–4 g/day in divided doses. Dosages >4 g/day are associated with an increased frequency of adverse effects. Efficacy of sulfasalazine for Crohn’s disease depends on the site of disease activity. (See Notes.) PO for desensitization of allergic patients reinstitute sulfasalazine at a total daily dosage of 50–250 mg; thereafter, double the daily dosage q 4–7 days until the desired therapeutic effect is achieved. If symptoms of sensitivity recur, discontinue sulfasalazine. Do not attempt desensitization in patients who have histories of agranulocytosis or anaphylactic reactions during previous sulfasalazine therapy. Consider mesalamine instead of desensitization in sulfasalazine-sensitive patients.

Special Populations. Pediatric Dosage. (<2 yr) contraindicated; (≥2 yr) PO for short-term treatment of active mild to moderate ulcerative colitis or Crohn’s disease 40–60 mg/kg/day in 3–6 equally divided doses. Dosages up to 75 mg/kg/day or up to 5 g/day in divided doses have been used. Additional age-related information is available. PO for maintenance of remission of ulcerative colitis 30 mg/kg/day in 4 equally divided doses.

Geriatric Dosage. No dosage reduction is necessary. However, older patients might have renal impairment.

Other Conditions. Consider dosage reduction in severe renal or hepatic impairment.

Dosage Forms. Tab 500 mg; EC Tab 500 mg.
**Patient Instructions.** Take each dose after meals or with food and drink at least 1 full glass of water with each dose; drink several additional glasses of water daily. This medication must be taken continually to be effective. It is often necessary to continue medication even when symptoms such as diarrhea and abdominal cramping have been controlled. Report any nausea, vomiting, abrupt change in character or volume of stools, or skin rashes. Sulfasalazine can cause orange-yellow discoloration of the urine or skin. Reversible infertility can occur in males. Contact your physician or pharmacist if whole tablets appear in the stool.

**Missed Doses.** If you miss a dose, take it as soon as possible. If it is almost time for your next dose, skip the missed dose and return to your usual dosage schedule. Do not double doses.

**Pharmacokinetics.**

**Onset and Duration.** Maximum effect is in 1–2 weeks; duration is 10 ± 2 hr after an oral dose.117

**Fate.** Sulfasalazine is 25–30% absorbed from the small intestine, but the absorbed drug is almost completely secreted unchanged in the bile. It is then metabolized in the large bowel by intestinal bacteria to sulfapyridine and mesalamine. Most of the sulfapyridine is absorbed from the bowel. Plasma protein binding: sulfasalazine (>99%); sulfapyridine (50%); mesalamine (55 ± 15%); N-acetylmesalamine (80%). Sulfapyridine is metabolized by acetylation to acetylsulfapyridine. Acetylsulfapyridine concentration depends on acetylator phenotype: slow acetylators have higher serum sulfapyridine concentrations, fast acetylators have lower serum sulfapyridine concentrations. After an oral dose of sulfasalazine, about 91% of sulfapyridine is recovered in the urine in 3 days as sulfapyridine, its metabolites, and small amounts of sulfasalazine. Mesalamine is eliminated primarily in the feces; only a small portion is absorbed, metabolized, and excreted in the urine as N-acetylmesalamine.117

$t_{1/2}$ (Sulfapyridine) 9 ± 4 hr, depending on acetylator phenotype.117 (See also Mesalamine Preparations.)

**Adverse Reactions.** Anorexia, nausea, vomiting, dyspepsia, and headache occur in about one-third of patients and are related to serum sulfapyridine concentrations. These side effects usually resolve with dosage reduction.154,175 Leukopenia occurs frequently. Mild allergic reactions such as rash, pruritus, and fever are common.175 Decreased folate absorption leading to anemia can occur, so folic acid supplementation is recommended.159,175 Rare toxic hypersensitivity reactions (caused by sulfapyridine) are neutropenia, agranulocytosis, hepatitis, pancreatitis, pericarditis, pneumonitis, peripheral neuropathy, and severe hemolytic anemia.159,174,175 Sulfasalazine can cause orange-yellow discoloration of the skin and precipitate acute attacks of porphyria. In men, sulfasalazine frequently leads to a reversible decrease in sperm count and abnormal sperm morphology and motility.154,175

**Contraindications.** Intestinal or urinary obstruction; porphyria; infants <2 yr; hypersensitivity to sulfasalazine, its metabolites, sulfonamides, or salicylates.

**Precautions.** Pregnancy, despite reports of safety; lactation. Use with caution in patients with renal or hepatic impairment, blood dyscrasias, slow acetylators, bronchial asthma, G-6-PD deficiency, or severe allergies.
Drug Interactions. Decreased digoxin bioavailability has been reported when sulfasalazine is concurrently administered.

Parameters to Monitor. Monitor therapeutic response (decrease in degree and frequency of diarrhea, rectal bleeding, abdominal cramping) and adverse effects (headache, anorexia, dyspepsia, nausea, hypersensitivity reactions). Obtain baseline and periodic serum electrolytes, liver function tests, CBC, reticulocyte counts, and urinalysis. Monitor serum folate periodically in patients on long-term therapy. Monitor serum digoxin levels during initiation and after discontinuation of sulfasalazine.

Notes. There appears to be no important therapeutic advantage of sulfasalazine over oral mesalamine when used to treat or maintain remission of ulcerative colitis; however, the higher sulfasalazine dosages used to treat active disease are associated with an increased frequency of adverse effects. Crohn’s disease patients with involvement of the ileum do not respond as well to sulfasalazine as those with only large bowel disease. Combining sulfasalazine with an oral or rectal corticosteroid or with rectal mesalamine might be beneficial in patients with ulcerative colitis who do not respond to single-drug therapy. In ulcerative colitis patients receiving maintenance therapy, there was less absorption and greater acetylation of 5-ASA with sulfasalazine or olsalazine than with mesalamine (Asacol). Sulfasalazine also has been used to treat ankylosing spondylitis and rheumatoid arthritis. Occasionally, the EC tablet can appear whole in the stool; if this occurs, consider switching the patient to the uncoated form of sulfasalazine or another mesalamine formulation.

Pharmacology. Ursodiol (ursodeoxycholic acid) is a hydrophilic bile acid used to dissolve small (<20 mm), noncalcified, radiolucent cholesterol gallstones in mildly symptomatic patients with functioning gallbladders who cannot undergo a cholecystectomy. It is also used to treat primary biliary cirrhosis. The exact mechanism of action of ursodiol is unclear, but it is thought to have a hepatoprotective effect by displacing accumulated toxic bile acids with hydrophilic bile acids, to promote secretion of toxic bile acid salts from the bile ducts and suppress the synthesis of chenodeoxycholic acid, and to act as an immunosuppressive agent by downregulating the antigen expression in hepatocytes in patients with primary biliary cirrhosis or primary sclerosing cirrhosis. Ursodiol improves liver function tests, liver histology, and certain immune markers; relieves pruritus in some patients; and can extend the period before death or to liver transplantation. Ursodiol also appears to be effective in decreasing episodes of rejection and improving 1-yr survival rates after liver transplantation. Patients undergoing bone marrow transplantation might benefit from ursodiol therapy through prevention of hepatic veno-occlusive disease.

Administration and Adult Dosage. All doses should be administered with food. PO for gallstone dissolution: 8–10 mg/kg/day in 2–3 divided doses. Complete gallstone dissolution usually requires 6–24 months of treatment, and treatment should be continued for at least 3 months after stones or sludge are not apparent.
on ultrasound. **PO for prevention of gallstones in patients with rapid weight loss** 300 mg bid. **PO for primary biliary cirrhosis** 13–15 mg/kg/day in 4 divided doses. **PO for prevention of hepatic veno-occlusive disease in bone marrow transplant** (<90 kg) 300 mg bid; (≥90 kg) 300 mg tid (or 300 mg q AM and 600 mg q PM).\(^{175}\) **PO as an adjunct to immunosuppressants after liver transplantation** 10–15 mg/kg/day in divided doses.\(^{176,177}\)

**Special Populations. Pediatric Dosage.** Safety and efficacy not established; pediatric use has been reported. **PO for cystic fibrosis in patients with liver disease** 5–20 mg/kg/day in divided doses. Higher doses might be required in this patient population.\(^{179}\) **PO for obese children with liver abnormalities** 10–12.5 mg/kg/day in 2 divided doses.\(^{180}\)

**Geriatric Dosage.** No dosage reduction is necessary.

**Dosage Forms.** Cap 300 mg; Tab 250 mg. Ursodiol can be formulated into a suspension.\(^{181,182}\)

**Adverse Reactions.** Ursodiol is relatively safe, with minimal side effects. The most common adverse effects are diarrhea, nausea, vomiting, dyspepsia, abdominal pain, and arthritis.

**Drug Interactions.** Bile acid sequestering agents (ie, cholestyramine, colestipol) and aluminum-containing antacids reduce ursodiol absorption; thus, the two drugs should be taken at least 2 hr apart. Oral contraceptives, estrogens, and lipid-lowering agents (eg, clofibrate) increase cholesterol secretion, thereby increasing the risk of developing cholesterol gallstones; using any of these agents can counteract the effectiveness of ursodiol.

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Coagulants and Anticoagulants

**ABCIXIMAB**

**Pharmacology.** Abciximab is a chimeric human-murine monoclonal antibody Fab fragment that binds to and irreversibly inhibits the platelet glycoprotein IIb/IIIa receptor. Blockade of the glycoprotein IIb/IIIa receptor prevents fibrinogen from binding, thereby inhibiting platelet aggregation. Abciximab also binds to the vitronectin receptor found on platelets, endothelial cells, monocytes, and smooth muscle cells; the clinical relevance of this is unknown. Abciximab inhibits platelet aggregation and prolongs bleeding time in a dose-dependent manner.1,2

**Administration and Adult Dosage.** IV for percutaneous coronary intervention 0.25 mg/kg as a bolus 10–60 min before starting percutaneous coronary intervention and then 0.125 µg/kg/min (up to 10 µg/min) by continuous infusion for 12 hr. IV for unstable angina and planned percutaneous intervention within 24 hr 0.25 mg/kg as a bolus and then 0.125 µg/kg/min (up to 10 µg/min) by continuous infusion for 18–24 hr, concluding 1 hr after the percutaneous coronary intervention. (See Parameters to Monitor and Notes.)

**Special Populations.** *Geriatric Dosage.* Same as adult dosage.

**Dosage Forms.** Inj 2 mg/mL.

**Pharmacokinetics.** Onset and Duration. Rapid inhibition of platelet function after IV administration. Platelet function gradually recovers after discontinuation of the IV infusion; bleeding time approaches baseline values within 24 hr and ex vivo platelet aggregation approaches baseline levels within 48 hr. Low levels of glycoprotein IIb/IIIa inhibition are detectable for up to 14 days after administration.1

**Fate.** Abciximab is rapidly cleared from the plasma after administration by rapid binding to the glycoprotein IIb/IIIa receptor.

\[ t_{1/2} \alpha \text{ phase } <10 \text{ min}; \beta \text{ phase } 30 \text{ min}. \]

**Adverse Reactions.** Bleeding, particularly from vascular access sites, occurs frequently. To minimize bleeding complications, care of the femoral artery access site is important and lower doses of unfractionated heparin are necessary during the percutaneous coronary intervention. (See Notes.) If serious bleeding complications occur, discontinue abciximab and transfuse platelets, if needed, to restore platelet function. Thrombocytopenia (<100,000/µL) has been reported in 2.6–5.6% of patients; severe thrombocytopenia (<50,000/µL) has occurred in 0.9–1.7% of patients. Thrombocytopenia can occur rapidly after administration and might require platelet transfusions if reversal is necessary.1
Contraindications. Active internal bleeding; recent (within 6 weeks) clinically significant GI or GU bleeding; history of CVA within 2 yr or CVA with significant residual neurologic deficit; bleeding diathesis; administration of oral anticoagulants within 7 days unless PT ≤ 1.2 times control; thrombocytopenia (<100,000/µL); recent (within 6 weeks) major surgery or trauma; intracranial neoplasm, AV malformation, or aneurysm; severe uncontrolled hypertension; presumed or documented history of vasculitis; use or planned use of IV dextran before or during percutaneous coronary intervention.

Precautions. Use with caution in patients being treated concomitantly with other antithrombotic drugs including thrombolytics, unfractionated heparin, low-molecular-weight heparin, oral anticoagulants; NSAIDs; and other drugs that increase bleeding risk.

Parameters to Monitor. Monitor CBC including platelet count; prothrombin time; aPTT; and activated clotting time at baseline. Maintaining the activated clotting time at 200–300 sec during percutaneous coronary intervention minimizes the risk of bleeding complications. Monitor platelet count 2–4 hr after the IV bolus and again at 24 hr or before hospital discharge, whichever occurs first. If prolonged infusion of unfractionated heparin is necessary after percutaneous coronary intervention, maintain aPTT at 60–85 sec.

Notes. Abciximab must be filtered using a sterile, nonpyrogenic, low-protein-binding 0.2 or 0.22 µ filter either at admixture or during administration with an in-line filter. To minimize the risk of bleeding complications, the following care for the arterial access site is recommended: maintain patient on complete bed rest with the affected limb restrained in a straight position while vascular sheaths are in place; discontinue unfractionated heparin immediately after percutaneous coronary intervention; remove vascular sheaths within 6 hr of completing the procedure if aPTT ≤ 50 sec or activated clotting time ≤ 175 sec; after sheath removal, apply pressure to the femoral artery for at least 30 min with manual compression or a mechanical device; and maintain bed rest for 6–8 hr after sheath removal. To minimize bleeding complications, the following periprocedural heparin dosage is recommended: if baseline activated clotting time ≤ 150 sec, administer 70 units/kg heparin IV bolus; if 150–199 sec, administer 50 units/kg heparin IV bolus; if ≥ 200 sec, do not administer heparin. During percutaneous coronary intervention, administer 20 units/kg heparin IV boluses as necessary to maintain activated clotting time at 200–300 sec.

Pharmacology. Alteplase (recombinant tissue-type plasminogen activator [rt-PA]) is a 1-chain tissue plasminogen activator (fibrinolytic) produced by recombinant DNA technology. It has a high affinity for fibrin-bound plasminogen, allowing activation on the fibrin surface. Most plasmin formed remains bound to the fibrin clot, minimizing systemic effects.

Administration and Adult Dosage. Accelerated IV infusion for clot lysis after MI (preferred) 15 mg as a bolus, followed by 0.75 mg/kg (up to 50 mg) over 30 min, and then 0.5 mg/kg (up to 35 mg) over the next 60 min. Start heparin infu-
sion (titrated to an aPTT of 1.5–2.0 times control) with or at completion of the alteplase infusion and continue for at least 48 hr. (See Notes.) Alternatively, IV infusion for clot lysis after MI 60 mg over 1 hr (6–10 mg in the first 1–2 min) and then 20 mg/hr for 2 hr to a total of 100 mg (for patients <65 kg, administer a dose of 1.25 mg/kg over 3 hr). Begin as soon as possible after acute MI symptoms. Adjunctive heparin is also recommended.5,7,8 IV infusion for pulmonary embolism 100 mg over 2 hr. Institute heparin infusion immediately after alteplase infusion when the aPTT or thrombin time returns to 2 times normal. Alternatively, 0.6 mg/kg as a single dose over 2 min in addition to heparin infusion has been used successfully.9 IV infusion for acute ischemic stroke 0.9 mg/kg, to a maximum of 90 mg; give 10% initially as a bolus, with the remainder given over the next 60 min. Avoid anticoagulants or antiplatelet drugs for 24 hr after treatment.10,11 IV for catheter clearance slowly inject 0.5 mg (1 mL) into the occluded catheter port. If catheter volume exceeds 1 mL, slowly inject a sufficient volume of NS to fill the catheter. Allow the solution to dwell for 60 min and then aspirate and flush the catheter with NS. If unsuccessful, repeat with escalating doses of alteplase (eg, 1 mg, 2 mg) to a maximum of 2 mg.12

Special Populations. Pediatric Dosage. Safety and efficacy not established.

Geriatric Dosage. Same as adult dosage.

Dosage Forms. Inj 50, 100 mg.

Pharmacokinetics. Onset and Duration. Duration is several hours because of binding with fibrin. However, rethrombosis after reperfusion appears to be inversely proportional to serum half-life.5

Fate. There is rapid uptake by hepatocytes and fibrin binding. \( V_c = 3.8–6.6 \text{ L} \) and \( V_d = 0.1 \pm 0.01 \text{ L/kg} \); \( Cl = 0.6 \pm 0.24 \text{ L/hr/kg} \).5,13

\( t_{1/2} \), \( \alpha \) phase 4.8 ± 2.4 min; \( \beta \) phase 26 ± 10 min.13

Adverse Reactions. Bleeding from GI and GU tracts and ecchymoses occur frequently. Retroperitoneal or gingival bleeding or epistaxis occur occasionally. Superficial bleeding from trauma sites also can occur. The overall risk of intracranial hemorrhage is 0.1–0.75%.10 In ISIS-3, the rates for definite or possible cerebral bleed were: rt-PA (duteplase, a 2-chain form of alteplase), 0.5%; streptokinase, 0.2%; anistreplase, 0.7%.14 Independent risk factors for thrombolytic-induced intracranial hemorrhage with alteplase are age >65 yr, body weight <70 kg, and hypertension on hospitalization.7

Contraindications. Active internal bleeding; history of CVA; recent (within 2 months) intracranial or intraspinal surgery or trauma; intracranial neoplasm, AV malformation, or aneurysm; bleeding diathesis; severe uncontrolled hypertension.

Precautions. Use with caution in the following: pregnancy; recent (within 10 days) major surgery, trauma, GI or GU bleeding; cerebrovascular disease; systolic blood pressure ≥180 mm Hg, diastolic blood pressure ≥110 mm Hg; high likelihood of left heart thrombus; acute pericarditis; subacute bacterial endocarditis; hemostatic defects; significant liver dysfunction; septic thrombophlebitis; age >75 yr; concurrent oral anticoagulants. Avoid IM injections and noncompressible
arterial punctures; minimize arterial and venous punctures and excessive patient handling. Stop immediately if severe bleeding or anaphylactoid reaction occurs.

**Drug Interactions.** Preliminary data from a nonrandomized study suggest that concurrent IV nitroglycerin therapy impairs the thrombolytic effect of alteplase in acute MI.\(^{15}\) Anticoagulants or antiplatelet drugs can increase the risk of bleeding.

**Parameters to Monitor.** For short-term thrombolytic therapy of MI, laboratory monitoring is of little value. No correlation has been made between clotting test results and likelihood of hemorrhage or efficacy.\(^{5}\)

**Notes.** Other than cerebral hemorrhage, no clear differences in bleeding risk have been observed with the various thrombolytics.\(^{5,7}\) Data from the ISIS-3 trial show the 5-week mortalities for **duteplase**, **streptokinase**, and **anistreplase** to be virtually identical.\(^{14}\) Based on the GUSTO trial, some investigators have suggested that the accelerated alteplase regimen be used for patients <75 yr with anterior or large infarctions presenting within 4 hr of symptoms. The absolute survival advantage over streptokinase was 0.9%, representing a 14% risk reduction.\(^{3,16}\) Double-bolus alteplase (50 mg IV over 1–3 min followed by 40–50 mg IV 30 min later) was compared with accelerated infusion alteplase to shorten and simplify administration. The double-bolus method was associated with a slightly higher rate of intracranial hemorrhage and is not recommended.\(^{17}\) In a study on catheter clearance, 96.5% of catheters were cleared successfully, 86.2% with a dose of 0.5 mg, 8.6% with 1 mg, and 1.7% with 2 mg.\(^{12}\)

**ANISTREPLASE**

**Pharmacology.** Anistreplase (anisoylated plasminogen-streptokinase activator complex) is an acylated form of the streptokinase–plasminogen complex that is temporarily inactive. After deacylation, the streptokinase–plasminogen complex promotes thrombolysis by converting plasminogen to the proteolytic enzyme plasmin. Thrombolysis occurs through the action of plasmin on fibrin.\(^{7,10,13,18,19}\)

**Adult Dosage.** IV for post-MI clot lysis 30 units over 2–5 min as a single injection. Adjunctive IV heparin is associated with higher bleeding rates than aspirin alone in anistreplase-treated patients and offers no additional improvement in outcome.

**Dosage Forms.** Inj 30 units.

**Pharmacokinetics.** Deacylation and thrombolysis begin immediately after injection. Duration of fibrinolytic activity is 4–6 hr. Anistreplase undergoes deacylation and local inactivation in the circulation by inhibitor complex formation and proteolysis and, to a lesser extent, is metabolized rapidly by the liver. \(V_d = 0.084 \pm 0.027\) L/kg, with a Cl of \(0.055 \pm 0.02\) L/hr/kg and half-life of 1.2 ± 0.4 hr.

**Adverse Reactions.** Data from ISIS-3 indicated that bleeding was slightly more common with anistreplase than with streptokinase or rt-PA; however, major bleeding rates were similar. In ISIS-3 the rates for definite or possible cerebral bleeding were: **anistreplase**, 0.7%; **streptokinase**, 0.2%; rt-PA (duteplase, a 2-chain form of alteplase), 0.5%.\(^{14}\) Allergic reactions similar to those reported with streptokinase are rash, erythema, bronchoconstriction, and, rarely, anaphylaxis. Precautions and monitoring parameters are the same as those for streptokinase.
**Notes.** Ease of administration (30 units given over 2–5 min as a single IV injection) is a potential advantage of anistreplase for the emergent treatment of acute MI in some settings (eg, in the field). However, anistreplase is more expensive than streptokinase, has the same allergy profile, offers no efficacy advantage, and might be associated with a slightly higher bleeding risk.

**ARGATROBAN**

**Pharmacology.** Argatroban is a modified amino acid that is a reversible, competitive, direct thrombin inhibitor used as an anticoagulant in patients with heparin-induced thrombocytopenia.20

**Administration and Adult Dosage.** IV as an anticoagulant 2–10 µg/kg/min, titrating aPTT to 1.5–3 times control. Dosage might need to be reduced in renal or hepatic impairment.

**Dosage Forms.** Inj 100 mg/mL.

**Pharmacokinetics.**

**Onset and Duration.** Onset <10 min after a bolus or 1–3 hr after start of infusion without a bolus; the effect dissipates with a half-life of 18–41 min after cessation.21

**Fate.** The drug is metabolized in the liver to three metabolites that are excreted renally.

**Adverse Reactions.** Bleeding is the most frequent complication but is usually minor. No specific reversal agent exists. Dose-related prolongation of PT occurs.20 Other common side effects include dyspnea, hypotension, and fever. Repeat exposure does not appear to predispose to immunologic reactions or excessive anticoagulation.

**Parameters to Monitor.** Monitor aPTT 2 hr after initiation of therapy or dosage adjustment and then once daily after stable anticoagulation has been achieved.

**Notes.** To initiate warfarin therapy, add the desired PT elevation to the argatroban-induced PT elevation (not to exceed 30 sec) and begin warfarin. Once this PT is achieved, stop argatroban. Argatroban has been used as an anticoagulant during extracorporeal circulation22 and percutaneous coronary intervention.

**CLOPIDOGREL BISULFATE**

**Pharmacology.** Clopidogrel is an antiplatelet agent that prevents platelet aggregation by direct inhibition of ADP binding to receptor sites, inhibiting subsequent activation of the glycoprotein IIb/IIIa complex. This action is irreversible; therefore, platelets exposed to clopidogrel are inhibited for their life spans.

**Adult Dosage.** PO for reduction of stroke, MI, or vascular death 75 mg once daily. A loading dose of 300 mg on the first day is often used to hasten the onset of action.

**Dosage Forms.** Tab 75 mg.

**Pharmacokinetics.** Clopidogrel is rapidly absorbed; bioavailability is about 50%; 98% is bound to plasma proteins. The parent compound has no platelet-inhibiting activity and undergoes extensive hepatic metabolism to a carboxylic acid derivative (main metabolite) and an unidentified active metabolite. The half-life of the carboxylic acid metabolite is about 8 hr.
Adverse Reactions. The most frequent side effects are diarrhea in 4.5%, rash in 4.2%, GI hemorrhage in 2%, and GI ulcers in 0.7% of patients. Serious, but less frequent, side effects are intracranial hemorrhage in 0.4% and severe neutropenia in 0.04%. Clopidogrel has been associated with the development of thrombotic thrombocytopenia purpura.23 The drug is contraindicated in active bleeding as in patients with peptic ulcer or intracranial hemorrhage. Use with caution in patients at increased risk of bleeding from trauma, surgery, or other conditions. Clopidogrel should be discontinued 7 days before elective surgery if an antiplatelet effect is not desired.

Drug Interactions. Use with caution in patients receiving anticoagulants or drugs that inhibit platelet function including NSAIDs.

Notes. The overall risk reduction for clopidogrel was 8.7% greater than that for aspirin in the CAPRIE study in patients at risk for ischemic events.24

Pharmacology. Dalteparin is a low-molecular-weight heparin (average mass 3000–8000 daltons) prepared by depolymerization and chromatographic purification of unfractionated porcine intestinal mucosa heparin. Other pharmacologic properties are similar to those of enoxaparin.25 (See Low-Molecular-Weight Heparins Comparison Chart.)

Adult Dosage. SC for prevention of ischemic complications in unstable angina and non–Q-wave MI 120 IU/kg (to a maximum of 10,000 IU) q 12 hr with concurrent oral aspirin 81–160 mg once daily. Continue treatment until patient is clinically stable, usually 5–8 days. SC for prevention of DVT and PE after abdominal surgery 2500 IU 1–2 hr before surgery and once daily for 5–10 days. SC for prevention of DVT and PE after abdominal surgery in high-risk patients (eg, with malignancy) 5000 IU the evening before surgery and then once daily postoperatively, or 2500 IU 1–2 hr before surgery, 2500 IU 4–8 hr postoperatively, and then 5000 IU once daily for 5–10 days. SC for prevention of DVT and PE after hip replacement surgery 2500 IU 2 hr before and 12 hr after surgery, and then 5000 IU once daily for 6–13 days; or 5000 IU 10–14 hr before surgery, 5000 IU 4–8 hr postoperatively, and then 5000 IU once daily thereafter. For postoperative initiation, give 2500 IU 4–8 hr postoperatively and then 5000 IU once daily.

Dosage Forms. Inj 2500 IU/0.2 mL, 5000 IU/0.2 mL; 95,000 IU.

Pharmacokinetics. Bioavailability after SC injection is about 87 ± 6%. After SC doses Vd is 0.04–0.06 L/kg; after a single IV dose Cl is 0.025 ± 0.0054 L/hr/kg and the terminal half-life is 2.1 ± 0.3 hr; after SC administration the apparent half-life is 3–5 hr. Dalteparin is eliminated primarily by the kidney.

Adverse Reactions. Overall, rates of major and minor bleeding complications are similar to those with unfractionated heparin. Hematoma or pain at the injection site occurs frequently. Thrombocytopenia occurs in fewer than 1% of patients; however, dalteparin should be used with extreme caution in patients with histories of heparin-induced thrombocytopenia (in vitro platelet testing is recommended before use). Rash, fever, skin necrosis, and anaphylactoid reactions occur rarely.
Contraindications. (See Enoxaparin Sodium.) Patients undergoing regional anesthesia should not receive dalteparin for unstable angina or non–Q-wave MI.

Precautions. (See Enoxaparin.)

Pharmacology. Enoxaparin is a low-molecular-weight heparin (average mass 3500–5500 daltons) prepared by depolymerization of unfractionated porcine intestinal mucosa heparin. Like unfractionated heparin, enoxaparin binds with antithrombin III, accelerating the rate at which antithrombin III neutralizes several activated clotting factors. However, enoxaparin has many biologic properties that differ from those of unfractionated heparin. Enoxaparin has a higher ratio of antifactor Xa to antifactor IIa activity, reduced interactions with platelets, and less lipoprotein lipase–releasing activity. It also has a lower affinity for platelet factor 4, von Willebrand factor (VIIIIR), and vascular endothelium. At recommended dosages, single injections do not markedly affect platelet aggregation, prothrombin time, or aPTT.26–29 (See Low-Molecular-Weight Heparins Comparison Chart.)

Administration and Adult Dosage. SC for prevention of DVT and PE after hip or knee replacement surgery 30 mg bid for 7–10 days started 12–24 hr postoperatively. SC for prevention of DVT and PE after abdominal surgery 40 mg once daily for 7–10 days started 2 hr before surgery. SC for active DVT treatment with and without PE 1 mg/kg q 12 hr initiated with warfarin therapy; continue for at least 5 days and until a warfarin target INR of 2.0 is achieved on 2 consecutive days. SC for unstable angina or non–Q-wave MI 1 mg/kg q 12 hr with concurrent aspirin 100–325 mg once daily. Continue treatment for at least 2 days or until patient is clinically stable, usually 2–8 days.

Special Populations. Pediatric Dosage. SC for treatment (neonates) 1.6 mg/kg bid; (older infants and children) 1 mg/kg bid dosages have been used.

Geriatric Dosage. Elderly patients might have reduced elimination; use with caution in these patients.

Other Conditions. Elimination can be delayed in renal insufficiency; use with caution in these patients.

Dosage Forms. Inj 30 mg/0.3 mL; 40 mg/0.4 mL; 60 mg/0.6 mL; 80 mg/0.8 mL; 100 mg/1 mL.

Pharmacokinetics. Onset and Duration. Peak antifactor Xa occurs 3–5 hr after SC injection and persists for about 12 hr after a 40 mg SC injection.

Fate. Mean absolute bioavailability after SC injection is about 92%. \( V_d \) is about 6 L and Cl is about 1.5 L/hr after IV administration. Some hepatic desulfation and depolymerization occur, but most of the drug is eliminated renally.

\( t_{1/2} \). The apparent half-life after SC administration is 4.5 hr.

Adverse Reactions. Overall, rates of major and minor bleeding complications in comparative studies with unfractionated heparin are similar. In clinical trials of enoxaparin in hip replacement surgery, major bleeding occurred in 4% of patients compared with 6% of patients treated with unfractionated heparin. Thrombocytopenia, fever, pain on injection, asymptomatic increases in transaminase levels,
hypochromic anemia, and edema occur frequently. Skin necrosis occurs occasionally.

**Contraindications.** Hypersensitivity to heparin or pork-derived products; active major bleeding; thrombocytopenia associated with positive in vitro testing for antiplatelet antibody in the presence of a low-molecular-weight heparin.

**Precautions.** If epidural or spinal anesthesia or spinal puncture is used, patients receiving low-molecular-weight heparins for prevention of thromboembolic complications are at risk for developing epidural or spinal hematoma, which can result in permanent paralysis. The risk of these events can increase when postoperative indwelling epidural catheters are used. Use with caution in patients with renal impairment.

**Drug Interactions.** Use with caution in patients receiving oral anticoagulants or drugs that inhibit platelet function, including NSAIDs.

**Parameters to Monitor.** Monitor CBC, including platelet count, and stool for occult blood periodically; aPTT monitoring is not required.
<table>
<thead>
<tr>
<th>DRUG</th>
<th>DOSAGE FORMS</th>
<th>ADULT DOSAGE</th>
<th>AVERAGE MASS (DALTONS)</th>
<th>AF-XA&lt;sup&gt;a&lt;/sup&gt; (IU/MG)</th>
<th>AF-XA/AF-IIA&lt;sup&gt;b&lt;/sup&gt; RATIO</th>
<th>HALF-LIFE (HR)</th>
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<tr>
<td>Dalteparin</td>
<td>Inj 2500, 5000, 10,000 IU/mL, SC for DVT prophylaxis, 2500–5000 IU/day</td>
<td>3000–8000</td>
<td>160</td>
<td>4:1</td>
<td>2.8–4</td>
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<td>Fragmin</td>
<td>Inj 10,000 IU/mL. SC for DVT treatment, unstable angina or non-Q-wave MI 120 IU/kg bid.</td>
<td>120 IU/kg bid.</td>
<td>6500</td>
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<td>3.35:1</td>
<td>18.3</td>
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<td>Danaparoid&lt;sup&gt;c&lt;/sup&gt;</td>
<td>Inj 750 units/0.6 mL. SC for DVT prophylaxis 750 units bid SC for DVT treatment 2000 units q 12 hr.</td>
<td>3500–5500</td>
<td>100</td>
<td>2.7:1</td>
<td>3.5–5.9</td>
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<tr>
<td>Orgaran</td>
<td>Inj 40 mg/0.4 mL, 60 mg/0.6 mL, 80 mg/0.8 mL, 100 mg/1 mL. SC for DVT treatment unstable angina or non-Q-wave MI 1 mg/kg q 12 hr.</td>
<td>30 mg bid; SC post-abdominal surgery 40 mg/day</td>
<td>(continued)</td>
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<tr>
<td>DRUG</td>
<td>DOSAGE FORMS</td>
<td>ADULT DOSAGE</td>
<td>AVERAGE MASS (DALTONS)</td>
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<td>AF-XA/AF-II(^a) RATIO</td>
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<td><strong>Fondaparinux</strong></td>
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<td>SC for DVT prophylaxis 1.5–3 mg once daily. SC for DVT treatment 7.5 mg once daily.</td>
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<td><strong>Arixtra</strong></td>
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<td>(Investigational—Sanofi)</td>
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<tr>
<td><strong>Nadroparin</strong></td>
<td>—</td>
<td>SC for DVT prophylaxis 4400 IU once daily SC for DVT treatment 90–92 IU/kg bid.</td>
<td>4500</td>
<td>85</td>
<td>3.2:1</td>
<td>2.3–5</td>
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<td>Fraxiparin</td>
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<td>(Investigational—Sanofi)</td>
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<tr>
<td><strong>Tinzaparin</strong></td>
<td>Inj 20,000 IU/mL.</td>
<td>SC for DVT prophylaxis 50–75 IU/kg once daily SC for DVT treatment 175 IU/kg once daily.</td>
<td>4900</td>
<td>86</td>
<td>1.9:1</td>
<td>1.85</td>
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<td>Innohep</td>
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\(^a\)Antifactor Xa activity.

\(^b\)Antifactor Xa:antifactor IIa ratio. The ratio for unfractionated heparin is 1.

\(^c\)A heparinoid; mixture of low-molecular-weight sulfated nonheparin glycosaminoglycans: heparan sulfate (84%), dermatan sulfate (12%), and chondroitin sulfate (4%).

\(^d\)Fondaparinux is a pure XA inhibitor.

From references 5, 25, 26, 28, 30, and 31.
Pharmacology. Eptifibatide is a synthetic, cyclic heptapeptide that reversibly binds to and inhibits the platelet glycoprotein IIb/IIIa receptor. Inhibition of the glycoprotein IIb/IIIa receptor prevents fibrinogen from binding, thereby preventing platelet aggregation. Eptifibatide inhibits platelet aggregation and prolongs bleeding time in a dose-dependent manner.1

Administration and Adult Dosage. IV for unstable angina or non–Q-wave MI (acute coronary syndrome) 180 µg/kg as a bolus and then 2 µg/kg/min by continuous infusion. Continue infusion for up to 72 hr, until hospital discharge or CABG surgery, whichever occurs first. Should percutaneous coronary intervention be performed, continue infusion for 18–24 hr after completing procedure (up to 96 hr total duration). Concomitant heparin therapy is recommended. (See Notes.) IV for percutaneous intervention 180 µg/kg as a bolus and then 2 µg/kg/min continuous infusion for 20–24 hr. Give a second bolus of 180 µg/kg 10 min after the first.

Special Populations. Other Conditions. In patients with renal insufficiency: for Crs 2–4 mg/dL, give 180 µg/kg as a bolus and then 1 µg/kg/min continuous infusion. For percutaneous intervention, give a second bolus of 180 µg/kg 10 min after the first.

Dosage Forms. Inj 0.75, 2 mg/mL.

Pharmacokinetics. Onset and Duration. Rapid inhibition of platelet function occurs after IV administration. Platelet function recovers soon after discontinuation of the IV infusion; bleeding time returns to baseline within 30 min and ex vivo platelet aggregation approaches baseline levels within 2–4 hr.1

Fate. Renal elimination accounts for about 50% of the total body clearance of eptifibatide. Cl is 0.055–0.058 L/kg/hr.

Adverse Reactions. Bleeding, particularly from vascular access sites, occurs frequently. Oropharyngeal, GI, and GU bleeding also can occur. The frequency of thrombocytopenia is equal to that of placebo.1,32,33

Contraindications. Active internal bleeding within previous 30 days; history of CVA within 30 days or any history of hemorrhagic CVA; bleeding diathesis; thrombocytopenia (<100,000/µL); recent (within 6 weeks) major surgery or trauma; severe uncontrolled hypertension; current or planned use of another parenteral glycoprotein IIb/IIIa inhibitor; dependency on hemodialysis.

Precautions. (See Abciximab.) Use caution in elderly patients because eptifibatide clearance might be reduced, increasing risk of bleeding.

Parameters to Monitor. Monitor CBC (including platelet count), PT, aPTT, and activated clotting time (if percutaneous coronary intervention performed). In clinical trials, the target activated clotting time (ACT) for patients treated with eptifibatide and undergoing percutaneous coronary intervention was 300–350 sec.32 If concomitant administration of unfractionated heparin is necessary, maintain aPTT at 50–70 sec.33
Notes. (See Abciximab, Notes for vascular access site care after percutaneous coronary intervention.) To minimize bleeding complications, use the following heparin dosages: **continuous IV heparin infusion** (≥70 kg) 5000 units as a bolus and then 1000 units/hr; (<70 kg) 60 units/kg as a bolus and then 12 units/kg/hr.  

**Initial IV heparin boluses during percutaneous coronary intervention** (baseline ACT ≤150 sec) 100 units/kg (up to 10,000 units); (baseline ACT 151–225 sec) 75 units/kg; (baseline ACT 226–299 sec) 50 units/kg; (baseline ACT ≥300 sec) do not administer heparin; then **IV boluses during percutaneous coronary intervention as needed to maintain ACT of 300–350 sec** (ACT 275–299 sec) 25 units/kg; (ACT <275 sec) 50 units/kg.

**Pharmacology.** A heterogeneous, unfractionated group of mucopolysaccharides derived from the mast cells of animal tissues. It binds with antithrombin III, accelerating the rate at which antithrombin III neutralizes activated forms of factors XII, XI, IX, X, VII, and II. It is active in vitro and in vivo.

**Administration and Adult Dosage.** Express dosage in units only; dosage must be individually titrated to desired effect (usually 1.5–2.5 times aPTT).\(^4,8,34\) Weight-based nomograms and computer-assisted dosages of heparin are effective, safe, and superior to “standard care” or empiric approaches.\(^35-37\) **IV for thrombophlebitis or PE (continuous infusion)** 50–100 units/kg initially and then 15–25 units/hr/kg; alternatively, 5000 units initially and then 1000 units/hr; (intermittent) 75–125 units/kg q 4 hr.\(^4,8,34,38\) Duration of therapy for thrombophlebitis or PE is 7–10 days, followed by oral anticoagulation (preferably initiated during the first 24 hr of heparin therapy).\(^34,39\) A 5-day course of heparin has been shown to be as effective as a 10-day course in treating DVT.\(^40\) **SC for thrombophlebitis or PE** 10,000–20,000 units initially (preceded by a 5000-unit IV loading dose) and then 8000–10,000 units q 8 hr or 15,000–20,000 units q 12 hr. **SC for prophylaxis of DVT (low dose)** 5000 units 2 hr before surgery, repeated q 8–12 hr for 5–7 days or until patient is ambulatory.\(^41\) **IV for heparin lock flush** inject sufficient solution (of 10 or 100 units/mL) into injection hub to fill the entire set after each heparin lock use. Some institutions reserve the 100 units/mL solution for flushing triple-lumen central catheters and use NS for all other catheters.

**Special Populations. Pediatric Dosage.** Same as adult dosage in units/kg.

**Geriatric Dosage.** Same as adult dosage.

**Other Conditions.** Patients with PE might require larger heparin doses than patients with thrombophlebitis.\(^38\) (See Administration and Adult Dosage.) Patients with severe renal dysfunction might require lower dosages.\(^5\) There is no good evidence that liver disease appreciably affects dosage requirements.

**Dosage Forms.** **Inj** 1000, 2000, 2500, 5000, 7500, 10,000, 20,000, 40,000 units/mL; 2, 50, 100 units/mL (prediluted); **Heparin Lock Flush** 10, 100 units/mL.

**Patient Instructions.** This drug is potentially harmful when taken with nonprescription or prescription drugs. Consult your physician or pharmacist when considering the use of other medications, in particular aspirin-containing products.

**Pharmacokinetics. Onset and Duration.** Onset immediate after IV administration.
Serum Levels. The relation between heparin serum concentrations and aPTT response can change between reagents and reagent lots. Each laboratory should establish a therapeutic aPTT range corresponding to heparin serum concentrations of 0.2–0.4 unit/mL using protamine titration. Circadian variation in heparin activity can occur, and aPTT response can change during the day at a given infusion rate.

Fate. SC bioavailability is 20–40% and is dose dependent. There is no biotransformation in plasma or liver; transfer and storage in reticuloendothelial cells have been suggested. Vd is 0.058 ± 0.011 L/kg (approximates plasma volume). Cl is dose dependent; Cl can be increased in PE, but this has not been a consistent finding. Shorter half-life has been reported in smokers vs nonsmokers.

\( t_{1/2} \) (Pharmacologic) 90 ± 60 min, dose related; higher doses lead to increased half-life; half-life can decrease in PE, but this has not been a consistent finding.

Adverse Reactions. Bleeding occurs in 3–20% of patients receiving short-term, high-dose therapy. Bleeding risk is increased by 3-fold when the aPTT is 2.0–2.9 and by 8-fold when the aPTT >3.0 times the control. Heparin administration by continuous IV infusion can cause a lower frequency of bleeding complications than intermittent IV administration. Renal dysfunction, liver disease, and other factors (serious cardiac illness, malignancy, age >60 yr, and maximum aPTT >2.2 times control) can increase bleeding risk.

Contraindications. Active bleeding; thrombocytopenia; threatened abortion; subacute bacterial endocarditis; suspected intracranial hemorrhage; regional or lumbar block anesthesia; severe hypotension; shock; and after eye, brain, or spinal cord surgery.

Precautions. Risk factors for hemorrhage are IM injections, trauma, recent surgery, age >60 yr, malignancy, peptic ulcer disease, potential bleeding sites, and acquired or congenital hemostatic defects.

Drug Interactions. Anticoagulants or antiplatelet drugs including aspirin and other NSAIDs can increase risk of bleeding.

Parameters to Monitor. Baseline aPTT, PT/INR, hematocrit, and platelet count. Obtain aPTT (therapeutic range 1.5–2.5 times control) 3 or 4 times (or until therapeutic range is achieved) on day 1 and at least daily thereafter. Monitor platelets and hematocrit every other day and signs of bleeding (melena, hematuria, ecchymoses, hematemesis, epistaxis) daily.

Notes. Heparin-induced thrombocytopenia is a potentially serious, and sometimes fatal, complication of heparin therapy. Lepirudin is approved for management of heparin-induced thrombocytopenia (see Lepirudin monograph). Other
agents including danaparoid, ancrod (Venacil, compassionate use—Knoll), dextran, and argatroban also have been used successfully in these patients.  

LEPIRUDIN  Refudan  
Pharmacology. Lepirudin is a recombinant hirudin analogue that binds to thrombin in a 1:1 stoichiometric complex, thereby inhibiting the thrombogenic activity of thrombin, including clot-bound thrombin. Inhibition of thrombin occurs independently of antithrombin III.  

Administration and Adult Dosage. IV for heparin-induced thrombocytopenia and associated thromboembolic disease 0.4 mg/kg (up to 44 mg) as a bolus and then 0.15 mg/kg/hr (up to 16.5 mg/hr) by continuous infusion for 2–10 days or longer, if necessary. IV in patients being treated concomitantly with thrombolytics 0.2 mg/kg as a bolus and then 0.1 mg/kg/hr by continuous infusion. Adjust dosage based on aPTT as follows: for supratherapeutic aPTT, hold infusion for 2 hr and then reduce infusion rate by 50%; for subtherapeutic aPTT, increase infusion in 20% increments, not to exceed 0.21 mg/kg/hr. 

Special Populations. Other Conditions. In renal insufficiency (Crs >1.5 mg/dL or Clcr <60 mL/min), reduce the IV bolus to 0.2 mg/kg and base the initial IV infusion on renal function: Clcr 45–60 mL/min or Crs 1.6–2 mg/dL, give 0.075 mg/kg/hr; Clcr 30–44 mL/min or Crs 2.1–3 mg/dL, give 0.045 mg/kg/hr; Clcr 15–29 mL/min or Crs 3.1–6 mg/dL, give 0.0225 mg/kg/hr; Clcr <15 mL/min or Crs >6 mg/dL, not recommended. 

Dosage Forms. Inj 50 mg. 

Pharmacokinetics. Fate. About 45% of the administered dose is eliminated in the urine, largely as unchanged drug (35%). Clearance is approximately 25% lower in women than in men and is also reduced about 20% in the elderly. 

\[ t_{\frac{1}{2}} = \alpha \text{ phase 10 min; } \beta \text{ phase 1.3 hr.} \] 

Adverse Reactions. Bleeding complications are the most frequent adverse reactions. Hypersensitivity reactions, primarily allergic skin reactions, occur frequently. 

Contraindications. Hypersensitivity to hirudins. 

Precautions. Use with caution in patients with active internal bleeding, history of recent major bleeding, or known bleeding diathesis; history of CVA or any history of intracranial hemorrhage; history of intracranial neoplasm, AV malformation, or aneurysm; recent puncture of large blood vessel or organ biopsy; recent (within 1 month) major surgery or trauma; severe uncontrolled hypertension; bacterial endocarditis; poor renal function; receiving concomitant antithrombotic therapy including thrombolytics. Up to 40% of patients with heparin-induced thrombocytopenia treated with lepirudin develop antihirudin antibodies, which can increase the anticoagulant effects of lepirudin, necessitating strict monitoring of aPTT values. 

Parameters to Monitor. Monitor aPTT 4 hr after initiating lepirudin therapy, 4 hr after each change in infusion rate, and daily once target aPTT has been achieved. Maintain aPTT approximately 1.5–2.5 times the control aPTT.
Notes. Desirudin (Revasc—Aventis) is a recombinant hirudin that is being studied for use in DVT prevention after hip replacement.

**Pharmacology.** Vitamin K is a required cofactor for the hepatic microsomal enzyme system that carboxylates glutamyl residues in precursor proteins to γ-carboxyglutamyl residues. These proteins are present in vitamin K–dependent clotting factors (II, VII, IX, and X), anticoagulation proteins (proteins C and S), bone (osteocalcin), some plasma proteins (protein Z), and the proteins of several organs (kidney, lung, and testicular tissue).51–53

**Administration and Adult Dosage.** The normal daily nutritional requirement is about 0.03–1.5 µg/kg.52,54,55 The adult RDAs are 70 µg/day for men 19–24 yr and 80 µg/day for men >25 yr and 60 µg/day for women 19–24 yr and 65 µg/day for women >25 yr.50 PO, SC, or IM to reverse bleeding 2.5–10 mg up to 25 mg initially. A single dose of 1–5 mg is usually sufficient to normalize PT during anticoagulant therapy, but in severe bleeding, 20–50 mg might be needed.52,56–58 The initial dose can be repeated, based on PT and clinical response, after 12–48 hr if given PO and 6–8 hr if given parenterally. Use the smallest dosage possible to reverse anticoagulants and obviate possible refractoriness to additional anticoagulant therapy.52,59 IV do not give AquaMephyton intravenously unless it is absolutely essential (e.g., INR >20, serious warfarin overdose or life-threatening bleeding). Give IV in 10 mg doses at infusion rates no greater than 1 mg/min.57 The drug can be diluted in preservative-free dextrose or saline solution just before IV use. PO for antenatal use in pregnant women receiving anticonvulsants 20 mg/day throughout the last 4 weeks of pregnancy.53

**Special Populations. Pediatric Dosage.** RDAs are (<6 months) 5 µg/day; (6 months–1 yr) 10 µg/day; (1–3 yr) 15 µg/day; (4–6 yr) 20 µg/day; (7–10 yr) 30 µg/day; (11–14 yr) 45 µg/day; (15–18 yr) 55 µg/day for females, 65 µg/day for males.56 IM for prophylaxis of hemorrhagic disease of the newborn 0.5–1 mg within 1 hr of birth. SC or IM for treatment of hemorrhagic disease of the newborn 1 mg; more if mother has been receiving an oral anticoagulant.

**Geriatric Dosage.** (>55 yr) RDAs are 65 µg/day for women and 80 µg/day for men.56

**Dosage Forms.** Tab (Mephyton) 5 mg; Inj (AquaMephyton) 2, 10 mg/mL.

**Pharmacokinetics.** Onset and Duration. Reversal of anticoagulant effect is variable among individuals; parenteral onset is often within 6 hr; peak and duration differ across individuals and doses. A 5 mg IV dose usually returns PT to normal in 24–48 hr.50 Large doses can cause prolonged refractoriness to oral anticoagulants.52,59

**Fate.** Absorbed from the GI tract via intestinal lymphatics only in the presence of bile; well absorbed after parenteral administration. Metabolized in the liver to hydroquinone and epoxide forms, which are interconvertible with the quinone.51 Little storage occurs in the body. Without bile, hypoprothrombinemia develops over several weeks.
Adverse Reactions. The drug itself appears to be nontoxic, but severe reactions (eg, flushing, dyspnea, chest pain) and, occasionally, deaths have occurred after IV administration of AquaMephyton, possibly caused by the emulsifying agents.52,55,62 This product should rarely be used IV, and only when other routes of administration are not feasible. A transient flushing sensation, peculiar taste, and pain and swelling at the injection site can occur. Large parenteral doses in neonates have caused hyperbilirubinemia.

Precautions. Temporary refractoriness to oral anticoagulants can occur, especially with large doses of vitamin K. Reversal of anticoagulant activity can restore previous thromboembolic conditions. No effect or worsening of hypoprothrombinemia can occur in severe liver disease, and repeated doses are not warranted if response to the initial dose is unsatisfactory.52,62

Drug Interactions. Mineral oil and cholesterol-binding resins can impair phytonadione absorption.

Parameters to Monitor. Monitor PT before and at intervals after administration of the drug; the interval depends on the route of administration, the condition being treated, and the patient’s status. (See Administration and Adult Dosage.)

Notes. Always protect the drug from light. Phytonadione reverses the effects of oral anticoagulant therapy but has no antagonist activity against heparin.

RETEPLASE

Pharmacology. Reteplase (recombinant plasminogen activator) is a nonglycosylated mutant of wild-type tissue plasminogen activator. In animals, this modification results in less high-affinity fibrin binding, longer half-life, and greater thrombolytic potency than alteplase (rt-PA).

Administration and Adult Dosage. IV for post-MI clot lysis two 10 IU boluses 30 min apart, with adjunctive IV heparin given as a 5000 unit bolus followed by 1000 units/hr (aPTT target 1.5–2.0 times control) for at least 24 hr.

Dosage Forms. Inj 10.8 IU.

Pharmacokinetics. Onset and Duration. Onset of fibrinolytic activity is immediate after IV administration; duration is about 48 hr as assessed by fibrinogen levels.

Fate. Vd is about 6 L. Elimination is primarily by the liver and kidneys.

$\alpha$ phase 14 ± 0.7 min (range 11–19 min); $\beta$ phase 173 ± 33 min.63

Adverse Reactions. (See Alteplase.)

Contraindications. (See Alteplase.)

Precautions. (See Alteplase.)

Notes. In the Reteplase Angiographic Phase II International Dose-finding study (RAPID) open-label MI trial, reteplase achieved more rapid, complete, and sustained thrombolysis than did standard-dose rt-PA, with comparable bleeding risk.64 The RAPID trial did not have sufficient power to detect differences in mortality between the groups. The GUSTO-III trial found reteplase equivalent to accelerated-infusion alteplase in MI for the combined endpoints of death or non-
fatal, disabling stroke. The International Joint Efficacy Comparison of Thrombolytics (INJECT) trial suggested that reteplase mortality rates were similar to those observed with streptokinase.

**STREPTOKINASE**

**Pharmacology.** A bacterial protein derived from group C β-hemolytic streptococci. It acts indirectly by forming a streptokinase-plasminogen activator complex that activates another plasminogen and converts it to the proteolytic enzyme plasmin. Plasmin then hydrolyzes fibrin, fibrinogen, factors II, V, and VIII, complement, and kallikreinogen.

**Administration and Adult Dosage.** IV for post-MI clot lysis 1.5 million IU over 60 min. IV for PE, DVT, arterial thrombosis, or embolism 250,000 IU over 30 min, followed by 100,000 IU/hr for 24–72 hr (72 hr if DVT suspected). Institute heparin therapy. (See Notes and Parameters to Monitor.) For arteriovenous cannula occlusion slowly instill 250,000 IU in 2 mL solution into each occluded limb of cannula; clamp for 2 hr, aspirate contents, and flush with NS. Selective intra-arterial infusion (investigational) 5000 IU/hr for 5–48 hr. (See Notes.)

**Special Populations.** **Pediatric Dosage.** Safety and efficacy not established.

**Geriatric Dosage.** Same as adult dosage.

**Dosage Forms.** Inj 250,000, 750,000, 1.5 million IU.

**Pharmacokinetics.** **Onset and Duration.** Onset of fibrinolytic activity immediately after IV administration; duration 8–24 hr after discontinuation of the infusion.

**Fate.** $V_d$ is 0.08 ± 0.04 L/kg; $Cl$ is 0.1 ± 0.04 L/hr/kg. Clearance results in part from formation of an antigen–antibody complex that remains soluble and is rapidly removed. Local inactivation in the circulation by inhibitor complex formation and proteolysis occurs. It is postulated that the reticuloendothelial system also contributes to clearance.

$\frac{1}{2}_\alpha$ phase averages 18–23 min and is related to antigen–antibody formation; $\beta$ phase averages 83 min and appears to be related to clearance by the reticuloendothelial system.

**Adverse Reactions.** Surface bleeding complications occur frequently and usually follow invasive procedures (eg, venous cutdowns, arterial punctures, and sites of surgical intervention). Severe internal bleeding is reported occasionally, but its prevalence is no greater than with other thrombolytics or standard anticoagulant therapy. Transient hypotension occurs occasionally. In ISIS-3 the rates for definite or possible cerebral bleeding were: streptokinase, 0.2%; rt-PA (duteplase, a 2-chain form of alteplase), 0.5%; anistreplase, 0.7%. Occasional allergic reactions are fever, urticaria, itching, flushing, and musculoskeletal pain. Anaphylactoid reactions occur rarely with preparations now in use.

**Contraindications.** (See Alteplase.)

**Precautions.** (See Alteplase.) Prior exposure to anistreplase or streptokinase within the last 12 months.
**Drug Interactions.** Anticoagulants or antiplatelet drugs can increase risk of bleeding.

**Parameters to Monitor.** For short-term thrombolytic therapy of MI, laboratory monitoring is of little value. For IV continuous infusion, monitor thrombin time, aPTT, or PT to detect activation of the fibrinolytic system, performed 3–4 hr after initiating therapy and q 12 hr throughout treatment. No correlation has been made between clotting test results and likelihood of hemorrhage or efficacy; however, prolongation of the thrombin time to 2–5 times normal control value has been recommended.

**Notes.** In addition to post-MI clot lysis, streptokinase is recommended for treatment of thrombosis involving the axillary–subclavian system, the popliteal vein or deep veins of the thigh and pelvis, and for patients in whom massive pulmonary emboli have caused obstruction of blood flow to one or more lung segments or when clinical shock is present. The risk of stroke and intracranial bleeding appears to be less with streptokinase (by a difference of ≤0.5%) than with other thrombolytic agents. Thrombolytic therapy can help prevent venous valvular damage and the development of venous or pulmonary hypertension. The recommended fixed dosage schedule results in sufficient activation of plasminogen in 95% of patients. However, consider using alteplase in those patients exposed to streptokinase or anistreplase within the last 12 months. The benefits of instituting anticoagulant therapy with heparin after completion of the thrombolytic infusion are not clear. However, it is recommended that heparin infusion (to an aPTT of 1.5–2.0 times control) be given only when there is high risk for systemic or venous thromboembolism (eg, anterior MI, CHF, previous embolus, atrial fibrillation). Heparin is initiated without a bolus 4 hr after the start of streptokinase infusion (or when aPTT is less than twice control) and continued for at least 48 hr.

**Pharmacology.** Tenecteplase is a recombinant tissue plasminogen activator, modified from human tissue plasminogen activator (t-PA). Genetic mutations of human t-PA resulted in greater thrombolytic potency, enhanced fibrin-specificity, decreased systemic activation of plasminogen, resistance to plasminogen activator inhibitor 1, and a longer half-life compared with t-PA.

**Administration and Adult Dosage.** IV bolus for myocardial infarction (<60 kg) 30 mg; (60–69 kg) 35 mg; (70–79 kg) 40 mg; (80–89 kg) 45 mg; (≥90 kg) 50 mg. Give tenecteplase in combination with continuous IV heparin infusion. (See Notes.)

**Dosage Forms.** Inj 50 mg.

**Pharmacokinetics.** Onset and Duration. Rapid onset of thrombolysis occurs after IV administration.

**Fate.** Hepatic metabolism is the primary mode of clearance. Cl is 5.94–7.14 L/hr. $t_{1/2}$ α phase 18–24 min; β phase 90–130 min.

**Adverse Reactions.** (See Alteplase.)

**Contraindications.** (See Alteplase.)
Precautions. (See Alteplase.)

Parameters to Monitor. (See Alteplase.) Monitor aPTT while on concomitant heparin therapy; target aPTT is 50–75 sec.77

Notes. In ASSENT-2, patients also were given concomitant IV heparin therapy as follows: (>67 kg) 4000 units as a bolus and then 800 units/hr by continuous infusion; (>67 kg) 5000 units as a bolus and then 1000 units/hr by continuous infusion. Heparin therapy was continued for 48–72 hr. This study demonstrated comparable patency rates, mortality, intracranial hemorrhage, and stroke compared with front-loaded alteplase.77

**Ticlopidine**

Pharmacology. Ticlopidine is an antiplatelet agent that inhibits most known stimuli (eg, ADP, collagen, epinephrine) for platelet aggregation. It prolongs bleeding time, normalizes shortened platelet survival, suppresses platelet growth factor release, and might block von Willebrand factor and fibrinogen interactions with platelets.78–80

Adult Dosage. PO for thrombotic stroke reduction in patients with stroke or stroke precursors, patients with unstable angina, or those undergoing coronary artery bypass graft or coronary angioplasty 250 mg bid.

Dosage Forms. Tab 250 mg.

Pharmacokinetics. The onset of clinical effect is delayed, with maximum efficacy being achieved in 3–8 days. Approximately 80% of the drug is absorbed orally, with peak serum concentrations occurring in about 2 hr. Ticlopidine undergoes extensive liver metabolism to possibly active metabolites, with only 2% excreted unchanged in urine. Half-life is 4–5 days with repeated dosages.

Adverse Reactions. Diarrhea and rash occur frequently. Minor bleeding such as bruising, petechiae, epistaxis, and hematuria occur occasionally. Severe neutropenia occurs in about 0.8% of patients and mild to moderate neutropenia in about 1.6% of patients during the first 3 months of therapy; neutropenia usually resolves within 3 weeks of discontinuation, although sepsis and death have been reported. Thrombocytopenia, thrombotic thrombocytopenic purpura, and cholestasis occur rarely.80,81 Obtain CBC and differential counts q 2 weeks during the first 3 months of therapy; more frequent monitoring is recommended if the ANC is consistently declining or is less than 30% of the baseline value or if patients demonstrate signs and symptoms of thrombotic thrombocytopenic purpura (weakness, pallor, petechiae, or purpura), dark urine, jaundice, or neurologic changes.

Notes. The Ticlopidine Aspirin Stroke Study trial found a 12% risk reduction in nonfatal stroke or cardiovascular death with ticlopidine compared with aspirin in high-risk (previous TIA or minor stroke) men and women. For secondary stroke prevention, the Canadian American Ticlopidine Study trial found that the risk of stroke, MI, or cardiovascular death was reduced by 23% with ticlopidine over placebo. Ticlopidine also was shown to markedly reduce MI, cardiovascular death, and ECG evidence of ischemia in patients with unstable angina.82,83 Reserve ticlopidine for patients intolerant to aspirin and clopidogrel.
Pharmacology. Tirofiban is a nonpeptide, tyrosine derivative that reversibly binds to and inhibits the platelet glycoprotein IIb/IIIa receptor. Inhibition of the glycoprotein IIb/IIIa receptor prevents fibrinogen from binding, thereby preventing platelet aggregation. Tirofiban inhibits platelet aggregation and prolongs bleeding time in a dose-dependent manner.\(^1\)

Administration and Adult Dosage. IV for unstable angina or non–Q-wave MI (acute coronary syndrome) 0.4 \(\mu\)g/kg/min infusion for 30 min and then 0.1 \(\mu\)g/kg/min by continuous infusion. Continue infusion until patient has clinically stabilized; infusion can be continued for up to 108 hr. Tirofiban can be administered to patients who undergo percutaneous coronary intervention. Should percutaneous coronary intervention be performed during tirofiban therapy, continue infusion for 12–24 hr after completing the procedure. Give tirofiban in combination with continuous IV heparin infusion. (See Notes.)

Special Populations. Other Conditions. (Cl\(_{cr}\) <30 mL/min) reduce maintenance infusion rate by 50%.

Dosage Forms. Inj 50, 250 \(\mu\)g/mL.

Pharmacokinetics. Onset and Duration. Rapid inhibition of platelet function occurs after IV administration. Platelet function recovers soon after discontinuation of the IV infusion; bleeding time and ex vivo platelet aggregation return to near baseline levels within 3–8 hr.\(^1\)

Fate. Renal elimination accounts for 39–69% of the total body clearance. About 65% of a dose is excreted in the urine, largely as unchanged drug; about 25% of an administered dose is excreted in the feces. Cl is 9.12–18.84 L/hr.

\(t_{1/2}\) 1.5–2 hr.\(^1\)

Adverse Reactions. Bleeding complications are the most frequent adverse reactions. Use care to minimize the risk of bleeding by minimizing vascular and other trauma and providing proper care of vascular access sites in patients having percutaneous coronary interventions performed. Thrombocytopenia occurs in <2% of patients and is reversible at discontinuation of the drug.\(^8\)

Contraindications. Active internal bleeding or bleeding diathesis within the previous 30 days; history of CVA within 30 days or any history of intracranial hemorrhage; history of intracranial neoplasm, AV malformation, or aneurysm; thrombocytopenia; recent (within 1 month) major surgery or trauma; history, symptoms, or findings suggestive of aortic dissection; severe uncontrolled hypertension; current or planned use of another parenteral glycoprotein IIb/IIIa inhibitor; acute pericarditis.

Precautions. (See Abciximab.)

Parameters to Monitor. Monitor CBC (including platelet count), prothrombin time, aPTT, and activated clotting time (if percutaneous coronary intervention performed). Maintain aPTT approximately 2 times control aPTT.\(^8\)

Notes. (See Abciximab for vascular access site care after percutaneous coronary intervention.) IV heparin during therapy 5000 units as a bolus and then
1000 units/hr continuous infusion adjusted to maintain aPTT 2 times control. **IV heparin if percutaneous coronary intervention was performed** discontinue IV infusion and give 5000–7500 units of heparin as a bolus and then 1000 units/hr by continuous infusion.84

**UROKINASE**

**Pharmacology.** Urokinase is a proteolytic enzyme produced by renal parenchymal cells that act to directly convert plasminogen to plasmin, with effects similar to those of streptokinase.5

**Administration and Adult Dosage.** **IV for pulmonary emboli** 4400 IU/kg loading dose over 10 min, followed by 4400 IU/kg/hr for 12 hr. Heparin therapy is initiated without a loading dose after discontinuation of the thrombolytic when the thrombin time or other coagulation test no longer exceeds 2 times normal control. **Selective intracoronary infusion** 6000 IU/min for up to 2 hr. **IV for catheter clearance** attach a 1 mL tuberculin syringe filled with 5000 IU reconstituted solution (Open-Cath) and slowly inject an amount equal to the catheter volume; aspirate and repeat q 5 min as necessary. If not successful, allow urokinase to remain in the catheter for 30–60 min before attempting to aspirate. For central venous catheters whose functions have not been restored by the bolus method, a 6- or 12-hr infusion of 40,000 IU/hr (5000 IU/mL at 8 mL/hr) in adults might be useful.85

**Dosage Forms.** **Inj** 5000, 9000, 250,000 IU.

**Pharmacokinetics.** The drug’s half-life is about 10–20 min.

**Adverse Reactions.** Side effects, contraindications, and precautions are similar to those of streptokinase, although allergic reactions occur much less frequently.

**WARFARIN SODIUM**

**Pharmacology.** Warfarin prevents the conversion of vitamin K back to its active form from vitamin K epoxide. This impairs formation of the vitamin K–dependent clotting factors II, VII, IX, and X (prothrombin) and proteins C and S (physiologic anticoagulants). The (S)-warfarin enantiomer is approximately 4-fold more potent an anticoagulant than (R)-warfarin.5,86

**Administration and Adult Dosage.** **PO or IV** 5–7.5 mg/day (range 2–10 mg/day), titrating dosage to an INR of 2.0–3.0 for treatment or prophylaxis of venous thrombosis, PE, systemic embolism, tissue heart valves, valvular heart disease, atrial fibrillation (except patients <60 yr with “lone atrial fibrillation”), and recurrent systemic embolism. Adjust dosage to an INR of 2.5–3.5 for management of mechanical prosthetic valves (upper end of range for caged-ball, tilting-disk, and mitral position valves);87 adding aspirin 100 mg/day offers additional protection but increases the risk of mild bleeding.7,34 For post-MI patients who are at increased risk of systemic or pulmonary embolism, maintain a warfarin dosage that achieves an INR of 2.5–3.5 for up to 3 months. Low-dose warfarin (1 mg/day) without measurable changes in PT/INR begun 3 days before central venous catheter placement and continued while the catheter remains in place is recommended to reduce the risk of axillary–subclavian venous thrombosis.8,88 (See Notes.)
Special Populations. Pediatric Dosage. (<18 yr) safety and efficacy not established. However, when used, dosage is titrated based on INR as in adult dosage.

Geriatric Dosage. Same as adult dosage. (See also Precautions.)

Other Conditions. Large variability in response requires that dosage be carefully individualized to each patient. Patients with liver disease, CHF, hyperthyroidism, or fever might be particularly sensitive to warfarin. Renal failure does not enhance the hypoprothrombinemic response to warfarin, but these patients might have compromised hemostatic mechanisms that predispose to bleeding.

Dosage Forms. Tab 1, 2, 2.5, 3, 4, 5, 6, 7.5, 10 mg; Inj 2 mg.

Patient Instructions. This drug is potentially harmful when taken with nonprescription or prescription drugs. Consult your physician or pharmacist when considering the use of other medications, in particular aspirin-containing products.

Missed Doses. Take this drug at the same time each day. It is important that you not miss any doses. If you do miss a dose, take it as soon as you remember. If it is about time for the next dose, take that dose only. Do not double the dose or take extra.

Pharmacokinetics. Onset and Duration. Peak PT effect is in 36–72 hr; at least 4–6 days of warfarin therapy are required before full therapeutic effect is achieved. Duration after discontinuation depends on resynthesis of vitamin K–dependent clotting factors II, VII, IX, and X (which requires about 4–5 days).

Fate. Completely absorbed orally; well absorbed after small bowel resection; 99 ± 1% is bound to plasma proteins; V₄ (racemic) is 0.14 ± 0.06 L/kg; Cl (racemic) is 0.0027 ± 0.0014 L/hr/kg. It undergoes oxidative P450 enzyme bio-transformation in the liver: (R)-warfarin, CYP1A2; (S)-warfarin, CYP2C subfamily, producing warfarin alcohols, which have minor anticoagulant activity. Less than 2% is excreted unchanged in urine.

t₁/₂. 37 ± 15 hr, unchanged in acute hepatic disease. Enantiomer half-lives: (R)-warfarin 43 ± 14 hr; (S)-warfarin 32 ± 12 hr.

Adverse Reactions. Bleeding (major and minor) occurs frequently (6–29%); fatal or life-threatening hemorrhage has been reported in 1–8% of patients. Risk factors for increased bleeding are PT ratio >2.0, age >60 yr, and other comorbid conditions. Skin necrosis (occurring early in therapy and involving the breast, buttocks, thigh, or penis), purple-toe syndrome (occurring after 3–8 weeks of therapy), and alopecia rarely occur.

Contraindications. Pregnancy; threatened abortion; blood dyscrasias; bleeding tendencies; unsupervised patients with senility, alcoholism, psychosis, or lack of cooperation; anticipated spinal puncture procedure; regional or lumbar anesthesia.

Precautions. Avoid all IM injections because of the risk of hematoma. Several other factors can influence response: diet, travel, and environment. Monitor patients with liver disease, CHF, atrial fibrillation, hyperthyroidism, or fever especially carefully. The elderly have a greater risk of major trauma (eg, hip fractures) and physiologic changes in subcutaneous tissues and joint spaces, which can allow bleeding to expand unchecked.
Drug Interactions. There are many important interactions that have potential clinical importance. Careful monitoring and appropriate dosage adjustment are recommended when any potential interacting drug is added or discontinued. Some agents commonly associated with increased warfarin effect are amiodarone, cimetidine, ciprofloxacin, clofibrate, erythromycin, fluconazole, fluvoxamine, lovastatin, metronidazole, quinidine, and trimethoprim-sulfamethoxazole. Some agents commonly associated with decreased warfarin effect are barbiturates, carbamazepine, cholestyramine, griseofulvin, and rifampin. (See references 99 and 100 for more comprehensive information regarding warfarin drug interactions.)

Parameters to Monitor. Monitor PT/INR daily while hospitalized and then weekly to monthly for therapeutic effect; hematocrit; stool guaiac; urinalysis (for hematuria) for toxicity. Also monitor for ecchymoses, hemoptysis, and epistaxis.

Notes. Loading dose has no therapeutic advantage and might be unsafe because of excessive depression of factor VII. Predictive techniques using small loading doses (eg, 10 mg/day for 2–3 days) were developed with a therapeutic range target much higher than current recommendations. With the current narrow and lower therapeutic target, the predictive error of these techniques might be unacceptable. Phytonadione begins to restore the PT toward normal within 4–8 hr, although large doses can induce subsequent resistance to anticoagulant effect lasting ≥1 week. A small oral dose (eg, 2.5 mg) or small slow IV injection (0.5–1 mg) of phytonadione can be used to bring an elevated PT/INR back into target range without resulting resistance. Treat the first episode of venous thrombosis for 6 weeks in patients with reversible risk factors and 6 months in others. Consider continuing warfarin for an indefinite period in patients with active cancer or recurrent venous thrombosis.

Hematopoietics

EPOETIN ALFA Epogen, Procrit

Pharmacology. Epoetin alfa (erythropoietin) is a recombinant human glycoprotein produced from mammalian cells and stimulates production of RBC. The product contains the identical amino acid sequence and produces the same biologic effects as natural erythropoietin.

Administration and Adult Dosage. IV or SC for dialysis or nondialysis chronic renal failure patients 50–100 units/kg 3 times/week initially, increasing or decreasing by 25 units/kg to maintain a target hematocrit of 30–36%. When the target hematocrit is reached (or when the increase >4% in any 2-week period), reduce the dosage to 25 units/kg 3 times/week. If at any time the hematocrit exceeds 36%, discontinue epoetin until the target hematocrit is achieved and then resume at a lower dosage. Individualize the maintenance dosage to maintain the target hematocrit. IV or SC for zidovudine-treated or HIV-infected patients 100 units/kg 3 times/week for 8 weeks initially, increasing or decreasing by 50–100 units/kg based on response; maximum effective dosage is 300 units/kg 3 times/week. If hematocrit exceeds 40%, discontinue epoetin until the hematocrit returns to 36% and then reduce dosage by 25%; adjust dosage to maintain desired hematocrit.
target. Patients with initial erythropoietin levels >500 units/L are unlikely to respond to epoetin. SC for anemia in chemotherapy-treated cancer patients 150 units/kg 3 times/week for 8 weeks initially, increasing to 300 units/kg 3 times/week if there is an unsatisfactory reduction in transfusion requirement or an unsatisfactory increase in hematocrit. If hematocrit exceeds 40%, discontinue epoetin until the hematocrit returns to 36% and then reduce dosage by 25%; adjust dosage to maintain desired hematocrit target. SC for reduction of allogenic blood transfusion in surgery patients 300 units/kg/day for 10 days before surgery, on day of surgery, and 4 days after surgery; alternatively, 600 units/kg/week 21, 14, and 7 days before surgery and on day of surgery. For use in anemic patients (hemoglobin >10 g/dL and ≤13 g/dL) undergoing noncardiac, nonvascular surgery with an anticipated large blood loss.

Special Populations. Pediatric Dosage. Safety and efficacy not established. SC for anemia of prematurity (preterm neonates) 200 units (140 units/kg) every other day for 10 doses; alternatively, 250 units/kg 3 times/week. SC or IV for anemia of end-stage renal disease (newborn–18 yr) 50 units/kg 3 times/week has been used.

Geriatric Dosage. Same as adult dosage.

Dosage Forms. Inj 2000, 3000, 4000, 10,000, 20,000, 40,000 units/mL.

Pharmacokinetics. Onset and Duration. In response to administration 3 times/week, reticulocyte count increases within 10 days followed by increases in RBC count, hematocrit, and hemoglobin in about 2–6 weeks.

Fate. Not orally bioavailable. Peak serum levels occur 5–24 hr after SC administration. Vd is 0.033–0.055 L/kg; Cl is about 0.00282 L/hr/kg. t¹⁄₂ 9.3 ± 3.3 hr initially; 6.2 ± 1.8 hr during long-term therapy.

Adverse Reactions. Hypertension, headache, tachycardia, nausea, vomiting, clotted vascular access, shortness of breath, hyperkalemia, and diarrhea occur frequently. Seizures occur occasionally; CVA, TIA, and MI occur rarely.

Contraindications. Uncontrolled hypertension. Hypersensitivity to mammalian cell-derived products or albumin.

Precautions. Pregnancy. Use cautiously with a known history of seizure or underlying hematologic diseases such as sickle cell anemia, myelodysplastic syndromes, and hypercoagulable states.

Drug Interactions. None known.

Parameters to Monitor. Evaluate iron stores before and during therapy. Supplemental iron might be required to maintain a transferrin saturation of at least 20% and ferritin levels of at least 100 µg/L. Determine hematocrit twice a week for 2–6 weeks or until stabilized in the target range; monitor at regular intervals thereafter. Monitor CBC with differential platelet count, BUN, Cr, serum uric acid, serum phosphorus, and serum potassium at regular intervals.

Notes. Darbepoetin (Aranesp—Amgen) is a highly glycosylated form of erythropoietin that is absorbed slowly and can be given once weekly or every 2 weeks. The weekly dose in µg/kg equals the total weekly dosage of epoetin alfa in IV/week divided by 200.
FERROUS SALTS

Pharmacology. Ferrous salts are soluble forms of iron, an essential nutrient that functions primarily as the oxygen-binding core of heme in red blood cells (as hemoglobin) and muscles (as myoglobin) and in the respiratory enzyme cytochrome C.

Administration and Adult Dosage. PO as a dietary supplement RDAs are 10 mg/day for men (19–51 yr) and 15 mg/day for women (19–51 yr).\textsuperscript{56} PO for treatment of iron deficiency 2–3 mg/kg/day of elemental iron in divided doses. (See Ferrous Salts Comparison Chart for usual dosage ranges for individual salts.) Dose-related adverse effects can be decreased by using suboptimal dosages, increasing the daily dosage gradually, or administering with a small amount of food (although this latter method reduces absorption). After hemoglobin is normalized, continue oral therapy for 3–6 months to replenish iron stores.

Special Populations. Pediatric Dosage. PO for prophylaxis RDA (infants) 6 mg/day; (1–10 yr) 10 mg/day; (11–18 yr, males) 12 mg/day; (11–18 yr, females) 15 mg/day.\textsuperscript{56} PO for treatment (infants) 10–25 mg of elemental iron in 3–4 divided doses; (6 months–2 yr) up to 6 mg/kg/day of elemental iron in 3–4 divided doses; (2–12 yr) 3 mg/kg/day of elemental iron in 3–4 divided doses.

Geriatric Dosage. Same as adult dosage, except dosage in women older than 51 yr is 10 mg/day of elemental iron.

Other Conditions. Iron requirement during pregnancy is approximately twice that of the normal, nonpregnant woman because of an expanding blood volume and the demands of the fetus and placenta. The RDA in pregnancy is 30 mg/day, and a prophylactic dose of 15–30 mg/day of elemental iron during the second and third trimesters has been recommended to prevent depletion of maternal iron stores. Iron-deficient patients might need higher doses.

Dosage Forms. (See Ferrous Salts Comparison Chart.)

Patient Instructions. Take this drug with a full glass of water on an empty stomach (1 hour before or 2 hours after meals) for best absorption. Take liquid preparations in water or juice and drink with a straw to minimize tooth staining. If gastric distress or nausea occurs, a small quantity of food can be taken with the drug but do not take with antacids because absorption is decreased. Iron preparations can cause constipation and black stools. Keep all iron products out of the reach of children.

Missed Doses. Take this drug at regular intervals. If you miss a dose, take it as soon as you remember. If it is about time for the next dose, take that dose only. Do not double the dose or take extra.

Pharmacokinetics. Onset and Duration. Responses to equivalent amounts of oral or parenteral therapy are essentially the same. Reticulocytes increase within 4–7 days and reach a peak on about the 10th day. An increase in hemoglobin of at least 2 g/dL and a 6% increase in hematocrit should occur in about 3–4 weeks. Three to 6 months of therapy are generally required for restoration of iron stores.\textsuperscript{110,111}

Serum Levels. Normal levels are 65–170 µg/dL (12–30 µmol/L) in men, 50–170 µg/dL (9–30 µmol/L) in women, and 50–120 µg/dL (9–21 µmol/L) in children. A decrease in the transferrin saturation (serum iron ÷ total iron-binding

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capacity × 100) indicates preanemic iron deficiency. A transferrin saturation <16% or plasma ferritin concentration <12 µg/L indicates probable iron deficiency. In overdosage, toxicity can occur at iron levels >350 µg/dL (63 µmol/L). Chelation therapy is indicated at these levels, especially if the patient is symptomatic.

**Fate.** Iron is absorbed primarily from the duodenum at a rate that depends on the amount of iron in storage sites. About 10% of dietary iron is absorbed in normal subjects, 20% in iron-deficient patients, and as much as 70% of medicinal iron is absorbed during marked iron deficiency or increased erythropoiesis. In the plasma, iron is oxidized to the ferric state, combined with transferrin, and used or stored as ferritin (mostly in the reticuloendothelial system and hepatocytes). The average loss in the healthy adult male is about 1 mg/day. GI loss of extravasated red cells, iron in bile, and exfoliated mucosal cells accounts for two-thirds of this iron. The other one-third is lost in the skin and urine. Menstruating women have an additional loss of about 0.5 mg/day.

**Adverse Reactions.** Side effects are related primarily to the dose of elemental iron. Frequent GI irritation, constipation, and stained teeth (liquid preparations only—dilute and use a drinking straw). An increased risk of cancer associated with excessive iron stores has been reported.

**Contraindications.** Hemochromatosis; hemosiderosis; hemolytic anemias in which no true iron deficiency exists.

**Precautions.** Use with caution in patients with peptic ulcer, regional enteritis, or ulcerative colitis. Serious acute poisoning (which can be fatal) occurs frequently in children: doses as low as 20 mg/kg of elemental iron can cause toxicity; 40 mg/kg is considered serious; and >60 mg/kg is potentially lethal.

**Drug Interactions.** Food, calcium carbonate, sodium bicarbonate, and possibly magnesium trisilicate can reduce iron absorption. Vitamin E can reduce utilization of iron in iron-deficiency anemia. Iron salts can reduce oral absorption of carbidopa/levodopa, methyldopa, penicillamine, quinolones, tetracyclines, and thyroid hormones.

**Parameters to Monitor.** Periodic reticulocyte count, hemoglobin, and hematocrit. (See Onset and Duration.)

**Notes.** Ferrous salts are used to prevent and treat iron-deficiency anemias. Such anemias occur most frequently with exceptional blood losses (eg, pathologic bleeding, menstruation) and during periods of rapid growth (eg, infancy, adolescence, pregnancy). Iron is ineffective in hemoglobin disturbances not caused by iron deficiency. Concurrent administration of high doses of vitamin C can enhance absorption (particularly when given with SR formulations), but cost/benefit might not warrant its use. Wide variations in dissolution and absorption exist among SR and EC products, and the frequency of adverse effects, although negligible, probably reflects the small amount of ionic iron available for absorption because of transport of the iron past the duodenum and proximal jejunum.
<table>
<thead>
<tr>
<th>DRUG</th>
<th>SOLID DOSAGE FORMS&lt;sup&gt;a&lt;/sup&gt;</th>
<th>ADULT DOSAGE (CAP OR TAB/DAY)</th>
<th>ELEMENTAL IRON/CAP OR TAB (%)</th>
<th>(mg Fe)</th>
<th>OTHER DOSAGE FORMS&lt;sup&gt;a&lt;/sup&gt;</th>
</tr>
</thead>
<tbody>
<tr>
<td>Carbonyl Iron</td>
<td>Cap 50 mg iron.</td>
<td>3</td>
<td>100</td>
<td>50</td>
<td>Susp 12 mg/mL iron.</td>
</tr>
<tr>
<td>Ferrous Fumarate</td>
<td>Chew Tab 100 mg</td>
<td>1–4</td>
<td>33</td>
<td>33</td>
<td>Drp 75 mg/mL.</td>
</tr>
<tr>
<td></td>
<td>Tab 63, 200, 324, 325, 350 mg.</td>
<td>1–4</td>
<td>33</td>
<td>20, 66</td>
<td>Susp 20 mg/mL.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>1–2</td>
<td>33</td>
<td>106, 106, 115</td>
<td></td>
</tr>
<tr>
<td>Ferrous Gluconate</td>
<td>Tab 240, 325 mg.</td>
<td>3–6</td>
<td>11</td>
<td>27, 36</td>
<td>Elxr 60 mg/mL.</td>
</tr>
<tr>
<td>Ferrous Sulfate</td>
<td>SR Tab 160 mg</td>
<td>1–2</td>
<td>30</td>
<td>50</td>
<td>Drp 125 mg/mL.</td>
</tr>
<tr>
<td>Exsiccated</td>
<td>Tab 187, 200 mg.</td>
<td>3–4</td>
<td>30</td>
<td>60, 65</td>
<td>Elxr 44 mg/mL.</td>
</tr>
<tr>
<td>Ferrous Sulfate</td>
<td>SR Cap/Tab various</td>
<td>—</td>
<td>20</td>
<td>—</td>
<td>Syrup 18 mg/mL.</td>
</tr>
<tr>
<td>Hydrous</td>
<td>Cap 250 mg</td>
<td>3</td>
<td>20</td>
<td>39</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Tab 195, 300, 324 mg.</td>
<td>3–6</td>
<td>20</td>
<td>50</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>3</td>
<td>20</td>
<td>60, 65</td>
<td></td>
</tr>
<tr>
<td>Polysaccharide-Iron</td>
<td>Cap 150 mg iron</td>
<td>1–2</td>
<td>—</td>
<td>150</td>
<td>Elxr 20 mg/mL iron.</td>
</tr>
<tr>
<td>Complex</td>
<td>Tab 50 mg iron.</td>
<td>2–4</td>
<td>—</td>
<td>50</td>
<td></td>
</tr>
</tbody>
</table>

<sup>a</sup>Doses listed represent total iron salt, not elemental iron, except for carbonyl iron and polysaccharide-iron complex.
Filgrastim is an Escherichia coli–derived (nonglycosylated) recombinant human granulocyte colony-stimulating factor (G-CSF). G-CSF is one of many glycoprotein hormones that regulate the proliferation and differentiation of hematopoietic progenitor cells and the function of mature blood cells. Specifically, G-CSF promotes proliferation and maturation and enhances the function and migration of neutrophil granulocytes. G-CSF also promotes pre–B-cell activation and growth and acts in synergy with interleukin-3 to support megakaryocyte and platelet production.

Administration and Adult Dosage. SC or IV for myelosuppressive cancer chemotherapy 5 µg/kg/day as a single injection. The drug is usually discontinued once the postnadir ANC reaches 1500–2000/µL. Based on severity of ANC nadir, dosage can be increased in 5 µg/kg/day increments for each chemotherapy cycle.

SC continuous infusion for chemotherapy-induced febrile neutropenia 12 µg/kg/day beginning within 12 hr of empiric antibiotic therapy and continued until ANC is >5000/µL and the patient is afebrile for 4 days. IV or SC for bone marrow transplant patients 10 µg/kg/day infused IV over 4 or 24 hr or as a continuous SC infusion and then decreasing to 5 µg/kg/day when ANC is >1000/µL for 3 consecutive days. Discontinue therapy if the ANC remains >1000/µL for 3 more consecutive days; resume at a dosage of 5 µg/kg/day when ANC becomes <1000/µL. SC for severe chronic neutropenia (congenital) 6 µg/kg bid; (idiopathic or cyclic) 5 µg/kg/day. Target ANC range is 1500–10,000/µL; decrease dosage if ANC is persistently >10,000/µL. SC with erythropoietin to decrease hematologic toxicity from zidovudine 3.6 µg/kg/day initially, increasing or decreasing weekly by 1 µg/kg/day to maintain a target ANC of 1500–5000/µL.

Special Populations. Pediatric Dosage. Safety and efficacy not established. IV or SC adult dosages in µg/kg are well tolerated.

Geriatric Dosage. Same as adult dosage.

Dosage Forms. Inj 300, 600 µg/mL.

Patient Instructions. Your pharmacist or physician should instruct you on proper dosage, administration, and disposal. Store vials in the refrigerator but do not freeze. Vials are designed for single use only; discard any unused portion. Bring vial to room temperature before administration; do not shake.

Pharmacokinetics. Onset and Duration. Increase in neutrophilic band forms occurs within about 60 min after administration. After therapy is discontinued, neutrophil counts return to baseline values in about 4 days.

Fate. Not orally bioavailable. Vd is about 0.15 L/kg; Cl is 0.03–0.042 L/hr/kg. t1/2 3.5–3.85 hr.

Adverse Reactions. Mild to moderate bone pain responsive to non-narcotic analgesics is reported frequently. Transient decreases in blood pressure occur occasionally. During long-term therapy, splenomegaly occurs frequently; occasional exacerbation of skin disorders, alopecia, hematuria, proteinuria, thrombocyto-
penia, and osteoporosis also are reported. Other adverse effects occur during administration of filgrastim that are likely the consequence of the underlying malignancy or cytotoxic chemotherapy. Acute reactions to sargramostim (eg, febrile episodes, flushing, hypotension, tachycardia, and hypoxia) appear to be more common than with filgrastim.104,115,117,118

**Contraindications.** History of hypersensitivity to *E. coli*–derived proteins. Do not use 24 hr before and 24 hr after administration of cytotoxic chemotherapy.

**Precautions.** Use with caution in any malignancy with myeloid characteristics because of the possibility of tumor growth. The efficacy of filgrastim has not been established in patients receiving nitrosoureas, mitomycin, fluorouracil, or cytarabine.

**Drug Interactions.** None known.

**Parameters to Monitor.** Perform CBC and platelet counts before chemotherapy and twice a week during filgrastim therapy. Regular monitoring of WBC count at the time of recovery from the postchemotherapy nadir is recommended to avoid excessive leukocytosis.

**Notes.** Other potential uses for filgrastim include AIDS-related neutropenia, myelodysplastic syndromes, and drug-induced neutropenia or aplastic anemia. Further clinical trials are needed to prove that use of filgrastim for these and other indications is beneficial, safe, and cost effective.

## Iron Dextran

**DexFerrum, InFeD, Various**

**Pharmacology.** (See Ferrous Salts.) The overall response to parenteral iron is no more rapid or complete than the response to orally administered iron, so iron dextran is indicated only when oral iron therapy is determined to be ineffective or impossible.

**Administration and Adult Dosage.** The total cumulative amount required for restoration of hemoglobin (Hb) in g/dL and body stores of iron can be approximated using lean body weight (LBW) in kg (or actual body weight if less than LBW) from the formula:

\[
\text{Total mg Iron} = (0.0442 \times [\text{Desired Hb} - \text{Observed Hb}] \times \text{LBW} + [0.26 \times \text{LBW}]) \times 50
\]

To calculate dose in mL, divide the result by 50. Usual Hb target for adults is 14.8 g/dL. The dose of iron required secondary to blood loss can be estimated from the formula:

\[
\text{Total mg Iron} = \text{Blood Loss (mL)} \times \text{Hematocrit (observed, as decimal fraction)}
\]

**Deep IM** (in upper outer quadrant of buttock only with the Z-track technique) 25 mg (0.5 mL) test dose the first day and then, if no adverse reaction occurs, administer a maximum daily dose of 100 mg (2 mL) until the total calculated amount is reached. **Slow IV** test dose of 25 mg (0.5 mL) over at least 30 sec the first day; if no adverse reaction occurs after at least 1 hr, proceed (until the total calculated amount is reached) by daily increments over 2–3 days, to a maximum
dose of 100 mg/day at a rate not to exceed 50 mg/min. IV in erythropoietin-treated dialysis patients 100–200 mg/week after dialysis. Total dose IV infusion is an off-label use and is discouraged by the FDA but is widely used. The total calculated dose of iron dextran is diluted in 500 mL of NS (dextrose solutions increase local phlebitis) and infused at a rate of 6 mg/min after a 30 mL test dose is delivered over 2 min.

Special Populations. Pediatric Dosage. (<4 months) safety and efficacy not established; (5–15 kg) total cumulative amount required for restoration of Hb (in g/dL) and body stores of iron can be estimated using body weight (W) in kg from the formula:

\[
\text{Total mg Iron} = (0.0442 \times [\text{Desired Hb} - \text{Observed Hb}] \times W + [0.26 \times W]) \times 50
\]

To calculate dose in mL, divide the result by 50. Usual Hb target for children ≤15 kg is 12 g/dL. Maximum daily dose is (infants <5 kg) 25 mg (0.5 mL), (children <10 kg) 50 mg (1 mL), (children >15 kg) same as adult dosage.

Geriatric Dosage. Same as adult dosage.

Dosage Forms. Inj 50 mg elemental iron/mL.

Pharmacokinetics. Onset and Duration. Hematologic response is the same as with oral therapy, although total body stores of iron are replaced when the above dosage regimens are used.

Serum Levels. (See Ferrous Salts.)

Fate. After IV administration, the inert complex is gradually cleared from the plasma by the reticuloendothelial cells of the liver, spleen, and bone marrow. With doses >500 mg, the rate of uptake is 10–20 mg/hr. Iron dextran is then dissociated and released as free ferric iron (at a rate controlled by the serum iron level), which combines with transferrin and is incorporated into hemoglobin within the bone marrow. Although all iron is eventually released in this manner, many months often are required for this process to be completed.

Adverse Reactions. Hypotension and peripheral vascular flushing occur with too-rapid IV administration. Mild, transient reactions including flushing, fever, myalgia, arthralgia, and lymphadenopathy usually occur only occasionally but have occurred in 80–90% of patients with active rheumatoid arthritis or active SLE. Immediate anaphylactoid reactions, which can be life-threatening, occur in 0.1–0.6% of patients. A predictive test for predisposition to anaphylaxis is not available. IM administration has been associated with variable degrees of soreness, sterile abscess formation, tissue staining, and sarcoma formation. Total dose infusion appears to be tolerated as well as divided doses.

Contraindications. Anemias other than iron-deficiency anemia; hemochromatosis; hemosiderosis; SC administration.

Precautions. Pregnancy. Use with extreme caution with serious liver impairment. Patients with rheumatoid arthritis might have an acute exacerbation or reactivation of joint pain and swelling after administration. History of allergies and/or asthma. Because of the potential for anaphylactoid reactions, have epinephrine, diphenhy-
dramine, and methylprednisolone immediately available during iron dextran administration. Use parenteral iron only in patients in whom an iron-deficient state has been clearly established and who are not amenable to oral therapy.

**Drug Interactions.** None known.

**Parameters to Monitor.** (See Ferrous Salts.)

**Notes.** Sodium ferric gluconate complex (Ferrlecit) is an injectable iron product containing 12.5 mg/mL of elemental iron. It is indicated for iron-deficiency anemia in chronic hemodialysis patients receiving epoetin alfa. A test dose is not required, but a test dose of 2 mL in 50 mL of NS given IV over 1 hr has been used. The standard dose is 10 mL/100 mL NS infused over 1 hr. Most patients require a total dosage of 1 g of elemental iron in 8 doses on sequential dialysis sessions to replete iron stores. It might be better tolerated than iron dextran, but it is much more expensive. It is a good alternative in patients intolerant to iron dextran.

**Iron sucrose** (Venofer) is an injectable iron product containing 20 mg/mL of elemental iron. It is indicated for iron-deficiency anemia in chronic hemodialysis patients receiving epoetin alfa. A test dose is not required, but a test dose of 2.5 mL in 50 mL of NS over 3–10 min has been used. The drug can be given by direct IV injection at a rate of 1 mL (20 mg of iron) per minute or by slow infusion by diluting one vial (100 mg iron) in no more than 100 mL of NS and infusing it over at least 15 min. The recommended dosage is 100 mg of iron (1 vial) no more than 3 times per week to a total of 1 g in 10 doses. This regimen can be repeated if necessary.

**SARGRAMOSTIM**

**Pharmacology.** Sargramostim is a yeast-derived (glycosylated) recombinant human granulocyte–macrophage colony-stimulating factor (GM-CSF). GM-CSF is one of many glycoprotein hormones that regulate the proliferation and differentiation of hematopoietic progenitor cells and the function of mature blood cells. Specifically, GM-CSF promotes proliferation, maturation, and function of neutrophils, eosinophils, monocytes, and macrophages. GM-CSF also stimulates production of cytokines such as interleukin-1 and tumor necrosis factor.

**Administration and Adult Dosage.** IV after autologous bone marrow infusion 250 µg/m²/day given as a 2-hr infusion beginning 2–4 hr after the autologous bone marrow infusion. Give the first dose no sooner than 24 hr after the last chemotherapy dose or 12 hr after the last dose of radiotherapy. Continue sargramostim until the ANC is >1500/µL for 3 consecutive days. IV for bone marrow transplantation failure or delay in engraftment 250 µg/m²/day for 14 days as a 2-hr infusion; repeat in 7 days if engraftment has not occurred. If there is no improvement, a third course with 500 µg/m²/day given for 14 days can be tried. IV for induction chemotherapy in acute myelogenous leukemia 250 µg/m²/day over 4 hr starting 4 days after completion of chemotherapy if bone marrow is hypoplastic (<5% blasts). Continue until ANC is >1500/µL for 3 consecutive days or at most 42 days. Discontinue or reduce dosage by 50% if ANC is >20,000/µL. Discontinue if leukemic regrowth occurs. SC for AIDS patients receiving ganciclovir 1–15 µg/kg/day has been used investigationaly.
Dosage Forms. Inj 250, 500 µg.

Adverse Reactions. Acute reactions (eg, febrile episodes, flushing, hypotension, tachycardia, and hypoxia) appear to be more common than with filgrastim. Other adverse reactions that occur frequently with sargramostim are bone pain, lethargy, rash, and fluid retention.

Parameters to Monitor. Obtain CBC with differential twice weekly. In patients with renal or hepatic insufficiency, monitor renal and hepatic functions q 2 weeks.

Notes. Sargramostim is indicated for myeloid reconstitution after autologous bone marrow transplantation. It has also been used with some success to maintain normal neutrophil counts in AIDS patients receiving ganciclovir. Other potential uses for sargramostim are AIDS-related neutropenia, myelodysplastic syndromes, and congenital, chronic, or drug-induced neutropenia, and aplastic anemia. Controlled clinical trials are needed to prove that use for these and other indications is beneficial, safe, and cost effective. Clinical and laboratory evidence suggest that sargramostim enhances the effect of zidovudine against HIV.104,115,118,123–125

REFERENCES


Adrenal Hormones

Class Instructions. Corticosteroids. (Systemic use) these drugs may be taken with food, milk, or an antacid to minimize stomach upset. Take single daily doses or alternate-day doses in the morning before 9 AM. Take multiple daily doses at evenly spaced intervals during the day. Report unusual weight gain, lower extremity swelling, muscle weakness, black tarry stools, vomiting of blood, facial swelling, menstrual irregularities, prolonged sore throat, fever, cold, infection, serious injury, fatigue, anorexia, nausea, vomiting, diarrhea, weight loss, dizziness, or low blood sugar. Consult your physician during periods of increased stress. If you are diabetic, you may have increased requirements for insulin or oral hypoglycemics. Carry appropriate identification if you are taking long-term corticosteroid therapy. Avoid immunizations with live vaccines.

Missed Doses. If a dose is missed and the proper schedule is every other day, take it as soon as possible and resume the schedule unless it is past noon. In that case, wait until the next morning and resume every-other-day administration. If the proper schedule is once a day, take the dose as soon as possible. If you do not remember until the next day, do not double that day’s dose; skip the missed dose. If the proper schedule is several times a day, take the dose as soon as possible and resume the normal schedule. If you do not remember until the next dose is due, then take the regular and missed doses and resume the normal dosage schedule.

Pharmacology. Cosyntropin is a synthetic polypeptide containing the first 24 of the 39 amino acids of natural corticotropin (adrenocorticotropic hormone; ACTH) and retaining the full activity of corticotropin with decreased antigenicity. Cosyntropin 250 μg is pharmacologically equivalent to corticotropin 25 units. Cosyntropin stimulates the adrenal cortex to produce and secrete gluco- and mineralocorticoids and androgens similar to corticotropin. Cosyntropin is used as a diagnostic agent to detect adrenocortical insufficiency but can be used therapeutically as a substitute for corticotropin.

Adult Dosage. IM or IV for diagnostic use hold all exogenous corticosteroids (except dexamethasone) on the test day because of assay cross-reactivity and, if the patient is not taking spironolactone or an estrogen, a baseline cortisol level (which should exceed 5 μg/dL) is drawn in the morning just before the dose. Then, cosyntropin 250 μg in 1 mL of NS is given IM, or 250 μg in 2–5 mL NS is given IV push over 2 min. Normal cortisol levels are >18 μg/dL (500 nmol/L)
30 min after the injection and \( \geq 7 \mu g/dL \) (190 nmol/L) above baseline. If the cortisol level is drawn 60 min postadministration, then an approximate doubling of the baseline cortisol value indicates a normal response. Alternatively, give an infusion of 250 \( \mu g \) in D5W or NS over 6 hr in the morning, with serum cortisol levels drawn before and after. The second cortisol level should be \( >18 \mu g/dL \) (500 nmol/L) and \( \geq 7 \mu g/dL \) (190 nmol/L) above baseline. **IV for therapeutic use**

- **Pediatric Dosage.** IM or IV as a diagnostic agent (≤2 yr) 125 \( \mu g \) given as above.
- **Geriatric Dosage.** Same as adult dosage.

**Dosage Forms.** **Inj** 250 \( \mu g \).

**Adverse Reactions.** (Diagnostic use) rare reports of hypersensitivity. (Therapeutic use) salt and water retention, and virilization occur.

**Pharmacology.** Dexamethasone is a potent, long-acting glucocorticoid lacking sodium-retaining activity with low to moderate doses. (See Prednisone Pharmacology and the Oral Corticosteroids Comparison Chart.)

**Administration and Adult Dosage.** Total daily dosage is variable depending on the clinical disorder and patient response. Not recommended for alternate-day administration because of prolonged duration of activity. **PO, IM, or IV for acute, self-limited allergic disorders or exacerbation of chronic allergic disorders** 4–8 mg on the first day in one dose, then taper over 5 days; give tapering dosage in 2 divided doses for 3 days, once daily for 2 days, then discontinue. **IV for cerebral edema** 10 mg (as the sodium phosphate) initially, followed by 4 mg IM or IV q 6 hr for several days until maximal response occurs; then decrease the dosage over 5–7 days and discontinue. **PO, IM, or IV for palliative management of recurrent or inoperable brain tumors** 2 mg bid-tid. **PO or IV as an antiemetic with cancer chemotherapy** (usually in combination with other antiemetics) 10–20 mg immediately before therapy; optionally, up to 40 mg may be given after chemotherapy.** PO as the dexamethasone suppression test to screen for Cushing’s disease** 1 mg at 11 PM; a measured serum cortisol at 8 AM the next morning <5 \( \mu g/dL \) (140 nmol/L) indicates a normal response. Alternatively, **PO 0.5 mg q 6 hr for 48 hr (8 doses), with a 24-hr urine collected for 17-hydroxycorticosteroids (17-OHCs) during the second 24-hr period. A normal response is \( \leq 2.5 \text{mg (6.9 \mu mol)} \) of 17-OHCs during the second 24-hr period.** **PO as a Cushing’s syndrome test to distinguish pituitary origin from other causes** 2 mg q 6 hr for 48 hr (8 doses). A normal response is a 24-hr urine concentration of <2.5 mg (6.9 \( \mu \text{mol}) \) of 17-OHCs. **IM (aqueous) in the mother for antenatal prevention of neonatal distress syndrome** starting 24 hr or more before premature delivery, give 6 mg q 12 hr for 4 doses; however, betamethasone may be the preferred agent. **IV for septic shock** not recommended because of a lack of efficacy and a possible increase in mortality. **IM (depot) for prolonged systemic effect** 8–16 mg q 1–3 weeks. (See also Inhaled Corticosteroids Comparison Chart in the Respiratory Drugs section.)
**Special Populations. Pediatric Dosage.** PO, IM, or IV for airway edema 0.25–0.5 mg/kg/dose q 6 hr prn for croup or beginning 24 hr before planned extubation, then for 4–6 doses; **PO, IM, or IV as an anti-inflammatory or immunosuppressive** 0.03–0.15 mg/kg/day divided q 6–12 hr; **PO, IM, or IV for cerebral edema** 1.5 mg/kg once, then 1.5 mg/kg/day divided q 4–6 hr for 5 days, then taper over the next 5 days and discontinue. IV to prevent hearing loss and other neurologic sequelae in *Haemophilus influenzae* bacterial meningitis (>2 months) 0.15 mg/kg q 6 hr, beginning no sooner than 20 min before (or with) the first dose of antibiotics and continued for 4 days.8–10

**Geriatric Dosage.** Consider using a lower dosage for decreased body size.

**Patient Instructions.** (See Class Instructions: Corticosteroids.)

**Dosage Forms.** Elxr 0.1 mg/mL; Soln 0.1, 1 mg/mL; Tab 0.25, 0.5, 0.75, 1, 1.5, 2, 4, 6 mg; Inj 4, 10, 20, 24 mg/mL (as sodium phosphate); Depot Inj 8, 16 mg/mL (as acetate).

**Pharmacokinetics. Onset and Duration.** (See Oral Corticosteroids Comparison Chart.)

**Serum Levels.** Serum concentration is not directly correlated with therapeutic effect.11

**Fate.** After oral administration, 78 ± 14% is absorbed; 68% is plasma protein bound. Vd is 0.82 ± 0.22 L/kg. The drug is eliminated primarily by hepatic metabolism, with about 2.6 ± 0.6% excreted unchanged in urine.2,12,13

\[ t_{1/2} \] (Males) 3.5 ± 0.87 hr; (females) 2.4 ± 0.16 hr.2,12,13

**Adverse Reactions.** (See Prednisone Adverse Reactions.) Perineal itching or burning can occur after IV administration.14

**Contraindications.** Systemic fungal infections (except as maintenance therapy in adrenal insufficiency); administration of live virus vaccines to patients receiving an immunosuppressive dosage of dexamethasone; IM use in idiopathic thrombocytopenic purpura.

**Precautions.** (See Prednisone.)

**Parameters to Monitor.** (See Prednisone.)

**Notes.** The dexamethasone suppression test for diagnosis of depression is of unproven value.15,16 Variations of the dexamethasone suppression test (for Cushing’s disease screening) have been used.5 Betamethasone appears to be the corticosteroid of choice for prevention of neonatal distress; the maternal dosage is 12 mg IM q 24 hr for 2 doses (usually as Celestone Soluspan).6,7

**Pharmacology.** Methylprednisolone sodium succinate is an injectable glucocorticoid that has about 1.25 times greater anti-inflammatory potency than prednisone or prednisolone and a similar duration of biologic activity. (See Prednisone Pharmacology, Oral Corticosteroids Comparison Chart.) It is commonly used when oral therapy is not possible and in situations in which large parenteral doses are necessary.
Administration and Adult Dosage. IV initial dosage 10–40 mg given over one to several minutes; dosages up to 30 mg/kg q 4–6 hr (high-dose therapy) for up to 48–72 hr are used for severe, acute conditions. Infuse large doses (≥500 mg) slowly (eg, over 30–60 min) because arrhythmias and sudden death have occurred with rapid infusions.17 IV for acute spinal cord injury 30 mg/kg over 15 min, begun within 8 hr of injury, followed 45 min later by a continuous IV infusion of 5.4 mg/kg/hr for 23 hr.18,19 (See Notes.)

Special Populations. Pediatric Dosage. IV as an anti-inflammatory or immunosuppressive 0.16–0.8 mg/kg/day in 2–4 divided doses.9

Geriatric Dosage. In the elderly, consider using lower dosages for decreased body size.

Dosage Forms. Inj 40, 125, 500 mg, 1, 2 g; Depot Inj (as acetate) 20, 40, 80 mg/mL. (See also Oral Corticosteroids Comparison Chart.)

Pharmacokinetics. Plasma protein binding is 78 ± 3%. Vd is 1.2 ± 0.2 L/kg; Cl is 0.37 ± 0.054 L/hr/kg. The drug is extensively metabolized, with 4.9 ± 2.3% excreted unchanged in urine. The serum half-life is 2.2 ± 0.5 hr and is not affected by renal function.2,13

Adverse Reactions. Side effects are similar to prednisone in equivalent dosages.

Drug Interactions. Ketoconazole and some macrolide antibiotics reduce methylprednisolone elimination, possibly leading to excessive corticosteroid effect. (See also Prednisone Drug Interactions.)

Notes. Evidence of efficacy in improving the outcome of septic shock is lacking, and, because of increased mortality in some patient groups, the use of methylprednisolone in septic shock is not recommended.8 Small, early studies in patients with acute spinal cord injury treated with high-dose methylprednisolone within 8 hr appeared to show improved neurologic recovery,2,18,20 although the bulk of evidence indicates no such benefit.21

PREDNISONE

Pharmacology. Prednisone is a synthetic glucocorticoid with less sodium-retaining activity than hydrocortisone. Prednisone is inactive until converted into prednisolone. At the cellular level, glucocorticoids appear to act by controlling the rate of protein synthesis mediated through gene transcription. Clinically, these drugs are used primarily for their anti-inflammatory and immunosuppressant effects.22

Administration and Adult Dosage. Total daily dosage is variable and must be individualized depending on the clinical disorder and patient response.2,5 Daily divided high-dose therapy for initial control of more severe disease states may be necessary until satisfactory control is obtained, usually 4–10 days for many allergic and collagen diseases. Administration of a short- or intermediate-acting preparation given as a single dose in the morning (before 9 AM) is likely to produce fewer side effects and less pituitary–adrenal suppression than a divided dosage regimen with the same agent or an equivalent dosage of a long-acting agent. Alternate-day therapy (ie, total 48-hr dosage administered every other morning) with intermediate-acting agents (eg, prednisone) further reduces the prevalence...
and degree of side effects. However, it might not be uniformly effective in treating all disease states, unless large doses are used (eg, 40–60 mg every other day for adults requiring long-term corticosteroid therapy for asthma). Complete adrenal suppression might not occur with single daily doses given in the morning if the prednisone dose is ≤15 mg, but Cushing's syndrome can still occur and patients should receive supplemental corticosteroids during periods of unusual stress.\(^5,20\)

**In times of stress** (eg, surgery, severe trauma, serious illness), patients on long-term corticosteroid therapy (>5 mg/day prednisone or equivalent) should receive supplemental IV hydrocortisone 100–300 mg/day or PO prednisone 25–75 mg/day in divided doses for 1–3 days.\(^1,2,5,20\) Guidelines for withdrawal from glucocorticoid therapy have been published.\(^1,5\)

**Common initial doses are:** PO for acute asthma exacerbations in adults and adolescents 40–60 mg/day in 1 or 2 doses for 3–10 days; hospitalized patients may require a parenteral preparation and a larger dosage (eg, methylprednisolone 120–180 mg/day in 3–4 divided doses for 48 hr, then 60–80 mg/day).\(^23\)

Reduce dosage to minimum effective maintenance dosage as soon as possible; PO as an adjunct therapy for *Pneumocystis carinii* pneumonia (with an arterial PO\(_2\) ≤70 mm Hg or an arterial–alveolar gradient ≥35 mm Hg) 40 mg bid for 5 days begun with antimicrobial therapy, then 20 mg bid for 5 days, then 20 mg/day for the duration of antimicrobial therapy;\(^22,24\) PO for rheumatoid arthritis 5–7.5 mg/day;\(^2\) PO for collagen diseases 1 mg/kg/day in divided doses;\(^2\) PO for acute gout 30–50 mg/day, gradually decreasing over 10 days;\(^25\)

PO for nephrotic syndrome 1–2 mg/kg/day;\(^2\) PO for skin disorders 40 mg/day, up to 120 mg/day in pemphigus;\(^2\) PO for ulcerative colitis 10–30 mg/day or, if severe, 60–120 mg/day;\(^2\) PO for thrombocytopenia 0.5 mg/kg/day;\(^2\) PO for organ transplantation (in combination with other immunosuppressants) 50–100 mg once, then taper dosage and make further dosage adjustments based on the clinical situation;\(^2\) PO for acute exacerbations of multiple sclerosis 200 mg/day for 1 week, then 80 mg every other day for 1 month.

**Special Populations. Pediatric Dosage.** Dosage depends on disease state and patient response rather than strict adherence to age or body weight. **Common initial doses are:** PO for acute asthma 1–2 mg/kg/day, to a maximum of 60 mg/day in 1–2 divided doses for 3–10 days; hospitalized patients may require a parenteral preparation and a larger dosage (eg, methylprednisolone 1 mg/kg q 6 hr for 48 hr, then 1–2 mg/kg/day, to a maximum of 60 mg/day, in 2 divided doses).\(^21\) PO for inflammation or immunosuppression 0.5–2 mg/kg/day in 1–4 divided doses.\(^26\)

**Geriatric Dosage.** Consider using lower dosages for decreased body size.

**Dosage Forms.** Soln 1, 5 mg/mL; Syrup 1 mg/mL; Tab 1, 2.5, 5, 10, 20, 50 mg.

**Patient Instructions.** (See Class Instructions: Corticosteroids.)

**Pharmacokinetics. Onset and Duration.** (See Oral Corticosteroids Comparison Chart.)

**Serum Levels.** Serum concentration is not directly correlated with therapeutic effect.\(^5,11\) A timed prednisolone serum drug level can be useful for estimating clearance and identifying abnormalities in absorption, elimination, or patient compliance.\(^27\)
**Fate.** Bioavailability is 80 ± 11%, prednisone is about 75% plasma protein bound and prednisolone is 90–95% plasma protein bound, depending on serum concentration. Liver disease does not impair conversion to active metabolite. In fact, patients with liver disease and hypoalbuminemia are more likely to suffer major side effects of prednisone as a result of decreased protein binding and reduced prednisolone clearance. Vₐ of prednisolone is 1.5 ± 0.2 L/kg; 3 ± 2% of a dose of prednisone is excreted unchanged in urine, with an additional 15 ± 5% excreted as prednisolone.²

\[ t_{1/2} \] (Prednisone) 3.6 hr; (prednisolone) 2.2 hr.² Biologic half-life exceeds serum half-life. (See Oral Corticosteroids Comparison Chart.)

**Adverse Reactions.** Dose- and duration-related side effects include fluid and electrolyte disturbances (with possible edema and hypertension), hyperglycemia and glycosuria, spread of herpes conjunctivitis, activation of tuberculosis, osteoporosis, bone fractures, myopathy, menstrual irregularities, behavioral disturbances (increasing with dosages >40 mg/day), poor wound healing, ocular cataracts, glaucoma, arrest of growth (in children), hirsutism, pseudotumor cerebri (primarily in children), and Cushing’s syndrome (moon face, buffalo hump, central obesity, easy bruising, acne, hirsutism, and striae).² ¹¹ ¹² ³⁰ ³¹ Prolonged therapy can lead to suppression of pituitary–adrenal function. Too rapid withdrawal of long-term therapy can cause acute adrenal insufficiency (eg, fever, myalgia, arthralgia, and malaise); adrenally suppressed patients cannot respond to stress.

**Contraindications.** Systemic fungal infections (except as maintenance therapy in adrenal insufficiency); administration of live virus vaccines in patients receiving immunsuppressive doses of corticosteroids.

**Precautions.** Pregnancy. Use with caution in diabetes mellitus; osteoporosis; peptic ulcer; esophagitis; tuberculosis; and other acute and chronic bacterial, viral, and fungal infections; hypertension or other cardiovascular diseases; hypothyroidism; immunizations; hypoalbuminemia; psychosis; and liver disease. Suppression of PPD and other skin test reactions can occur.

**Drug Interactions.** Corticosteroids can increase serum glucose levels, and an increase in the dosage of antidiabetic drugs might be required. Corticosteroids can decrease isoniazid and salicylate serum levels. Amphotericin B and loop and thiazide diuretics can enhance corticosteroid-induced potassium depletion. Carbamazepine, phenobarbital (and possibly other barbiturates), phenytoin (best documented with dexamethasone), rifampin, and possibly aminoglutethimide increase the metabolism of corticosteroids.

**Parameters to Monitor.** Observe for behavioral disturbances and signs or symptoms of Cushing’s syndrome. With short-term, high-dose therapy, frequently monitor serum potassium and glucose and blood pressure. With long-term therapy, monitor these parameters occasionally and perform periodic eye examinations and possibly stool guaiac. Monitor growth in infants and children on prolonged therapy.

**Notes.** Other, more expensive glucocorticoids offer minimal advantages over prednisone in most clinical situations.³ Dosage ranges for prednisolone are the...
same as those for prednisone. Patients who have received daily glucocorticoid therapy for less than 2 weeks do not require dosage tapering to prevent acute adrenal insufficiency; however, dosage tapering may be required to maintain an adequate clinical response.\textsuperscript{1,2,5} Efficacy in patients with stable COPD is controversial.\textsuperscript{31,33,34}
### ORAL CORTICOSTEROIDS COMPARISON CHART

<table>
<thead>
<tr>
<th>DURATION AND DRUG</th>
<th>DOSAGE FORMS</th>
<th>EQUIVALENT ANTI-INFLAMMATORY DOSE (MG)*</th>
<th>RELATIVE ANTI-INFLAMMATORY POTENCY*</th>
<th>RELATIVE MINERALOCORTICOID ACTIVITY</th>
<th>SERUM HALF-LIFE (HR)</th>
<th>COMMENTS</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>SHORT-ACTING GLUCOCORTICOIDS (BIOLOGIC ACTIVITY 8–12 HR)</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Cortisone</td>
<td>Tab 5, 10, 25 mg.</td>
<td>25</td>
<td>0.8</td>
<td>2</td>
<td>0.5</td>
<td>Must be metabolized to active form (hydrocortisone).</td>
</tr>
<tr>
<td>Various</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Hydrocortisone</td>
<td>Tab 5, 10, 20 mg Susp (as cypionate)</td>
<td>20</td>
<td>1</td>
<td>2</td>
<td>1.5</td>
<td>Daily secretion in adults is 20 mg.</td>
</tr>
<tr>
<td>Various</td>
<td></td>
<td></td>
<td></td>
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<td></td>
<td></td>
</tr>
<tr>
<td><strong>INTERMEDIATE-ACTING GLUCOCORTICOIDS (BIOLOGIC ACTIVITY 18–36 HR)</strong></td>
<td></td>
<td></td>
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<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Methylprednisolone</td>
<td>Tab 2, 4, 8, 16, 24, 32 mg.</td>
<td>4</td>
<td>5</td>
<td>0</td>
<td>2.2</td>
<td>Minimal sodium-retaining activity.</td>
</tr>
<tr>
<td>Medrol</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Various</td>
<td></td>
<td></td>
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<td></td>
<td></td>
</tr>
<tr>
<td>Prednisolone</td>
<td>Tab 5 mg Syrup 1, 3 mg/mL.</td>
<td>5</td>
<td>4</td>
<td>1</td>
<td>2.2</td>
<td>Minimal sodium-retaining activity.</td>
</tr>
<tr>
<td>Various</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Prednisone</td>
<td>Tab 1, 2.5, 5, 10, 20, 50 mg Soln 1, 5 mg/mL. Syrup 1 mg/mL.</td>
<td>5</td>
<td>4</td>
<td>1</td>
<td>3.6</td>
<td>Must be metabolized to active form (prednisolone).</td>
</tr>
<tr>
<td>Various</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Triamcinolone</td>
<td>Tab 4, 8 mg Syrup 0.8 mg/mL.</td>
<td>4</td>
<td>5</td>
<td>0</td>
<td>2.6</td>
<td></td>
</tr>
<tr>
<td>Aristocort Kenacort</td>
<td>Various</td>
<td></td>
<td></td>
<td></td>
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<td></td>
</tr>
</tbody>
</table>

(continued)
### ORAL CORTICOSTEROIDS COMPARISON CHART (continued)

| DURATION AND DRUG | DOSAGE FORMS | EQUIVALENT ANTI-INFLAMMATORY DOSE (MG)
<table>
<thead>
<tr>
<th></th>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>LONG-ACTING GLUCOCORTICOIDS (BIOLOGIC ACTIVITY 36–54 HR)</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Betamethasone</td>
<td>Tab 0.6 mg</td>
<td>0.6–0.75</td>
</tr>
<tr>
<td>Celestone</td>
<td>Syrup 0.12 mg/mL.</td>
<td>25</td>
</tr>
<tr>
<td>Dexamethasone</td>
<td>Tab 0.25, 0.5, 0.75, 1, 1.5, 2, 4, 6 mg</td>
<td>0.75</td>
</tr>
<tr>
<td>Decadron</td>
<td>Exr 0.1 mg/mL.</td>
<td>25</td>
</tr>
<tr>
<td>Hexadrol</td>
<td>Soln 0.1, 1 mg/mL.</td>
<td>0</td>
</tr>
<tr>
<td>Various</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>RELATIVE ANTI-INFLAMMATORY POTENCY</th>
<th>RELATIVE MINERALOCORTICOID ACTIVITY</th>
<th>SERUM HALF-LIFE (HR)</th>
<th>COMMENTS</th>
</tr>
</thead>
<tbody>
<tr>
<td>25</td>
<td>0</td>
<td>5+</td>
<td>Minimal sodium-retaining activity, but with high doses, retention may occur.</td>
</tr>
<tr>
<td>25</td>
<td>0</td>
<td></td>
<td>No sodium-retaining activity with low to moderate doses.</td>
</tr>
</tbody>
</table>

| **MINERALOCORTICOID (BIOLOGIC ACTIVITY 18–36 HR)** | | |
| Fludrocortisone | Tab 100 µg. | — | 10 |
| Florinef | | | 125 |

| | | | 3.5+ |

+ Anti-inflammatory potency does not correlate with immunosuppressive effects. From references 2, 12, 13, and 36.
Pharmacology. Topical corticosteroids have nonspecific, local anti-inflammatory effects in the dermal and epidermal skin layers that probably occur by inhibiting mediators in the arachidonic acid pathway in cells, by suppressing DNA synthesis at the cellular level, and by decreasing the influx of WBCs into the local area. Potency is dependent on the characteristics and concentration of the drug and the vehicle used and is usually measured by the assessment of the relative degree of skin blanching (vasoconstrictor assay).

Administration and Adult Dosage. Uses for the nonprescription hydrocortisone preparations include relief of itching, inflammation, and rashes caused by eczema; insect bites; poison oak, ivy, or sumac; soaps, detergents, or cosmetics; jewelry; seborrheic dermatitis, psoriasis; and external genital or anal itching. Prescription indications include relief of inflammatory and pruritic manifestations of corticosteroid-responsive dermatoses including contact or atopic dermatitis; nummular, stasis, or astematotic eczema; lichen planus; lichen simplex chronicus; insect and arthropod bite reactions; and first- and second-degree localized burns and sunburns. These products are usually applied sparingly in a light film, 2-4 times/day; however, with continuous use, a repository effect may make 1-2 applications/day as effective. High-potency agents should be reserved for short-term or intermittent use only but may be more effective and cause fewer adverse effects than continuous therapy with lower potency products. Treatment with very high-potency agents should not exceed 2 consecutive weeks and the total dosage should not exceed 50 g/week because of the hypothalamic-pituitary-adrenal (HPA) axis suppressing potential.

Dosage Forms. (See Topical Corticosteroids Comparison Chart.)

Patient Instructions. Avoid prolonged use around the eyes (or contact with the eyes); in the genital and rectal areas; on the face, armpits, and in skin creases. Do not use with occlusive dressings unless directed.

Missed Doses. Apply a missed dose as soon as you remember unless it is almost time for the regular schedule. If it is almost time for the regular application, then continue on the regular schedule. Do not apply a double dose.

Pharmacokinetics. The absorption of these drugs depends on the physical properties of the drug itself, the surface area of use, the thickness of the skin (greater absorption from the face, in skin folds, in the perineum, and on denuded skin; lesser absorption from the palms and soles), skin temperature or hydrational state (greater with increased skin temperature or increased hydration), the age of the patient (children have a greater surface area:mass ratio and increased systemic effects), the use of occlusive dressings, the vehicle, application frequency, and length of treatment. Approximately 12-30 g is sufficient to cover the adult body one time. (See Topical Corticosteroids Comparison Chart.)

Adverse Reactions. Adverse reactions occur more frequently with increasing product potency and include local burning, itching, irritation, erythema, dryness, folliculitis, hypertrichosis, acneiform eruptions, hypopigmentation, rosacea, skin atrophy, striae, telangiectasias, purpura, perioral dermatitis, overgrowth of skin bacteria and fungi, allergic contact dermatitis, and cataracts or glaucoma with pro-
longed application around the eye. Systemically, there can be enough absorption of potent steroids to cause suppression of the HPA axis, causing symptoms of Cushing’s syndrome, and growth retardation, particularly in young children.

### Adrenal Hormones

**TOPICAL CORTICOSTEROIDS COMPARISON CHART**

<table>
<thead>
<tr>
<th>CLASS AND DRUG</th>
<th>BRAND NAMES</th>
<th>DOSAGE FORMS</th>
<th>STRENGTHS</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>LOW-POTENCY AGENTS</strong> <em>(Modest anti-inflammatory effects. Safest for chronic application, face and intertriginous areas, use under occlusion, and use on young children and infants.)</em></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Aclometasone Dipropionate</td>
<td>Aclovate</td>
<td>Cream, Ointment</td>
<td>0.05%</td>
</tr>
<tr>
<td>Desonide</td>
<td>Tridesilon, Various</td>
<td>Cream, Ointment</td>
<td>0.05%</td>
</tr>
<tr>
<td>Fluocinolone Acetonide</td>
<td>Various</td>
<td>Cream, Ointment</td>
<td>0.01%</td>
</tr>
<tr>
<td>Hydrocortisone</td>
<td>Hytone, Various</td>
<td>Cream, Lotion, Ointment, Solution, Spray</td>
<td>0.5–2.5%</td>
</tr>
<tr>
<td>Hydrocortisone Acetate</td>
<td>Various</td>
<td>Cream, Ointment</td>
<td>0.5, 1%</td>
</tr>
<tr>
<td><strong>MEDIUM-POTENCY AGENTS</strong> <em>(Effective for moderate inflammatory dermatoses [eg, chronic eczematous dermatoses]. May be used for a limited duration of time on the face and intertriginous areas.)</em></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Betamethasone Valerate</td>
<td>Valisone, Various</td>
<td>Cream, Foam, Lotion, Ointment</td>
<td>0.01–0.1%, 0.12%, 0.1%</td>
</tr>
<tr>
<td>Desoximetasone</td>
<td>Topicort, Various</td>
<td>Cream, Gel</td>
<td>0.05%</td>
</tr>
<tr>
<td>Fluocinolone Acetonide</td>
<td>Synalar, Various</td>
<td>Cream, Ointment, Solution</td>
<td>0.01, 0.025%, 0.01%</td>
</tr>
<tr>
<td>Hydrocortisone Valerate</td>
<td>Westcort</td>
<td>Cream, Ointment</td>
<td>0.2%</td>
</tr>
<tr>
<td>Mometasone Furoate</td>
<td>Elocon</td>
<td>Cream, Lotion, Ointment</td>
<td>0.1%</td>
</tr>
<tr>
<td>Triamcinolone Acetonide</td>
<td>Aristocort, Kenalog</td>
<td>Cream, Lotion, Ointment</td>
<td>0.025%, 0.1%</td>
</tr>
<tr>
<td><strong>HIGH-POTENCY AGENTS</strong> <em>(May be used for more severe eczematous dermatoses [eg, lichen simplex chronicus, psoriasis]. May be used for intermediate duration, with the exception of areas of thickened skin and chronic conditions. May be used on the face or in intertriginous areas for short periods of time.)</em></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Amcinonide</td>
<td>Cyclocort</td>
<td>Cream, Lotion</td>
<td>0.1%</td>
</tr>
<tr>
<td>Betamethasone</td>
<td>Diprosone, Various</td>
<td>Cream, Lotion, Ointment</td>
<td>0.05%, 0.1%</td>
</tr>
<tr>
<td>Dipropionate</td>
<td>Various</td>
<td>Cream, Lotion, Ointment, Aerosol</td>
<td>(continued)</td>
</tr>
</tbody>
</table>
Antidiabetic Drugs

**ACARBOSE**

_Acetate (Precose)_

**Pharmacology.** Acarbose is an oral α-glucosidase inhibitor indicated for the management of hyperglycemia caused by type 2 diabetes mellitus. Inhibition of this gut enzyme system effectively reduces the rate of complex carbohydrate digestion and the subsequent absorption of glucose, thereby lowering postprandial glucose excursions in type 2 diabetes. In obese and nonobese patients with type 2 diabetes, acarbose monotherapy is associated with a 0.5–1% decrease in hemoglobin A1c.

**Administration and Adult Dosage.** PO for type 2 diabetes (as monotherapy or with a sulfonylurea) 25 mg, tid initially, just before meals. Increase to 50 mg tid after 4–8 weeks and, if necessary, to 100 mg tid after 4–8 additional weeks. Dosages >100 mg tid are not recommended because of increased risk of hepatotoxicity, and patients weighing ≤60 kg should not receive >50 mg tid.

**Special Populations.** _Pediatric Dosage._ Safety and efficacy not established.
Geriatric Dosage. Same as adult dosage.

Dosage Forms. Tab 25, 50, 100 mg.

Patient Instructions. Take acarbose at the beginning of each meal. When a meal is skipped, also skip taking this medication. If a dose is missed, do not take it unless it is just before the next meal. If hypoglycemia occurs, dextrose (glucose) needs to be ingested; sucrose (table sugar) is not effective.

Pharmacokinetics. Fate. The drug is poorly absorbed from the GI tract (<2%). It undergoes extensive metabolism in the GI tract via intestinal flora and digestive enzymes. The clinical effect is not dependent on the serum level achieved. All absorbed acarbose and metabolites are renally excreted. In patients with renal impairment, plasma acarbose concentrations are elevated in relation to the degree of renal dysfunction.

\[ t_{1/2} \] About 2 hr with normal renal function.

Adverse Reactions. The major side effects of acarbose are flatulence, diarrhea, and abdominal pain. Acarbose monotherapy is not associated with hypoglycemia; however, patients managed with combination therapy (with a sulfonylurea or insulin) can experience hypoglycemia secondary to the other drug. In this setting, manage hypoglycemia with oral glucose (if the patient is conscious) or IV glucose or glucagon (if the patient is unconscious) rather than with a complex carbohydrate (eg, sucrose). Attempting to manage hypoglycemia with oral sugar sources other than glucose is not effective in acarbose-treated patients and might have grave consequences.

Contraindications. Inflammatory bowel disease; colonic ulceration; obstructive bowel disorders; cirrhosis; type 1 diabetes; history of diabetic ketoacidosis.

Precautions. Use with caution in patients with disorders of digestion or absorption or with medical conditions that might deteriorate with increased intestinal gas formation. Not recommended in patients with Cr >2 mg/dL.

Drug Interactions. Charcoal and other intestinal adsorbents as well as digestive preparations containing amylase, pancreatin, and related enzymes should not be taken concurrently with acarbose.

Parameters to Monitor. Monitor clinical symptoms of hyperglycemia (mainly polyphagia, polyuria, polydipsia, or numbing or tingling of feet) or hypoglycemia (hunger, nervousness, sweating, palpitations, headaches, confusion, drowsiness, anxiety, or blurred vision) when taken concurrently with insulin or insulin secretagogues (eg, sulfonylureas). Self-monitoring of fasting and selected postprandial blood glucose levels by the patient is also helpful. (See Blood Glucose Monitors Comparison Chart.) Long-term diabetic control may best be monitored using hemoglobin A\(_1c\).\(^{44}\)

Notes. Miglitol (Glyset—Bayer) is an \(\alpha\)-glucosidase inhibitor that has similar indications, uses, and side effects as acarbose. The dosage of the two drugs is the same. The clinical benefits, if any, of miglitol over acarbose have not been determined. Voglibose (Takeda America) is another \(\alpha\)-glucosidase inhibitor in clinical trials.
Aldose Reductase Inhibitors

Prolonged hyperglycemia causes excess flux of glucose into tissues, and glucose is shunted to the polyol pathway, resulting in excess sorbitol production. Excess intracellular sorbitol causes a reduction in the uptake of myoinositol and ultimately a down-regulation in the Na+/K+-ATPase system. This process is thought to be one of the biochemical mechanisms leading to the development of neuropathy, collagen disorders, cataracts, and possibly retinopathy in patients with diabetes. Because aldose reductase is the rate-limiting enzyme in this pathway, aldose reductase inhibitors are being studied as a possible means of decreasing the sorbitol-linked sequelae of diabetes. Although this is a promising class of drugs, the side effects, dosage regimens, and long-term benefits are to be determined. Aldose reductase inhibitors currently under investigation are fidarestat and zopolrestat (Alond—Pfizer).

Pharmacology.

Insulin promotes cellular uptake of glucose, fatty acids, and amino acids and their conversion to glycogen, triglycerides, and proteins. Beef and pork insulins are extracted and purified from the animal’s pancreas. Human insulin is produced by recombinant DNA technology or enzymatic conversion of pork insulin. No differences in side effects or long-term control of diabetes have been observed between human insulin and highly purified pork insulin.

Administration and Adult Dosage. SC for type 1 diabetes usual initial dosage ranges of 0.6–0.75 unit/kg/day in divided doses. During the first week of therapy, the dosage requirement might escalate to 1 unit/kg/day in divided doses because of insulin resistance and the usual age group (adolescents) being treated. The dosage requirement can temporarily decrease to 0.1–0.5 unit/kg/day if the patient experiences a “honeymoon phase.” Dosage adjustments are made on the basis of clinical symptoms, blood glucose levels, and hemoglobin A1c values. Insulin can be administered by various methods depending on a number of factors. Single daily SC injections of intermediate-acting insulin are often used but should
not usually be relied on to adequately control blood glucose levels in the type 1 patient because they are not sufficient to prevent long-term complications even though they can offer protection from diabetic ketoacidosis. Intensive forms of insulin therapy, which may provide better glycemic control, include the split-and-mixed regimen (2 SC injections daily of mixed short- and long-acting insulin), multiple daily SC doses of short-acting insulin in combination with a single injection of long-acting insulin, and insulin pump therapy. IV, SC, or IM for diabetic ketoacidosis (IV preferred for patients in shock) 0.1 unit/kg, followed by a continuous infusion of 0.1–0.2 unit/kg/hr. If the serum glucose does not change in the first hour, double the insulin rate, with further adjustments in insulin dosage based on glucose levels. Fluid and electrolyte repletion must accompany insulin therapy. SC for type 2 diabetes (patients unresponsive to oral agent therapy or with extreme hyperglycemia: fasting serum glucose >200–225 mg/dL) may need as little as 5–10 units/day or >100 units/day. Patients who require <30 units/day may be well controlled with 1 injection/day of intermediate-acting insulin; patients who require >30 units/day should be treated with ≥2 injections/day. Insulin resistance in the type 2 population is usually associated with obesity. Weight reduction and improved glycemic control usually improve insulin response.

**Special Populations. Pediatric Dosage.** (See Administration and Adult Dosage.) Common maintenance dosages are 0.6–0.9 unit/kg/day in divided doses in prepubertal children, up to 1.5 units/kg/day during puberty, and <1 unit/kg/day after puberty. Requirements occasionally can be as high as 200 units/day during growth spurts.

**Geriatric Dosage.** Same as adult dosage.

**Other Conditions.** Insulin requirements may be decreased in patients with renal or hepatic impairment or hypothyroidism. Requirements may be increased during pregnancy (especially in the second and third trimesters), in patients with high fever, hyperthyroidism, or severe infections; and after trauma or surgery.

**Dosage Forms.** (See Insulins Comparison Chart.)

**Patient Instructions.** Instruct patients in the following areas: use of insulin syringes and needles; storage, mixing, and handling of insulin; urine ketone testing; blood glucose testing; adherence to proper diet and regular meals; personal hygiene (especially the feet); and recognition and treatment of hypoglycemia and hyperglycemia. (See Sulfonylurea Agents.)

**Pharmacokinetics. Onset and Duration.** Human insulin is more soluble than animal-source insulins and may have a shorter onset and duration of action. (See Insulins Comparison Chart.)

**Serum Levels.** Patients with diabetes vary widely in their responses to insulin, and serum levels are not normally monitored clinically.

**Fate.** The rate of absorption depends on the insulin type. (See Insulins Comparison Chart.) Serum levels are affected by obesity, diet, degree of activity, pancreatic β-cell activity, growth hormone, and circulating antibodies. Insulin is metabolized primarily in the liver, although the kidneys are responsible for the metabolism of up to 40% of the daily insulin output.
t½. (Regular insulin) 4–5 min after IV administration.

**Adverse Reactions.** Hypoglycemia is dose related. Patients being treated with intensive insulin regimens of ≥3 injections/day are more prone to hypoglycemic episodes than are patients treated with the conventional 1–2 injections/day. Local allergic reactions, with an onset of 15 min–4 hr, are usually caused by insulin impurities; 70% of these patients have histories of interrupted treatment. Immune or nonimmune insulin resistance occurs occasionally. Lipohypertrophy at the injection site can occur, especially with repeated use of the same site. Lipohypertrophy also can occur at the injection site and be less frequent with the highly purified animal or human insulins. Allergy, resistance, and lipohypertrophy can be overcome by switching to a more highly purified product (eg, human insulin). In general, pork insulin is less antigenic than beef-pork or pure beef insulin (neither is available in the United States) because it is structurally more similar to human insulin, which is the least immunogenic.

**Contraindications.** Hypoglycemic episodes.

**Precautions.** Use with caution in patients with renal or hepatic disease or hypothyroidism. Insulin requirements can change with exercise or infection, or when switching animal sources or to more purified products.

**Drug Interactions.** Alcohol can produce hypoglycemia, especially in fasting patients; moderate increases in blood glucose can occur in nonfasting patients. Oral contraceptives, corticosteroids, furosemide, niacin (large doses), diazoxide, thiazide diuretics, and thyroid hormones (large doses) can increase insulin requirements. Anabolic steroids can decrease the insulin requirement. Avoid MAOIs in patients with diabetes because they can interfere with the normal adrenergic response to hypoglycemia by prolonging the action of antidiabetic agents. β-Blockers prolong hypoglycemic episodes and inhibit tachycardia and tremors, which are signs of hypoglycemia (sweating is not inhibited); hypertension can occur during hypoglycemia; cardioselective β-blockers (eg, atenolol, metoprolol) are less likely than nonselective types (eg, nadolol, propranolol) to cause problems.

**Parameters to Monitor.** Monitor blood glucose routinely. (See Blood Glucose Monitors Comparison Chart.) Long-term diabetic control is best monitored using hemoglobin A1c. The patient should continually watch for subjective symptoms of hypoglycemia and hyperglycemia. Observe for signs of lipohypertrophy, lipohypertrophy, and allergic reactions.

**Notes.** Human insulin is the insulin of choice for patients with insulin resistance, pregnancy, or allergy; new insulin-dependent patients; or any patient taking insulin intermittently. Insulin is stable for 1–2 months at constant room temperature and up to 24 months under refrigeration. Insulin is adsorbed by glass and plastic IV infusion equipment, with little difference between glass and plastic; maximal adsorption occurs within 15 sec. Adsorption can be minimized by the addition of small amounts (1–2%) of albumin to the infusion container; however, this may be costly and unnecessary because patient response is generally adequate without addition of albumin. Variation can be minimized by flushing all new IV administration equipment with 50 mL of the insulin-containing solution (thereby saturating “binding sites”) before it is used.
Pump devices are available to deliver insulin depending on or independent of a measured serum glucose level. “Open-loop” devices can deliver insulin at a constant rate and be manually controlled. “Closed-loop” devices (the “artificial pancreas”) can deliver insulin at variable rates in response to serum glucose but are used only in experimental settings.

Novel forms of insulin delivery are currently under investigation, with inhaled insulin showing the most promise. Pulmonary tissue provides a large absorptive surface for insulin. Insulin can be delivered effectively as an inhaled aerosol and in one study had an onset of action that was 23 min earlier than SC regular insulin and sustained its action for >3 hr. Ninety-nine units of inhaled insulin was metabolically equivalent to 10 units of SC insulin.54 Pulmonary-delivered insulins are currently in clinical trials.

**INSULIN ANALOGUES**

Insulin injected SC does not result in serum insulin concentrations that mimic normal physiologic insulin response. More than 30 human insulin analogues with different pharmacokinetic profiles have been produced using recombinant DNA technology. The goal of insulin analogue research is to produce a human insulin analogue with a rapid action (to provide bolus postprandial insulin) and a slow, extended release pattern (to provide basal insulin). Insulin lispro (Humalog) is a rapid-acting analogue with a pharmacokinetic profile between that of IV and SC regular human insulin. It has an onset in ≤15 min, a peak at 1–2 hr, and a duration of 2–4 hr. Insulin lispro offers a pharmacokinetic profile that is superior to regular human insulin when used to cover postprandial glycemic excursions. Another short-acting analogue, insulin aspart (Novolog), has a pharmacokinetic profile similar to that of insulin lispro.55 The long-acting analogue, insulin glargine (Lantus), has an onset of action of approximately 1 hr, with a sustained peak activity beginning at 4–5 hr and persisting for 24 hr. Insulin glargine is a basal insulin that is administered once daily at bedtime; it must be used in combination with a rapid-acting premeal insulin such as insulin lispro to achieve optimum results.55,56

<table>
<thead>
<tr>
<th>INSULINS COMPARISON CHARTa</th>
</tr>
</thead>
</table>

<table>
<thead>
<tr>
<th>PRODUCT</th>
<th>MANUFACTURER</th>
<th>STRENGTH</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>RAPID-ACTING (ONSET, &lt;0.25 HR; PEAK, 1–2 HR; DURATION 2–4 HR)</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Biosynthetic</td>
<td></td>
<td></td>
</tr>
<tr>
<td>NovoLog (Aspart)</td>
<td>Novo Nordisk</td>
<td>U-100</td>
</tr>
<tr>
<td>Humalog (Lispro)</td>
<td>Lilly</td>
<td>U-100</td>
</tr>
</tbody>
</table>

| **SHORT-ACTING (ONSET, 0.5–2 HR; PEAK, 3–4 HR; DURATION, 4–8 HR)** |
| Pork | | |
| Iletin II Regular | Lilly | U-100 |
| Purified Pork Regular | Novo Nordisk | U-100 |

| Human | | |
| Humulin R | Lilly | U-100, U-500 |
| Novolin R | Novo Nordisk | U-100 |
| Velosulin | Novo Nordisk | U-100 |

(continued)
## Insulins Comparison Chart (continued)

<table>
<thead>
<tr>
<th>Product</th>
<th>Manufacturer</th>
<th>Strength</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Intermediate-Acting (Onset, 2–4 HR; Peak, 8–14 HR; Duration, 14–24 HR)</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Pork</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Iletin II Lente</td>
<td>Lilly</td>
<td>U-100</td>
</tr>
<tr>
<td>Iletin II NPH</td>
<td>Lilly</td>
<td>U-100</td>
</tr>
<tr>
<td>Purified Pork Lente</td>
<td>Novo Nordisk</td>
<td>U-100</td>
</tr>
<tr>
<td>Purified Pork NPH</td>
<td>Novo Nordisk</td>
<td>U-100</td>
</tr>
<tr>
<td><strong>Human</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Humulin L (Lente)</td>
<td>Lilly</td>
<td>U-100</td>
</tr>
<tr>
<td>Humulin N (NPH)</td>
<td>Lilly</td>
<td>U-100</td>
</tr>
<tr>
<td>Novolin L (Lente)</td>
<td>Novo Nordisk</td>
<td>U-100</td>
</tr>
<tr>
<td>Novolin N (NPH)</td>
<td>Novo Nordisk</td>
<td>U-100</td>
</tr>
<tr>
<td><strong>Long Acting (Onset, 6–14 HR; Peak, None; Duration, 20–30 HR)</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Human</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Humulin U (Ultralente)</td>
<td>Lilly</td>
<td>U-100</td>
</tr>
<tr>
<td>Lantus (Glargine)</td>
<td>Aventis</td>
<td>U-100</td>
</tr>
<tr>
<td><strong>Fixed Combinations (Onset, 0.5–1 HR; Peak, 3–10 HR; Duration, 14–18 HR)</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Human</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Humalog Mix 75/25</td>
<td>Lilly</td>
<td>U-100</td>
</tr>
<tr>
<td>Humulin 70/30</td>
<td>Lilly</td>
<td>U-100</td>
</tr>
<tr>
<td>Novolin 70/30</td>
<td>Novo Nordisk</td>
<td>U-100</td>
</tr>
<tr>
<td>Humulin 50/50</td>
<td>Lilly</td>
<td>U-100</td>
</tr>
</tbody>
</table>

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There can be variations within the ranges of onset, peak, and duration among manufacturers. Onset and duration may be prolonged in long-standing diabetes, and large doses may have prolonged durations of action. Site of injection, depth of injection, and whether site is exercised, massaged, or has heat applied to it also affect rate of insulin absorption. Human insulins have a slightly more rapid onset and a shorter duration of action than animal-derived insulins.

These products contain isophane and regular insulin in the specified proportions; the first number designates the percentage of isophane insulin and the second designates the percentage of regular insulin.

Suspension of insulin lispro protamine and soluble insulin lispro. Onset is within 0.25 hr.

From references 51, and 57, and product information.

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### Metformin

**Glucophage**

**Pharmacology.** Metformin is a biguanide antihyperglycemic agent used in the management of type 2 diabetes mellitus. It does not affect insulin secretion; rather, it reduces hepatic glucose production and enhances glucose utilization by muscle. Reported increases in glucose utilization in muscle are 7–35%. In addition to blood glucose reductions (mean 53 mg/dL), metformin may have beneficial effects on serum lipids.58 (See Notes.)

**Administration and Adult Dosage.** PO for type 2 diabetes (immediate-release) initiate 500 mg tablets with a dosage of 1 tablet bid with morning and evening meals. Increase dosage in 500 mg/day increments at weekly intervals, to a maxi-
mum of 2.5 g/day. Initiate 850 mg tablets with 1 tablet/day before the morning meal. Increase dosage in 850 mg/day increments q 2 weeks, to a maximum of 850 mg tid. Individualize maintenance dosage based on glycemic response. Give all dosages up to 2 g/day in 2 divided doses; larger dosages require a tid regimen to reduce GI discomfort. PO (SR Tab) 500 mg/day with the evening meal initially, increasing in 500 mg/day increments at weekly intervals to a maximum of 2 g/day with the evening meal. To switch from the immediate-release to the SR formulation, give the same daily dosage of SR as a single dose with the evening meal.

**Special Populations.**

**Pediatric Dosage.** Safety and efficacy not established.

**Geriatric Dosage.** Initial and maintenance dosages should be lower in the elderly. Avoid usual maximum adult dosage. Do not start metformin in patients ≥80 yr unless renal function is normal.

**Dosage Forms.** Tab 500, 850, 1000 mg; SR Tab 500 mg (Glucophage XR); Tab 250 mg with glyburide 1.25 mg, 500 mg with glyburide 2.5 or 5 mg (Glucovance).

**Patient Instructions.** Take metformin just before meals to reduce gastrointestinal side effects (diarrhea, nausea, and heartburn). Contact your physician if gastrointestinal side effects persist. Do not take metformin if you develop a serious medical condition such as myocardial infarction, stroke, or serious infection; require surgery; consume excessive amounts of alcohol; or require x-ray procedures with contrast dyes. Discontinue metformin and contact your health care provider immediately if hyperventilation, muscle pain, malaise, unexplained drowsiness, or other unusual symptoms occur that might indicate the development of lactic acidosis.

**Missed Doses.** Take as soon as possible, unless the time for the next dose is near. Do not double doses.

**Pharmacokinetics.**

**Fate.** Absorption half-life is 0.9–2.6 hr for immediate-release tablets; peak levels occur at 4–8 hr (median 7 hr) with the SR formulation. Absolute bioavailability is 50–60% for both products. With immediate-release tablets, peak serum levels are 1–2 mg/L in patients with type 2 diabetes; with the SR formulation, peak levels are 20% lower. Plasma protein binding is negligible; $V_d$ is 654 ± 358 L after a single 850 mg oral dose; Cl is proportional to renal function. Metformin is excreted in the urine unchanged.\(^5^8\)

$t_{1/2}$ (Immediate-release) 1.7–4.5 hr with normal renal function.\(^5^8\)

**Adverse Reactions.** Acute side effects occur in as many as 30% of patients treated with metformin. Side effects include primarily GI complaints, such as diarrhea, abdominal discomfort, nausea, anorexia, and metallic taste. GI side effects are usually transient and dose related and can be mitigated by giving the drug just before meals, initiating therapy with small doses and slowly increasing the dosage. Metformin reduces serum vitamin $B_{12}$ levels in approximately 7% of patients but is rarely associated with anemia. Vitamin $B_{12}$ deficiency anemia can be treated with vitamin $B_{12}$ supplementation or by discontinuing metformin. Diminished vitamin $B_{12}$ absorption and transport can be improved with oral calcium supplementation.\(^5^9\) Lactic acidosis has been reported; however, almost all cases occur in patients in whom metformin was contraindicated or in patients who at-
tempted suicide by overdose. Lactic acidosis occurs in 0.03 case/1000 patient-yr, with fatalities in about 50% of cases.58

**Contraindications.** Acute or chronic metabolic acidosis; patients undergoing radiographic studies requiring contrast media (withhold metformin just before the radiographic study and do not reinstate for 48 hr after contrast media administration and upon documentation of normal renal function); abnormal Clcr or CrCl >1.5 mg/dL in males or >1.4 mg/dL in females; any disease that can cause hypoxia and result in accumulation of lactate (eg, CHF requiring pharmacologic treatment, MI, severe infections, stroke); hepatic dysfunction.

**Precautions.** Avoid in pregnancy and lactation.

**Drug Interactions.** Furosemide and nifedipine increase serum levels of metformin, the clinical relevance of which is unknown. Cimetidine reduces the tubular secretion of metformin and can increase peak serum concentrations by as much as 60%.50 (See also Insulin Drug Interactions.)

**Parameters to Monitor.** Monitor renal function, hepatic function, and CBC before initiation of therapy and at least annually thereafter. Monitor renal function more closely in the elderly because of the age-related changes in renal function and greater risk for acute renal failure. (See Contraindications.) The goal of therapy is to reduce fasting blood glucose and glycosylated hemoglobin levels to normal or near normal by using the lowest effective dosage of the drug.

**Notes.** Because of its effect on weight and lipids, metformin is an appropriate choice for initial monotherapy in obese, new-onset type 2 diabetic patients, whereas sulfonylureas are usually a better choice for nonobese patients. In patients who do not respond to metformin monotherapy, combination therapy with a sulfonylurea or a thiazolidinedione might be effective. Weight loss has been associated with metformin therapy (mean 0.8 kg); weight gain (mean 2.8 kg) has been found in patients treated with sulfonylureas.61 Reductions in total cholesterol, LDL cholesterol, and triglycerides of 5%, 8%, and 16%, respectively, and an increase of 2% in HDL cholesterol have been reported.

### NATEGLINIDE

**Pharmacology.** Nateglinide is a meglitinide similar to repaglinide that is a rapid-acting oral insulin secretagogue that stimulates insulin secretion in relation to serum blood glucose levels.

**Adult Dosage.** PO for type 2 diabetes (alone or in combination with metformin) 120 mg tid before each meal. For patients near their HbA1c goals, a dose of 60 mg can be used. Dosage adjustment is not necessary in the elderly or those with mild to severe renal impairment or mild to moderate hepatic impairment.

**Dosage Forms.** Tab 60, 120 mg.

**Pharmacokinetics.** Oral bioavailability is 72%. Peak plasma concentrations occur within 0.5–1.9 hr. Plasma protein binding is 97%; Cl is 8.4 L/hr. The drug is metabolized in the liver primarily by CYP3A4 and somewhat by CYP2C9, with 80% of the parent drug and glucuronide metabolites eliminated in the urine. Mild
to moderate hepatic cirrhosis does not markedly alter single-dose pharmacokinetics of nateglinide. Half-life is 1.4 hr. Administration with metformin does not alter the pharmacokinetics of either drug.

**Adverse Reactions.** The most frequent side effect is mild hypoglycemia manifested by increased sweating, tremor, dizziness, and increased appetite. Headache has occurred. Because of nateglinide’s hepatic metabolism and extensive protein binding, interactions with other drugs affecting CYP3A4 and CYP2C9 or drugs extensively protein bound might result in pharmacokinetic interactions.

**Contraindications, Precautions, and Parameters to Monitor.** *(See Repaglinide.)*

**REPAGLINIDE**

**Pharmacology.** Repaglinide is a meglitinide agent that stimulates insulin release from the pancreas, although it is structurally unrelated to sulfonylureas. Compared with the sulfonylureas, repaglinide has a quicker onset and shorter duration of action, resulting in a lower risk of prolonged hypoglycemia.

**Administration and Adult Dosage.** PO for type 2 diabetes newly treated patients with HbA1c <8% should start with 0.5 mg within 30 min before each meal. Patients previously treated with antidiabetic agents should start with 1 or 2 mg within 30 min before each meal. Increase dosage based on glycemic response, to a maximum of 4 mg/dose or 16 mg/day. Starting doses of repaglinide are unchanged when taken concurrently with metformin.

**Special Populations.** *Pediatric Dosage.* Safety and efficacy not established. *Geriatric Dosage.* Dosage adjustment is not needed unless renal function is compromised. However, the elderly are more sensitive to hypoglycemia and should be monitored closely with initiation of therapy. *Other Conditions.* No adjustment of the initial dosage is required in renal impairment but use caution with subsequent dosage increases. In patients with hepatic abnormalities, wait longer before increasing the dosage.

**Dosage Forms.** Tab 0.5, 1, 2 mg.

**Patient Instructions.** Take each dose 0 to 30 minutes before each meal, usually 15 minutes. Recognize signs and symptoms of hypoglycemia and treat accordingly. Skip your dose if you will miss a meal. Add a dose when you eat an extra meal.

**Missed Doses.** If you miss a dose, take your regular dose at your next scheduled meal. Do not double the dose.

**Pharmacokinetics.** *Fate.* Oral bioavailability is 56%. Peak plasma concentrations occur within 1 hr; food reduces mean peak concentration by 20%, although time to peak concentration is not altered. Serum concentrations are higher and prolonged in those with liver impairment. Plasma protein binding is >98%. $V_d$ is 31 L; $Cl$ is 38 L/hr. The drug is metabolized primarily in the liver by CYP3A4 to inactive metabolites excreted in the feces. $t_{1/2}$ About 1 hr.
Adverse Reactions. The most frequent side effect is hypoglycemia. Upper respiratory infections, sinusitis, nausea, diarrhea, constipation, arthralgia, and headache have been reported, but their frequencies are equal to or only slightly higher than that of placebo.

Contraindications. Diabetic ketoacidosis; type 1 diabetes.

Precautions. Pregnancy; lactation. Use cautiously in patients with renal impairment and those at increased risk of hypoglycemia, including those with hepatic or adrenal insufficiency and in debilitated, elderly, or malnourished patients. Hypoglycemia is more frequent in treatment of previously untreated patients and those with HbA1c <8%.65

Drug Interactions. Inhibitors of CYP3A4 (eg, ketoconazole, miconazole, erythromycin) inhibit the metabolism of repaglinide. CYP3A4 inducers (eg, rifampin, barbiturates, carbamazepine) might reduce serum levels of repaglinide.

Parameters to Monitor. Monitor fasting and selected postprandial blood glucose levels regularly and HbA1c periodically.

SULFONYLUREA AGENTS

Pharmacology. Sulfonylureas enhance insulin secretion from pancreatic β-cells and potentiate insulin action on several extrahepatic tissues. Long-term, sulfonylureas increase peripheral utilization of glucose, suppress hepatic gluconeogenesis, and possibly increase the sensitivity and/or number of peripheral insulin receptors. Second-generation sulfonylureas (eg, glyburide, glipizide, glimeperide) are more potent than first-generation agents and are used in much smaller dosages, with lower resultant blood levels. These lower serum concentrations decrease the likelihood of protein-binding displacement and hepatic metabolic interference.

Administration and Adult Dosage. (See Sulfonylurea Agents Comparison Chart.)

Special Populations. Pediatric Dosage. Safety and efficacy not established.

Geriatric Dosage. Start at the lower end of the dosage range and slowly titrate upward if needed. Observe precautions with renal or hepatic impairment.

Other Conditions. Dosage alterations may be necessary with all sulfonylureas in patients with severe hepatic dysfunction. With renal disease, especially in geriatric patients, there is an increased duration of action with chlorpropamide, acetohexamide, and possibly glyburide.66

Dosage Forms. (See Sulfonylurea Agents Comparison Chart.)

Patient Instructions. Eat a recommended diet consistently on a day-to-day basis. Take this medication at the same time each day (in the morning for once-daily medications). Report factors that might alter blood glucose levels (eg, infection, fasting states) and any side effects.

Missed Doses. Take a missed dose as soon as you remember unless it is near time of the next dose. Do not double doses.

Pharmacokinetics. (See Sulfonylurea Agents Comparison Chart.)

Adverse Reactions. Hypoglycemic reactions (especially with chlorpropamide), anorexia, nausea, vomiting, diarrhea, allergic skin reactions, and cholestatic jaun-
dice occur occasionally. Hematologic disorders, mild disulfiram-like reaction to alcohol, hyponatremia (most common with chlorpropamide but can occur with tolbutamide), and bone marrow suppression occur rarely.67

**Contraindications.** Pregnancy; type 1 diabetes; juvenile, unstable, or brittle diabetes; diabetes complicated by acidosis, ketosis, diabetic coma, major surgery, severe infection, or severe trauma.

**Precautions.** Patients sensitive to one sulfonylurea might experience cross-sensitivity to other sulfonylureas. Chlorpropamide can cause hyponatremia, particularly in elderly women taking diuretics.68

**Drug Interactions.** Drugs that have been reported to enhance sulfonylurea effects include chloramphenicol (chlorpropamide and tolbutamide), dicumarol, fluconazole (glipizide, glyburide, tolbutamide, and possibly others), sulfonamides, and high-dose salicylates. Rifampin stimulates the metabolism of tolbutamide and possibly other sulfonylureas. Drugs that impair glucose tolerance include oral contraceptives, corticosteroids, thiazide diuretics, furosemide, thyroid hormones (large doses), and niacin.69 Acute ingestion of alcohol in combination with sulfonylureas can produce severe hypoglycemia. (See also Insulin Drug Interactions.)

**Parameters to Monitor.** Monitor clinical symptoms of hyperglycemia (mainly polyphagia, polyuria, polydipsia, or numbing or tingling of feet) or hypoglycemia (hunger, nervousness, warmth, sweating, palpitations, headaches, confusion, drowsiness, anxiety, blurred vision, or paresthesias of lips). Monitor fasting serum glucose levels frequently at the initiation of therapy to gauge the adequacy of the dosage. Self-monitoring of fasting and selected postprandial blood glucose levels by the patient is also helpful. (See Blood Glucose Monitors Comparison Chart.) Long-term diabetic control may best be monitored using hemoglobin A1c.

**Notes.** Sulfonylureas are usually an appropriate choice for nonobese, new-onset type 2 diabetic patients, whereas metformin is more appropriate for obese type 2 patients.70 Individualize the choice of sulfonylurea based on the patient’s characteristics (eg, renal function, hepatic function, likelihood of hypoglycemia) and the pharmacokinetics of the drugs. Glyburide, glipizide, glimepiride, and chlorpropamide are more effective at lowering blood glucose than acetohexamide, tolazamide, or tolbutamide.52 Acetohexamide, tolazamide, and tolbutamide probably should be reserved for mild hyperglycemia or in those likely to develop hypoglycemia (eg, the elderly). In patients who do not respond to sulfonylurea monotherapy, combination therapy with insulin, metformin, rosiglitazone, pioglitazone, or acarbose may be effective.71 Glimeperide is the most potent sulfonylurea agent, has the lowest rate of hypoglycemia, and does not affect potassium channels in the heart.72
<table>
<thead>
<tr>
<th>DRUG</th>
<th>DOSAGE FORMS</th>
<th>DAILY DOSAGE</th>
<th>FATE</th>
<th>DURATION (HR)</th>
<th>COMMENTS</th>
</tr>
</thead>
<tbody>
<tr>
<td>Acetohexamide</td>
<td>Tab 250, 500 mg.</td>
<td>250 mg–1.5 g in 2 divided doses.</td>
<td>65% converted to an active metabolite (hydroxyhexamide).</td>
<td>12–18</td>
<td>May be useful in the elderly and others prone to hypoglycemia but avoid in patients with renal dysfunction.</td>
</tr>
<tr>
<td>Dymelor</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Chlorpropamide</td>
<td>Tab 100, 250 mg.</td>
<td>100–500 mg in a single dose.</td>
<td>Metabolized, and 20% excreted unchanged.</td>
<td>24–72</td>
<td>Avoid in elderly and in patients with renal dysfunction. Causes disulfiram-like reaction in 30% of patients.</td>
</tr>
<tr>
<td>Diabinese</td>
<td></td>
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<td></td>
<td></td>
</tr>
<tr>
<td>Various</td>
<td></td>
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</tr>
<tr>
<td>Tolazamide</td>
<td>Tab 100, 250, 500 mg.</td>
<td>100 mg–1 g in 1–2 divided doses.</td>
<td>Converted to weakly active metabolites.</td>
<td>16–24</td>
<td>Delayed onset of action (3–4 hr). May be useful in the elderly and others prone to hypoglycemia.</td>
</tr>
<tr>
<td>Tolinase</td>
<td></td>
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<td></td>
<td></td>
</tr>
<tr>
<td>Various</td>
<td></td>
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<td></td>
</tr>
<tr>
<td>Tolbutamide</td>
<td>Tab 500 mg.</td>
<td>500 mg–3 g in 2–3 divided doses.</td>
<td>Converted to inactive compounds.</td>
<td>6–12</td>
<td>May be useful in the elderly and others prone to hypoglycemia.</td>
</tr>
<tr>
<td>Orinase</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Various</td>
<td></td>
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<td></td>
</tr>
</tbody>
</table>

(continued)
### SULFONYLUREA AGENTS COMPARISON CHART (continued)

<table>
<thead>
<tr>
<th>DRUG</th>
<th>DOSAGE FORMS</th>
<th>DAILY DOSAGE</th>
<th>FATE</th>
<th>DURATION (HR)</th>
<th>COMMENTS</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>SECOND-GENERATION</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Glimepiride</td>
<td>Tab 1, 2, 4 mg.</td>
<td>1–8 mg in a single dose.</td>
<td>Converted to inactive and active metabolites.</td>
<td>24</td>
<td>Similar to glyburide. Lowest rate of hypoglycemia and does not affect cardiac potassium channels.</td>
</tr>
<tr>
<td>Amaryl</td>
<td></td>
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<td></td>
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</tr>
<tr>
<td>Glipizide</td>
<td>Tab 5, 10 mg</td>
<td>Non-SR 5–40 mg in 1–2 divided doses.</td>
<td>Converted to inactive metabolites.</td>
<td>10–24 (non-SR)</td>
<td>Take non-SR product on an empty stomach.</td>
</tr>
<tr>
<td>Glucotrol SR</td>
<td>Tab 2.5, 5, 10 mg.</td>
<td>SR 5–20 mg in a single dose.</td>
<td></td>
<td>18–24 (SR)</td>
<td></td>
</tr>
<tr>
<td>Glucotrol XL</td>
<td></td>
<td></td>
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<td></td>
<td></td>
</tr>
<tr>
<td>Glyburide</td>
<td>Tab 1.25, 2.5, 5 mg</td>
<td>1.25–20 mg in 1–2 divided doses.</td>
<td>Converted to inactive and active metabolites.</td>
<td>18–24</td>
<td>The micronized product (Glynase, various) offers no advantage over the nonmicronized products.</td>
</tr>
<tr>
<td>DiaBeta</td>
<td>Tab (micronized)</td>
<td>Micronized 0.75–12 mg in 1–2 divided doses.</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Glynase</td>
<td>1.5, 3, 4.5, 6 mg.</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Micronase</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Various</td>
<td></td>
<td></td>
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<td></td>
</tr>
<tr>
<td><strong>COMBINATION PRODUCT</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Glyburide and Metformin</td>
<td>Tab 1.25 mg glyburide plus 250 mg metformin, 2.5 mg or 5 mg glyburide plus 500 mg metformin.</td>
<td>1.25 mg/250 mg daily-bid initially, to a maximum of 20 mg/2000 mg daily in 1–2 divided doses.</td>
<td>(See individual agents.)</td>
<td>18–24</td>
<td></td>
</tr>
<tr>
<td>Glucovance</td>
<td></td>
<td></td>
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</tr>
</tbody>
</table>

From references 67, and 68 and product information.
Pharmacology. Pioglitazone is a thiazolidinedione antihyperglycemic agent used to improve insulin sensitivity in patients with type 2 diabetes. Insulin-dependent glucose disposal in skeletal muscle is improved and hepatic glucose production is decreased; both actions contribute to pioglitazone’s glucose-lowering effects. Pioglitazone is only effective in the presence of insulin; by itself it does not lead to hypoglycemia and does not increase insulin secretion. Because insulin is required for its action, pioglitazone should not be used in patients with type 1 diabetes. Rosiglitazone (Avandia) is another thiazolidinedione antidiabetic agent that acts similarly to pioglitazone.

Administration and Adult Dosage. PO for type 2 diabetes (monotherapy) 15–30 mg once daily with food; after a 4-week trial dosage can be increased to a maximum of 45 mg/day. If no response occurs at the maximum dose of 45 mg/day, other therapeutic options should be considered; (combination therapy with insulin, sulfonylurea, or metformin) 15–30 mg once daily initially, increasing q 4 weeks to a maximum of 45 mg/day. Dosages of insulin or sulfonylurea may need to be decreased based on the glucose-lowering response. For those on insulin, decrease the insulin dosage when fasting plasma glucose levels are <100 mg/dL or if hypoglycemic symptoms occur.

Special Populations. Pediatric Dosage. Safety and efficacy not established.

Geriatric Dosage. (>65 yr) no differences in efficacy or safety; dosage adjustments are not required.

Other Conditions. No dosage adjustment is required in renal impairment.

Dosage Forms. Tab 15, 30, 45 mg.

Patient Instructions. Take once daily without regard to meals. If you are taking insulin, a sulfonylurea, or other glucose-lowering agent, you should understand the signs and symptoms of hypoglycemia and its appropriate treatment. Report nausea, vomiting, abdominal pain, loss of appetite, or dark urine immediately to your health care provider. Because pioglitazone’s effect on oral contraceptives has not been established, other means of contraception may be required.

Missed Doses. If you forget to take a dose, take your regular dose the next day. Do not double the dose on the next day.

Pharmacokinetics. Fate. Oral absorption is rapid, with the peak plasma concentration in 2 hr. Steady-state serum levels are achieved in 7 days. Extensively bound to serum albumin (>99%). Vd is 10.5–26.5 L/kg. Metabolized extensively by hydroxylation and oxidation in the liver and principally by CYP2C8 and CYP3A4. Renal elimination is negligible (15–30%), with most of the oral dose believed to be excreted into the bile unchanged or as metabolites and subsequently eliminated in the feces.

\( t_{1/2} \) 16–24 hr.

Adverse Reactions. Mild to moderate hypoglycemia when used concurrently with a sulfonylurea or insulin. Headache, anemia (mean hemoglobin value decrease of 2–4%), edema, weight gain.73
**Contraindications.** Active liver disease or ALT levels exceeding 2.5 times the upper limit of normal.

**Precautions.** Premenopausal anovulatory individuals might resume ovulation, placing them at risk for pregnancy. Use cautiously in patients with edema.

**Drug Interactions.** Ethinyl estradiol and norethindrone plasma concentrations might be reduced, resulting in possible loss of contraceptive efficacy. Ketoconazole and possibly other drugs that inhibit CYP3A4 might inhibit the metabolism of pioglitazone. Additionally, CYP3A4-metabolized drugs such as calcium-channel blockers, corticosteroids, cyclosporine, and HMG-CoA reductase inhibitors have not been specifically studied but might affect the metabolism of pioglitazone.

**Parameters to Monitor.** Monitor serum ALT levels at the start of therapy, q 2 months for the first 12 months, and then periodically. If ALT is elevated 1–2.5 times the upper limit of normal at any time (before initiation or during therapy), the cause of the enzyme elevation should be determined. If ALT levels exceed 3 times the upper limit of normal, pioglitazone should be discontinued. Monitor fasting blood sugars and HbA1c. *(See Sulfonylureas.)*

**Notes.** In patients with type 2 diabetes on insulin therapy, pioglitazone often results in a decreased requirement for insulin. Patients on concomitant therapy with insulin or sulfonylureas should not alter the doses of the latter medications until positive changes in fasting plasma glucose levels are obtained (fasting blood sugars <100 mg/dL) or if symptoms of hypoglycemia are experienced. *(See Thiazolidinedione Comparison Chart.)*

### THIAZOLIDINEDIONE COMPARISON CHART

<table>
<thead>
<tr>
<th>DRUG</th>
<th>MONOTHERAPY INITIATION DOSE</th>
<th>COMBINATION INITIATION DOSE</th>
<th>MAXIMUM DOSE</th>
<th>COMMENTS</th>
</tr>
</thead>
<tbody>
<tr>
<td>Pioglitazone</td>
<td>15–30 mg once daily.</td>
<td>15–30 mg once daily.</td>
<td>45 mg once daily.</td>
<td>Approved for use with insulin, a sulfonylurea, or metformin. May improve lipid profile.</td>
</tr>
<tr>
<td>Actos</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Rosiglitazone</td>
<td>2 mg bid or 4 mg once daily.</td>
<td>2 mg bid or 4 mg once daily.</td>
<td>4 mg bid or 8 mg once daily.</td>
<td>Approved for use with a sulfonylurea or metformin. May increase LDL and HDL cholesterol.</td>
</tr>
<tr>
<td>Avandia</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>NAME MANUFACTURER</td>
<td>NAME RANGE (MG/DL)</td>
<td>TEST TIME (SEC)</td>
<td>FEATURES</td>
<td></td>
</tr>
<tr>
<td>-----------------------------------</td>
<td>--------------------</td>
<td>-----------------</td>
<td>------------------------------------------------------------------------------------------------------------------------------------------</td>
<td></td>
</tr>
<tr>
<td>Accu-Chek Advantage (Roche Diagnostics)</td>
<td>Advantage or Comfort Curve</td>
<td>10–600</td>
<td>40</td>
<td>No cleaning, wiping, or timing; touchable test strips; time and date; large target area; 100-value memory; PC downloading.</td>
</tr>
<tr>
<td>Accu-Chek Complete (Roche Diagnostics)</td>
<td>Advantage or Comfort Curve</td>
<td>10–600</td>
<td>40</td>
<td>2-step procedure; pushbutton selection; stores and analyzes up to 1000 values.</td>
</tr>
<tr>
<td>Accu-Chek Instant (Roche Diagnostics)</td>
<td>Instant Glucose</td>
<td>20–500</td>
<td>12</td>
<td>No wiping or timing; 9-value memory; Spanish version available.</td>
</tr>
<tr>
<td>Accu-Chek Simplicity (Roche Diagnostics)</td>
<td>Simplicity or Comfort Curve</td>
<td>20–500</td>
<td>25–30</td>
<td>Large target area requires small blood sample; 30-value memory.</td>
</tr>
<tr>
<td>Accu-Chek Voice Mate (Roche Diagnostics)</td>
<td>Comfort Curve</td>
<td>10–600</td>
<td>40</td>
<td>For visually impaired and the blind; voice guidance; no need to clean.</td>
</tr>
<tr>
<td>Assure (Chronimed)</td>
<td>Assure</td>
<td>30–550</td>
<td>35</td>
<td>Large touch-screen display; 180-test memory.</td>
</tr>
<tr>
<td>AtLast Blood Glucose System (Amira Medical)</td>
<td>AtLast</td>
<td>40–400</td>
<td>15</td>
<td>Sample taken from the forearm, upper arm, thigh; 10-test memory with 14-day average.</td>
</tr>
<tr>
<td>CheckMate Plus (Questar Medical)</td>
<td>CheckMate Plus</td>
<td>25–500</td>
<td>15–70</td>
<td>Display provides words for guidance; no wiping or timing; 6 language prompts; data port allows downloading to a PC.</td>
</tr>
<tr>
<td>ExactTech (Abbott Laboratories)</td>
<td>ExactTech</td>
<td>40–450</td>
<td>30</td>
<td>Credit-card size and shape; no wiping, timing, or cleaning; last reading recall.</td>
</tr>
<tr>
<td>ExactTech RSG (Abbott Laboratories)</td>
<td>ExactTech RSG</td>
<td>40–450</td>
<td>30</td>
<td>No calibration required; no cleaning or maintenance.</td>
</tr>
<tr>
<td>FastTake (LifeScan)</td>
<td>FastTake</td>
<td>20–600</td>
<td>15</td>
<td>Very small blood sample needed; compact; 150-test memory; warning to test ketones when range is 240–600; PC downloading.</td>
</tr>
</tbody>
</table>

(continued)
<table>
<thead>
<tr>
<th>NAME MANUFACTURER</th>
<th>TEST STRIP USED</th>
<th>RANGE (MG/DL)</th>
<th>TEST TIME (SEC)</th>
<th>FEATURES</th>
</tr>
</thead>
<tbody>
<tr>
<td>Glucometer DEX (Bayer Diagnostics)</td>
<td>Glucometer DEX</td>
<td>10–600</td>
<td>30</td>
<td>10-test cartridge; 100-test memory; PC downloading.</td>
</tr>
<tr>
<td></td>
<td>Test Sensors</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Glucometer Elite Diabetes Care System (Bayer Diagnostics)</td>
<td>Elite</td>
<td>20–600</td>
<td>30</td>
<td>No buttons; turns on when test strip inserted; blood touched to tip of test strip is automatically drawn; lancing device included; videotape available; 20-test memory.</td>
</tr>
<tr>
<td>Glucometer Elite XL (Bayer Diagnostics)</td>
<td>Elite</td>
<td>20–600</td>
<td>30</td>
<td>No buttons; 120-test memory; 14-day average; lancing devices and lancets included; 120-test memory; video available.</td>
</tr>
<tr>
<td>Glucometer Encore Diabetes Care System (Bayer Diagnostics)</td>
<td>Encore</td>
<td>20–600</td>
<td>15</td>
<td>Automatic 3-min shutoff; 10-test memory; Spanish instructions available.</td>
</tr>
<tr>
<td>Medisense 2 Card (Abbott Laboratories)</td>
<td>Medisense 2 or</td>
<td>20–600</td>
<td>20</td>
<td>No cleaning, wiping, timing; individually wrapped test strips; credit-card size; large display window.</td>
</tr>
<tr>
<td></td>
<td>Precision Q.I.D.</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Medisense 2 Pen (Abbott Laboratories)</td>
<td>Medisense 2 or</td>
<td>20–600</td>
<td>20</td>
<td>No cleaning, wiping, timing; individually wrapped test strips; pen size.</td>
</tr>
<tr>
<td></td>
<td>Precision Q.I.D.</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>One Touch Basic (LifeScan)</td>
<td>Genuine One Touch</td>
<td>0–600</td>
<td>45</td>
<td>75-test memory; large easy-to-handle test strips; single-button coding.</td>
</tr>
<tr>
<td>One Touch FastTake (LifeScan)</td>
<td>FastTake</td>
<td>20–600</td>
<td>15</td>
<td>Very small blood sample; warning to test ketones when range is 240–600; 150-test memory; PC downloading.</td>
</tr>
<tr>
<td>One Touch Profile (LifeScan)</td>
<td>One Touch</td>
<td>0–600</td>
<td>45</td>
<td>No timing, wiping or blotting; large display in English, Spanish, and 17 other languages; time and date; cleaning notification; 250-test memory; 14- and 30-day test averages.</td>
</tr>
</tbody>
</table>

(continued)
<table>
<thead>
<tr>
<th>NAME</th>
<th>MANUFACTURER</th>
<th>TEST STRIP USED</th>
<th>RANGE (MG/DL)</th>
<th>TEST TIME (SEC)</th>
<th>FEATURES</th>
</tr>
</thead>
<tbody>
<tr>
<td>One Touch SureStep</td>
<td>(LifeScan)</td>
<td>SureStep</td>
<td>0–500</td>
<td>15–30</td>
<td>Large display; touchable test strips; 150-test memory; PC downloading.</td>
</tr>
<tr>
<td>Precision Extra</td>
<td>(Abbott Laboratories)</td>
<td>Precision Extra</td>
<td>20–600</td>
<td>20</td>
<td>Also measures ketones; 450-test memory.</td>
</tr>
<tr>
<td>Precision Q.I.D.</td>
<td>(Abbott Laboratories)</td>
<td>Precision Q.I.D.</td>
<td>20–600</td>
<td>20</td>
<td>Automatic start when small blood sample applied; OK to touch strip; compact size; large display; 150-test memory; data-downloading capacity.</td>
</tr>
<tr>
<td>Prestige</td>
<td>(Home Diagnostics)</td>
<td>Prestige</td>
<td>25–600</td>
<td>10</td>
<td>Nonwipe system; blood applied to strip outside the monitor; large display; universal symbols guide user; English and Spanish videos available.</td>
</tr>
<tr>
<td>Prestige XL</td>
<td>(Home Diagnostics)</td>
<td>Prestige</td>
<td>25–600</td>
<td>10</td>
<td>365-test memory; large display; English and Spanish videos available.</td>
</tr>
<tr>
<td>Select GT</td>
<td>(Chronimed)</td>
<td>Select GT</td>
<td>30–600</td>
<td>50</td>
<td>Large display; blood application inside or outside the meter; 100-test memory; universal symbols.</td>
</tr>
<tr>
<td>Supreme II</td>
<td>(Chronimed)</td>
<td>Supreme</td>
<td>30–600</td>
<td>50</td>
<td>Large display; blood application inside or outside the meter; 100-test memory; universal symbols.</td>
</tr>
<tr>
<td>SureStep</td>
<td>(LifeScan)</td>
<td>SureStep</td>
<td>0–500</td>
<td>15–30</td>
<td>Large display; touchable test strip; 10-test memory; PC downloading.</td>
</tr>
</tbody>
</table>

Adapted from reference 75.
Contraceptives

Class Instructions. Oral Contraceptives. Take this drug at approximately the same time each day for maximum efficacy. This drug may be taken at bedtime or with food, milk, or an antacid if stomach upset occurs. Use an additional form of contraception concurrently during the first 7 days of the oral progestin-only products or if you do not start your oral contraceptives on day 1 of menses. If spotting occurs and no oral doses have been missed, continue to take tablets even if spotting continues. Report immediately if any of the following occur: new severe or persistent headache; blurred or loss of vision; shortness of breath; severe leg, chest, or abdominal pain; or any abnormal vaginal bleeding. Hormonal contraceptives do not protect against HIV infection or other sexually transmitted diseases.

Pharmacology. These products contain an estrogen, ethinyl estradiol or mestranol, and one of several 19-nortestosterone progestins, which are taken in a cyclic fashion, usually 21 of 28 days. As contraceptives, estrogens suppress follicle-stimulating hormone (FSH) and luteinizing hormone (LH) to inhibit ovulation, cause edematous endometrial changes that are hostile to implantation of the fertilized ovum, accelerate ovum transport, and produce degeneration of the corpus luteum (luteolysis). Progestins inhibit ovulation by suppression of LH, inhibit sperm capacitation, slow ovum transport, produce a thinning endometrium that hampers implantation, and cause cervical mucus changes that are hostile to sperm migration. Induction of a pseudopregnancy state and anovulation improves symptoms of endometriosis. Anovulatory dysfunctional uterine bleeding caused by unopposed estrogen or estrogen withdrawal responds to progestins. (See Contraception Efficacy, Risks and Benefits of Oral Contraceptives Comparison Charts.)

Administration and Adult Dosage. PO for contraception (monophasic combinations) 1 tablet daily beginning on the first day of menses and continue for 21 days;

### BLOOD GLUCOSE TEST STRIPS COMPARISON CHART: STRIPS FOR VISUAL READING

<table>
<thead>
<tr>
<th>NAME (MANUFACTURER)</th>
<th>COLOR CHART INCREMENTS (MG/DL)</th>
<th>PROCEDURE</th>
</tr>
</thead>
<tbody>
<tr>
<td>Chemstrip bG (Roche Diagnostics)</td>
<td>20, 40, 80, 120, 180, 240, 400, 800.</td>
<td>Wipe after 1 min, read after 2 min.</td>
</tr>
<tr>
<td>Glucostix Reagent Strips (Roche Diagnostics)</td>
<td>20, 40, 70, 110, 140, 180, 250, 400, 800.</td>
<td>Blot after 30 sec, read 90 sec later.</td>
</tr>
<tr>
<td>Select GT Strips (Chronimed)</td>
<td>Low, 40, 70, 120, 180, 240, 400, high.</td>
<td>Wait 60 sec, turn strip over and read.</td>
</tr>
<tr>
<td>Supreme Strips (Chronimed)</td>
<td>Low, 40, 70, 120, 180, 240, 400, high.</td>
<td>Wait 60 sec, turn strip over and read.</td>
</tr>
</tbody>
</table>

Adapted from reference 75.
stop for 7 days and start the next cycle of 21 tablets. Combination 28-day products (7 inert or iron tablets) are taken, 1 tablet daily continuously; (multiphasic combinations) 1 tablet daily beginning on the first day of menses (Triphasil only) or manufacturer states first Sunday after the beginning of menstruation (if menstruation begins on Sunday, take first tablet on that day), although day-1 start is most effective, then 1 tablet daily for 21 or 28 days as above. **PO for contraception postpartum** start 6 weeks postpartum if not breastfeeding; lactation prolongs period of infertility. **PO for contraception postabortion** start immediately if gestation is terminated at 12 weeks or earlier; start in 1 week if gestation is terminated at 13–28 weeks. **PO for emergency postcoital contraception** (Ovral, Preven) 2 tablets taken as soon as possible after coitus and 2 more tablets taken 12 hr later, but within 72 hr after coitus; or (Lo-Ovral) 4 tablets taken as soon as possible after coitus and 4 more taken 12 hr later, but within 72 hr after coitus.76–78 (See Notes.) **PO for dysfunctional uterine bleeding (anovulatory cycles)** (any combination agent) 1 tablet daily to qid for 5–7 days for acute bleeding, then 1 tablet daily cyclically as for contraception for 3 months to prevent further bleeding.79 **PO for dysmenorrhea or endometriosis** (any combination tablet) 1 tablet daily continuously for 15 weeks, followed by 1 drug-free week; repeat 16-week cycle for 6–12 months to induce a pseudopregnant state.80

**Special Populations. Geriatric Dosage.** Same as adult dosage.

**Other Conditions.** Discontinue oral contraceptives at least 2 weeks before elective major surgery and do not reinstitute until at least 2 weeks afterward. Stop immediately in patients undergoing emergency surgery or immobilization for long periods; institute low-dose SC heparin or other appropriate thromboembolitic prophylaxis in the postoperative period and restart cycle 4 weeks after returning to normal activities.78 Start with an agent containing at least 50 μg estrogen in women receiving rifampin or any cytochrome P450–inducing anticonvulsant.76–78

**Dosage Forms.** (See Oral Contraceptive Agents Comparison Chart.)

**Patient Instructions.** (See Class Instructions: Oral Contraceptives.) (Contraception) any menstrual irregularities and bothersome side effects should diminish after the first 3–4 cycles. Report if no menses occur for 2 months. (Acute anovulatory bleeding) expect heavy and severely cramping flow 2–4 days after stopping therapy, with normal periods thereafter.

**Missed Doses.** If you miss 1 active dose, take it as soon as you remember it and take the next tablet at the correct time even if you take 2 tablets on the same day or at the same time. If you miss 2 active doses in week 1 or 2, take 2 tablets on the day you remember and 2 tablets the next day. If you miss 2 active doses in week 3 or miss 3 or more active tablets, then (if you start on day 1) start a new pack the same day or (if you start on Sunday) take 1 tablet daily until Sunday and then start a new pack that day. Use an alternative form of contraception for the next 7 days after you miss 2 or more active doses in weeks 1, 2, or 3 or abstain from sex for the next 7 days.

**Pharmacokinetics. Onset and Duration.** Onset of contraception after 1 week of oral regimen. Dysfunctional uterine bleeding should decrease within 12–24 hr of starting the regimen.79
Serum Levels. No correlation of estradiol or mestranol serum levels with pharmacologic activity.

Fate. There are marked intra- and interpatient variabilities in the pharmacokinetics of all these agents. All are concentrated in body fat and endometrium and penetrate poorly into breast milk.\textsuperscript{81–90} Desogestrel is a prodrug that undergoes extensive first-pass and possibly gut-wall metabolism to its active form, 3-ketodesogestrel. Bioavailability is 63 ± 7%; 65% bound to albumin and 35% to sex hormone–binding globulin (SHBG); SHBG increases by about 200% during long-term use. V\(_d\) is 2.4 ± 1.1 L/kg; Cl is 0.2 ± 0.1 L/hr/kg. About 45% is recovered in urine as glucuronides (38–61%), sulfates (23–29%), and unconjugated forms (14–28%); 31% is recovered in feces.\textsuperscript{81,82,91–93}

Ethinyl estradiol is rapidly absorbed, with peak concentrations in 60 ± 30 min; bioavailability is 59 ± 13%.\textsuperscript{81,86,87} It undergoes extensive small intestine and hepatic first-pass metabolism and conjugation to sulfates and hydroxylation to active 2-hydroxyethinyl estradiol and other hydroxylated metabolites. Ethinyl estradiol is 98.5% bound to albumin and not bound to SHBG. V\(_d\) is 5 ± 2 L/kg; reported Cl has ranged from 0.4 ± 0.2 to 1 ± 0.3 L/hr/kg. About 23–59% is excreted in urine, 30–53% in feces as glucuronides and sulfates, and 28–43% undergoes enterohepatic circulation with a rebound in estradiol levels 10–14 hr after administration.\textsuperscript{78,81,83–87,91,93} (See Estradiol and Its Esters monograph for estrogen replacement.)

Ethynodiol diacetate undergoes rapid absorption and hydrolysis to norethindrone and its metabolites in vivo. (See Progestin-Only Contraceptives.)

Mestranol is approximately 54% demethylated to ethinyl estradiol; serum levels of ethinyl estradiol after oral administration of 50 μg of mestranol are equivalent to those after 35 μg of ethinyl estradiol.\textsuperscript{87,90} Norgestrel/levonorgestrel, norethindrone, and norethindrone acetate (see Progestin-Only Contraceptives).

Norethynodrel is rapidly converted to norethindrone in vitro.\textsuperscript{83,90} Norgestimate undergoes hepatic and gut metabolism to levonorgestrel (15.4 ± 5.4%), norgestrel acetate (9.5 ± 1.7%), norgestrel oxime (10.6 ± 1.8%), and 8.1 ± 4.5% as other conjugated metabolites.\textsuperscript{84} It is not bound to sex hormone-binding globulin. From 35% to 49% is excreted in urine (57% conjugated sulfates and glucuronides and 12% unconjugated) and 16–49% in feces.

\(t_{1/2}\) (Desogestrel) 24 ± 5 hr;\textsuperscript{91,92} (ethinyl estradiol) 15 ± 3 to 33 ± 10 hr;\textsuperscript{91,91} (levonorgestrel) 31.4 ± 18.5 hr;\textsuperscript{80–86,89,90} (norgestimate) 16 hr, (norethindrone) 7.6 ± 1.9 hr.\textsuperscript{81–85,90,94}

Adverse Reactions. The risk of major congenital malformations is not increased if oral contraceptives are taken during pregnancy.\textsuperscript{95,96} Most of the risks of oral contraceptives are minimal with the lower dosages of estrogens and progestins currently available.\textsuperscript{78,96–100} (See Risks and Benefits of Oral Contraceptives and Hormone Excess and Deficiency Symptomatology Comparison Charts.)

Contraindications. Known or suspected pregnancy; presence or history of thrombophlebitis or thromboembolic disorders; presence or history of carcinoma of breast or genitals, or other estrogen-dependent tumors; cerebral vascular or coronary artery disease; uncontrolled hypertension; focal migraine; markedly impaired liver function; hepatic adenoma or carcinoma; cholestatic jaundice of pregnancy
or jaundice with prior oral contraceptive use; undiagnosed abnormal genital bleeding; malabsorption syndrome, heavy smoking (>15 cigarettes/day) in women ≥35 yr;78,98 polycythemia vera because of greater tendency for deep vein thrombosis. (See Notes.)

**Precautions.** Use with caution in patients with hyperlipidemia, diabetes, conditions that might be aggravated by fluid retention (eg, hypertension, convulsions, migraine, and cardiac or renal dysfunction), or severe varicosities, in adolescents in whom regular menses are not established, and during lactation.

**Drug Interactions.** Oral contraceptives might be less effective, resulting in increased breakthrough bleeding or pregnancy, when given with some antibiotics (eg, ampicillin, griseofulvin, metronidazole, nitrofurantoin, neomycin, penicillin, rifampin, tetracycline), or anticonvulsants (eg, barbiturates, carbamazepine, phenytoin). Administer doses of vitamin C ≥1 g/day at least 4 hr before or after oral contraceptives to avoid increasing the bioavailability of ethinyl estradiol; use caution if long-term vitamin C intake is discontinued.78,101

**Parameters to Monitor.** Complete pretreatment physical examination with special reference to blood pressure, breasts, abdomen, pelvic organs, and Pap smear at least q 1–2 yr.

**Notes.** The initial oral contraceptive prescribed should be a combined product (eg, Ovcon-35, Ortho-Cept, Desogen, Ortho-Novum 7/7/7, Tri-Norinyl, Ortho-Cyclen) containing the smallest effective dose of estrogen (≤35 μg ethinyl estradiol) and progestin (≤0.15 mg desogestrel or levonorgestrel, ≤1 mg norethindrone, or ≤0.25 mg norgestimate) that provides an acceptable pregnancy rate and minimizes side effects.76,78 Prescribing oral contraceptives to smokers ≥35 yr requires adequate informed consent because of a doubled risk of cardiovascular disease.97,99 The health risks of pregnancy in healthy, nonsmoking women in their forties is greater than the risks of taking sub-50 μg estrogen or progestin-only contraceptives.96,99 (See Risks and Benefits of Oral Contraceptives Comparison Chart.) For emergency postcoital contraception, 2 doses of a combination contraceptive might be somewhat less effective than high-dose estrogens.102

<table>
<thead>
<tr>
<th>PROGESTIN-ONLY CONTRACEPTIVES:</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>LEVONORGESTREL/NORGESTREL</strong></td>
</tr>
<tr>
<td><strong>MEDROXYPROGESTERONE ACETATE</strong></td>
</tr>
<tr>
<td><strong>NORETHINDRONE</strong></td>
</tr>
</tbody>
</table>

**Pharmacology.** Norgestrel and norethindrone are 19-nortestosterone derivatives; only the 1-isomer of norgestrel (levonorgestrel) is active. Medroxyprogesterone acetate is a 17α-acetoxypregesterone derivative with greater progestational activity and oral efficacy than native progesterone. These compounds share the actions of progestins, although progestin-only contraceptives suppress ovulation in only about 50% of cycles. (See Combination Oral Contraceptives.)

**Administration and Adult Dosage.** **PO for contraception** (norethindrone) 0.35 mg/day or (norgestrel) 0.075 mg/day continuously at the same time each day,
starting on the first day of menses or immediately postpartum. IM (medroxyprogesterone acetate) 150 mg q 3 months, starting within 5 days of menses or immediately postabortion or postpartum (within 5 days to 6 weeks of delivery). In breastfeeding mothers, the first dose is recommended at 6 weeks postpartum, although some clinicians give it 3–6 weeks postpartum. PO for emergency postcoital contraception (Plan B) 1 tablet taken as soon as possible after coitus and 1 more tablet taken 12 hr later, but within 72 hr after coitus; Subdermal (levonorgestrel) 216 mg (6 Norplant implants) q 5 yr; insert within 7 days of onset of menstruation, immediately postabortion, or no earlier than 6 weeks postpartum if breastfeeding; insertion and removal require a simple surgical procedure performed by trained personnel. (See also Progesterone.)

**Dosage Forms.** Tab (norethindrone) 0.35 mg (Micronor, Nor-Q.D.); (norgestrel) 0.075 mg (Ovrette); (levonorgestrel) 0.75 mg (Plan B). Implant Pellet (levonorgestrel) kit of 6 capsules, each containing 36 mg (Norplant). Inj (medroxyprogesterone acetate) use only the 150 mg/mL dosage form of Depo-Provera for contraception.

**Patient Instructions.** (See Class Instructions: Oral Contraceptives.) Spotting and breakthrough bleeding occur more frequently than with the combination oral contraceptives during the first few months of use; notify prescriber if this persists through the third month. (Plan B) If you vomit within 1 hour of taking a tablet, call your health care provider to discuss whether to repeat the dose. You might experience spotting during use of this medication, and your next menstrual period might be delayed. If it is delayed more than 7 days, you might be pregnant. (Depo-Provera) use an alternative form of contraception for the first 2 weeks if your first injection is more than 5 days after the start of menses. Cessation of menses is common after 1 to 2 years. (Norplant) use an alternative form of contraception for the first 24 hours if inserted more than 7 days after the start of menses. Irregular bleeding patterns should become more regular 9 to 12 months after insertion. The implants may be visible under the skin. Removal at 5 years must be done by trained personnel.

**Missed Doses.** (Oral contraceptives) If you miss a dose, even if it is taken only 3 hours late, use an additional backup method for the next 48 hours. Take the missed dose as soon as you remember. If menses do not occur within 45 days, discontinue the contraceptive, use an alternate nonhormonal method of contraception, and make sure you are not pregnant. Because of the higher risk of failure if 1 tablet is missed every 1 to 2 cycles, consider changing the time of tablet taking or using a different contraceptive.

**Pharmacokinetics.** Onset and Duration. (Oral contraceptives) onset after 1 week; duration 24 hr. (Depo-Provera) onset is within 24 hr if given within 5 days of menses; the drug prevents ovulation the first month of use; ovulation is inhibited for at least 14 weeks after 150 mg IM; mean interval before return of ovulation after last injection is 9 months; 70% of former users conceive within the first 12 months after stopping. (Norplant) onset is within 24 hr after subdermal implantation if inserted within 7 days of menses; immediately reversible once removed, normal ovulatory cycles return during the first month after removal.
Serum Levels. (Ovulation inhibition) levonorgestrel 0.2 μg/L (0.64 nmol/L); medroxyprogesterone >0.1 μg/L (0.25 nmol/L); norethindrone 0.4 μg/L (1.34 nmol/L).

Fate. Levonorgestrel is completely absorbed orally with no first-pass metabolism. Peak serum levels occur in 1.1 ± 0.4 hr, are dose dependent, and exhibit considerable interindividual variations. Oral administration of 30 μg yields peak levels of 0.9 ± 0.7 μg/L (2.9 ± 2.2 nmol/L); 150 μg yields 3.6 ± 0.5 μg/L (11.5 ± 1.6 nmol/L); 250 μg yields 5 ± 0.5 μg/L (16 ± 1.6 nmol/L). Within 24 hr after implantation of Norplant, levonorgestrel produces serum levels >0.3 μg/L (0.96 nmol/L). Release of 80 μg/day of levonorgestrel during the first 6–12 months yields levels of 0.4 ± 0.1 μg/L (1.1 ± 0.4 nmol/L); thereafter, release of 25–35 μg/day yields levels that remain above 0.28 ± 0.16 μg/L (0.9 ± 0.5 nmol/L) for the remainder of the 5 yr. Levels are unmeasurable within 48 hr after removal of the implant. Levonorgestrel is concentrated in body fat and endometrium but penetrates poorly into breast milk (approximately 10% of serum levels); it is bound 69.4% to sex hormone-binding globulin and 30% to albumin. V_d is 1.5 ± 0.4 L/kg; Cl is 0.05 ± 0.01 L/hr/kg. Conjugated glucuronides, sulfates, and unconjugated levonorgestrel and its metabolites are excreted 45% in urine and 32% in feces. (See Medroxyprogesterone Acetate and Norethindrone.)

t_1/2. (Levonorgestrel) 31.4 ± 18.5 hr; (medroxyprogesterone acetate) about 50 days, reflecting slow IM absorption from depot; (norethindrone) 6.4 ± 3 hr.

Adverse Reactions. (Oral contraceptives) menstrual irregularities, including spotting, breakthrough bleeding, prolonged cycles, and amenorrhea, are frequent. Because ovulation is suppressed in only about 50% of cycles, functional ovarian cysts might occur. Most resolve spontaneously within 4 weeks, and surgical intervention is usually not necessary. Ectopic pregnancy occurs in 6% of all pregnancies. Low doses of progestins have minimal effects on the following: serum glucose, insulin or lipid levels; coagulation; liver or thyroid function; blood pressure; or cardiovascular complications. (Plan B) nausea, abdominal pain, fatigue, headache and menstrual changes occur frequently. (Depo-Provera) menstrual irregularities, spotting, and breakthrough bleeding are frequent in the first 12 months after IM injection; amenorrhea (after 1 yr), infertility (up to 18 months), and weight gain of 1–1.5 kg also occur. Reversible reduced bone density changes occur with >5 yr of use as contraceptive, but there is no clinical evidence of fractures. Long-term use (>5 yr) does not increase the overall risk of ovarian, liver, breast, or cervical cancer but reduces the risk of endometrial cancer for at least 8 yr after stopping. (Norplant) the most frequent adverse effects are irregular menstrual bleeding, headaches, weight gain, mood changes and depression, premenstrual bilateral mastalgia, galactorrhea (especially after discontinuation of lactation), acne, outbreaks of genital herpes in patients with a history. Rarely, rash, implant expulsions, and local complications (eg, infection, hematoama formation, irritation, allergic reactions to adhesives) occur.

Contraindications. Thrombophlebitis or history of deep vein thrombophlebitis or thromboembolic disorders; known or suspected carcinoma of the breast or en-
dometrium, or other estrogen-dependent tumors; undiagnosed abnormal genital bleeding. Known or suspected pregnancy is a contraindication, but the risk of congenital malformations is not increased when progestin-only contraceptives are taken during pregnancy. Although acute liver disease, benign or malignant liver tumors, history of cholestatic jaundice of pregnancy, or jaundice with prior hormonal contraceptive use are listed as contraindications by manufacturers, liver disease is not considered by others to be a contraindication to progestin-only contraceptives.

**Precautions.** Use with caution in patients with histories of depression, diabetes, gestational diabetes, coronary artery disease, cerebrovascular disease, hyperlipidemia, liver disease, or hypertension. Although progestins are not harmful to the fetus during the first 4 months of pregnancy, confirm a negative pregnancy test before reinjecting women >2 weeks late for their IM injection. Progestin-only contraceptives used during breastfeeding pose no risk to the infant and they usually do not decrease breastmilk production if begun ≥6 weeks postpartum.

**Drug Interactions.** Rifampin and cytochrome P450–inducing anticonvulsants can decrease efficacy. Long-term use of griseofulvin can increase menstrual irregularities.

**Parameters to Monitor.** Complete pretreatment physical examination with special reference to blood pressure, breasts, abdomen, pelvic organs, and Pap smear at least q 1–2 yr.

**Notes.** Progestin-only contraceptives are the hormonal contraceptives of choice during breastfeeding or in patients with contraindications to estrogen therapy (eg, hypertension, diabetes, hyperlipidemia, smokers). Long-term noncontraceptive benefits of IM medroxyprogesterone acetate are decreases in menstrual blood loss, anemia, candidal vulvovaginitis, pelvic inflammatory disease, and endometrial cancer. A 30% reduction in seizure frequency was observed in a small group of women with uncontrolled seizures who became amenorrheic with medroxyprogesterone.
<table>
<thead>
<tr>
<th>METHOD</th>
<th>AVERAGE PREGNANCY RATES PER 100 WOMAN-YR</th>
</tr>
</thead>
<tbody>
<tr>
<td>Oral Monophasic Combination</td>
<td></td>
</tr>
<tr>
<td>&lt;30 µg ethinyl estradiol (EE)ᵃ</td>
<td>0.75</td>
</tr>
<tr>
<td>35–49 µg EE</td>
<td>0.27</td>
</tr>
<tr>
<td>50 µg EE</td>
<td>0.16</td>
</tr>
<tr>
<td>Oral Multiphasic Combination</td>
<td>0.33</td>
</tr>
<tr>
<td>Oral Progestin Onlyᵃ</td>
<td></td>
</tr>
<tr>
<td>Age 25–30 yr</td>
<td>3.1</td>
</tr>
<tr>
<td>Age 30–34 yr</td>
<td>2.0</td>
</tr>
<tr>
<td>Age 35–39</td>
<td>1.0</td>
</tr>
<tr>
<td>Age ≥40 yr</td>
<td>0.3</td>
</tr>
<tr>
<td>Lactating</td>
<td>0.3</td>
</tr>
<tr>
<td>Subdermal Progestin Implant</td>
<td>0.2</td>
</tr>
<tr>
<td>Injectable Depot Progestin</td>
<td>0.3</td>
</tr>
<tr>
<td>Emergency Postcoital (within 72 hr of intercourse)</td>
<td>0.2–2.5ᵇ</td>
</tr>
<tr>
<td>Oral 2 tablets q 12 hr for 2 doses</td>
<td></td>
</tr>
<tr>
<td>Ethinyl estradiol 5 mg/day for 5 days</td>
<td>0.5–1.6ᵇ</td>
</tr>
<tr>
<td>Intrauterine Device</td>
<td></td>
</tr>
<tr>
<td>Copper T 380</td>
<td>0.5</td>
</tr>
<tr>
<td>Progestasert</td>
<td>2.9</td>
</tr>
<tr>
<td>Barrier Method</td>
<td></td>
</tr>
<tr>
<td>Diaphragm</td>
<td>1.9</td>
</tr>
<tr>
<td>Condomᶜ</td>
<td>3.6</td>
</tr>
<tr>
<td>Vaginal Sponge</td>
<td>10</td>
</tr>
<tr>
<td>Cervical Cap</td>
<td>13</td>
</tr>
<tr>
<td>Vaginal Spermicide (cream, foam, jelly)</td>
<td>11.9</td>
</tr>
<tr>
<td>Vaginal Spermicide (cream, foam, jelly)</td>
<td></td>
</tr>
<tr>
<td>Other</td>
<td></td>
</tr>
<tr>
<td>Tubal Sterilization</td>
<td>&lt;1</td>
</tr>
<tr>
<td>Coitus Interruptus</td>
<td>6.7</td>
</tr>
<tr>
<td>Rhythm</td>
<td>15.5</td>
</tr>
<tr>
<td>Abstinence Method</td>
<td>70</td>
</tr>
</tbody>
</table>

ᵃMestranol 50 µg is approximately equal to 35 µg of ethinyl estradiol.
ᵇPostcoital contraception numbers represent the percentage of women in whom pregnancies occur.
ᶜProtects against most sexually transmitted diseases.

## Oral Contraceptive Agents Comparison Chart

<table>
<thead>
<tr>
<th>PRODUCT</th>
<th>CYCLE</th>
<th>ESTROGEN</th>
<th>PROGESTIN</th>
<th>POTENCY</th>
<th>BREAKTHROUGH BLEEDING AND SPOTTING (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>Estrogenic</td>
</tr>
<tr>
<td>Alesse, Levilite</td>
<td>21, 28</td>
<td>Ethinyl estradiol 20 µg.</td>
<td>Levonorgestrel 0.1 mg.</td>
<td>+</td>
<td>+</td>
</tr>
<tr>
<td>Loestrin 1/20</td>
<td>21, 28</td>
<td>Ethinyl estradiol 20 µg.</td>
<td>Norethindrone acetate 1 mg.</td>
<td>+</td>
<td>++</td>
</tr>
<tr>
<td>Desogen, Ortho-Cept</td>
<td>21, 28</td>
<td>Ethinyl estradiol 30 µg.</td>
<td>Desogestrel 0.15 mg.</td>
<td>+</td>
<td>+</td>
</tr>
<tr>
<td>Loestrin 1.5/30</td>
<td>21, 28</td>
<td>Ethinyl estradiol 30 µg.</td>
<td>Norethindrone acetate 1.5 mg.</td>
<td>+</td>
<td>+++</td>
</tr>
<tr>
<td>Levlen, Levora 0.15/30, Nordette</td>
<td>21, 28</td>
<td>Ethinyl estradiol 30 µg.</td>
<td>Levonorgestrel 0.15 mg.</td>
<td>+</td>
<td>+</td>
</tr>
<tr>
<td>Low/Ovral, Low-Ogestrel</td>
<td>21, 28</td>
<td>Ethinyl estradiol 30 µg.</td>
<td>Norgestrel 0.3 mg.</td>
<td>+</td>
<td>+</td>
</tr>
<tr>
<td>Yasmin</td>
<td>28</td>
<td>Ethinyl estradiol 30 µg.</td>
<td>Drospirenone 3 mg.</td>
<td>+</td>
<td>+</td>
</tr>
<tr>
<td>Brevicon, ModiCon, Various</td>
<td>21, 28</td>
<td>Ethinyl estradiol 35 µg.</td>
<td>Norethindrone 0.5 mg.</td>
<td>++</td>
<td>+</td>
</tr>
<tr>
<td>Ovcon-35</td>
<td>21, 28</td>
<td>Ethinyl estradiol 35 µg.</td>
<td>Norethindrone 0.4 mg.</td>
<td>++</td>
<td>+</td>
</tr>
<tr>
<td>Demulen 1/35, Zovia 1/35E</td>
<td>21, 28</td>
<td>Ethinyl estradiol 35 µg.</td>
<td>Ethynodiol diacetate 1 mg.</td>
<td>+</td>
<td>++</td>
</tr>
</tbody>
</table>

(continued)
## ORAL CONTRACEPTIVE AGENTS COMPARISON CHART (continued)

<table>
<thead>
<tr>
<th>PRODUCT</th>
<th>CYCLE</th>
<th>ESTROGEN</th>
<th>PROGESTIN</th>
<th>POTENCY</th>
<th>BREAKTHROUGH BLEEDING AND SPOTTING (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>BIPHASIC COMBINATION PRODUCTS CONTAINING &lt;50 µG OF ESTROGEN</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Jenest-28</td>
<td>28</td>
<td>Ethinyl estradiol 35 µg (days 1–21).</td>
<td>Norethindrone 0.5 mg (days 1–7); 1 mg (days 8–21).</td>
<td>++</td>
<td>+</td>
</tr>
<tr>
<td>Mircette</td>
<td>28</td>
<td>Ethinyl estradiol 20 µg (days 1–21); 10 µg (days 24–28).</td>
<td>Desogestrel 0.15 mg (days 1–21).</td>
<td>+</td>
<td>+</td>
</tr>
<tr>
<td>Neocon 10/11, Nelova 10/11, Ortho-Novum 10/11</td>
<td>Various</td>
<td>Ethinyl estradiol 35 µg (days 1–21).</td>
<td>Norethindrone 0.5 mg (days 1–10); 1 mg (days 11–21).</td>
<td>++</td>
<td>+</td>
</tr>
</tbody>
</table>

(continued)
## Oral Contraceptive Agents Comparison Chart (continued)

<table>
<thead>
<tr>
<th>PRODUCT</th>
<th>CYCLE</th>
<th>ESTROGEN</th>
<th>PROGESTIN</th>
<th>POTENCY</th>
<th>BREAKTHROUGH BLEEDING AND SPOTTING (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>Estrogenic</td>
</tr>
<tr>
<td><strong>Triphasic Combination Products Containing &lt;50 µg of Estrogen</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Cyclessa 28</td>
<td>28</td>
<td>Ethinyl estradiol 25 µg (days 1–21)</td>
<td>Desogestrel 0.1 mg (days 1–7); 0.125 mg (days 8–14); 0.15 mg (days 15–21)</td>
<td>+</td>
<td>+</td>
</tr>
<tr>
<td>Estrostep 21, 28</td>
<td>21, 28</td>
<td>Ethinyl estradiol 20 µg (days 1–5); 30 µg (days 6–12); 35 µg (days 13–21)</td>
<td>Norethindrone acetate 1 mg (days 1–21)</td>
<td>+</td>
<td>+</td>
</tr>
<tr>
<td>Ortho-Novum 7/7/7</td>
<td>21, 28</td>
<td>Ethinyl estradiol 35 µg (days 1–21)</td>
<td>Norethindrone 0.5 mg (days 1–7); 0.75 mg (days 8–14); 1 mg (days 15–21)</td>
<td>++</td>
<td>+</td>
</tr>
<tr>
<td>Ortho Tri-Cyclen 21, 28</td>
<td>21, 28</td>
<td>Ethinyl estradiol 35 µg (days 1–21)</td>
<td>Norgestimate 0.18 mg (days 1–7); 0.215 mg (days 8–14); 0.25 mg (days 15–21)</td>
<td>++</td>
<td>+</td>
</tr>
</tbody>
</table>

(continued)
### ORAL CONTRACEPTIVE AGENTS COMPARISON CHART (continued)

<table>
<thead>
<tr>
<th>PRODUCT</th>
<th>CYCLE¹</th>
<th>ESTROGEN²</th>
<th>PROGESTIN³</th>
<th>POTENCY⁴</th>
<th>BREAKTHROUGH BLEEDING AND SPOTTING (%)⁵</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Tri-Norinyl</strong></td>
<td>21, 28</td>
<td>Ethinyl estradiol 35 µg (days 1–21).</td>
<td>Norethindrone 0.5 mg (days 1–7); 1 mg (days 8–16); 0.5 mg (days 17–21).</td>
<td>++</td>
<td>+</td>
</tr>
<tr>
<td><strong>Tri-Levlen, Triphasil, Trivora-28</strong></td>
<td>21, 28</td>
<td>Ethinyl estradiol 30 µg (days 21–28); 40 µg (days 7–11); 30 µg (days 12–21).</td>
<td>Levonorgestrel 0.05 mg (days 1–6); 0.075 mg (days 7–11); 0.125 mg (days 12–21)</td>
<td>+</td>
<td>+</td>
</tr>
</tbody>
</table>

#### MONOPHASIC COMBINATION AGENTS CONTAINING 50 µG OF ESTROGEN

<table>
<thead>
<tr>
<th>PRODUCT</th>
<th>CYCLE¹</th>
<th>ESTROGEN²</th>
<th>PROGESTIN³</th>
<th>POTENCY⁴</th>
<th>BREAKTHROUGH BLEEDING AND SPOTTING (%)⁵</th>
</tr>
</thead>
<tbody>
<tr>
<td>Norinyl 1 + 50, Ortho-Novum 1/50, Various</td>
<td>21, 28</td>
<td>Mestranol 50 µg.</td>
<td>Norethindrone 1 mg.</td>
<td>++</td>
<td>++</td>
</tr>
<tr>
<td>Demulen 1/50, Zovia 1/50E</td>
<td>21, 28</td>
<td>Ethinyl estradiol 50 µg.</td>
<td>Ethynodiol diacetate 1 mg.</td>
<td>+</td>
<td>++</td>
</tr>
<tr>
<td>PRODUCT</td>
<td>CYCLE&lt;sup&gt;2&lt;/sup&gt;</td>
<td>ESTROGEN&lt;sup&gt;b&lt;/sup&gt;</td>
<td>PROGESTIN&lt;sup&gt;c&lt;/sup&gt;</td>
<td>POTENCY&lt;sup&gt;d&lt;/sup&gt;</td>
<td>BREAKTHROUGH BLEEDING AND SPOTTING (%)&lt;sup&gt;h&lt;/sup&gt;</td>
</tr>
<tr>
<td>----------------</td>
<td>-------------------</td>
<td>---------------------------</td>
<td>-----------------------</td>
<td>--------------------</td>
<td>-----------------------------------------------</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Ethinyl estradiol</td>
<td>Norethindrone</td>
<td>Estrogenic&lt;sup&gt;e&lt;/sup&gt;</td>
<td>Progestational&lt;sup&gt;f&lt;/sup&gt;</td>
</tr>
<tr>
<td>Ovcon-50</td>
<td>21, 28</td>
<td>50 µg.</td>
<td>1 mg.</td>
<td>++</td>
<td>++</td>
</tr>
<tr>
<td>Ovral</td>
<td>21, 28</td>
<td>Ethinyl estradiol</td>
<td>Norgestrel</td>
<td>++</td>
<td>+++</td>
</tr>
<tr>
<td>PROGESTIN ONLY</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Micronor,</td>
<td>Continuous</td>
<td>None.</td>
<td>Norethindrone 0.35 mg.</td>
<td>0</td>
<td>+++</td>
</tr>
<tr>
<td>Nor-Q.D.</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Ovrette</td>
<td>Continuous</td>
<td>None.</td>
<td>Norgestrel 0.075 mg.</td>
<td>0</td>
<td>+</td>
</tr>
<tr>
<td>POSTCOITAL</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Plan B</td>
<td>2 doses of 1 tablet each</td>
<td>—</td>
<td>Levonorgestrel 0.75 mg.</td>
<td>—</td>
<td>—</td>
</tr>
<tr>
<td>Preven</td>
<td>2 doses of 2 tablets each</td>
<td>Ethinyl estradiol 50 µg.</td>
<td>Levonorgestrel 0.25 mg.</td>
<td>—</td>
<td>—</td>
</tr>
</tbody>
</table>

+++ = High; ++ = Moderate; + = Low; ± = Very Low; 0 = None.

<sup>2</sup>28-day cycles contain 7 inert or iron tablets to complete the 28-day cycle.

<sup>3</sup>Estrogen equivalent potency: ethinyl estradiol is about 1.5 times as potent as mestranol. Inhibition of ovulation requires 50 µg of ethinyl estradiol or 80 µg of mestranol.
Most products contain either norethindrone or norgestrel. Norethindrone may be preferred over norgestrel, which has a marked adverse effect on lipid profile (decreased HDL, increased LDL). Only levonorgestrel is biologically active and exists in newer preparations. Older preparations contain norgestrel, which also has an inactive D-isomer. Desogestrel and norgestimate have positive effects on lipids.

Potency designations are based on laboratory tests of individual components. Applicability of these methods for combination products used clinically has been questioned.

Overall estrogenic effect as modified by antiestrogenic or estrogenic effect of progestational component. Relative estrogenic potency as measured by affinity for estrogen receptor (all are relatively weak): norethynodrel > ethinodiol diacetate > norethindrone acetate > norethindrone > levonorgestrel/norgestimate/desogestrel. Antiestrogenic potency: norethindrone acetate > levonorgestrel > norethindrone > ethinodiol diacetate > norethynodrel > norgestimate > desogestrel.

Progestational potency as measured by delay of menses test. Relative progestogenic potency: norgestimate > desogestrel > levonorgestrel > norethindrone > norethindrone acetate > ethinodiol diacetate > norethynodrel.

Relative androgenic potency (prostate growth in rats): levonorgestrel > norethindrone > norethindrone acetate > ethinodiol diacetate > norethynodrel > norgestimate > desogestrel. Drospirenone is antiandrogenic.

Prevalence of breakthrough bleeding (BTB) decreases from the first cycle to third cycle by 50–66% per cycle; these figures represent data submitted to FDA on prevalence of BTB in the third cycle of use. BTB can result from either estrogen or progestin deficiency. Bleeding decreases after the first 6 months of use regardless of the formulation used.

Bi- and triphasic compounds are overall estrogen dominant.

Drospirenone is a spironolactone analogue that has antiandrogenic and antimineralocorticoid activity. As such, it can cause mild diuresis and potassium retention. Use with caution in patients predisposed to potassium retention (eg, renal insufficiency, ACE inhibitors, angiotensin receptor blockers, potassium-sparing diuretics).

From references 76, 78, 85, and 97.
<table>
<thead>
<tr>
<th>CONDITION</th>
<th>CLINICAL INFORMATION</th>
<th>COMMENTS</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>RISKS</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Breast Cancer</td>
<td>Controversial. Overall, lifetime risk is not increased. A meta-analysis of 27 studies indicates a relative risk of 1.16 after 4–12 yr of use. The relative risk is increased to 3 if started in teenage years and duration is &gt;10 yr.</td>
<td>Further information required regarding risk with progestin-only contraceptives.</td>
</tr>
<tr>
<td>Cerebrovascular Accidents</td>
<td>Risk of hemorrhagic stroke is increased 2.5-fold compared with nonusers; ever-users have a 1.5-fold risk compared with never-users. Risk is mostly in heavy smokers ≥35 yr and with predisposing risk factors (eg, hypertension, diabetes, hyperlipidemia). Odds ratio is 2.9 with combined oral contraceptives (OCs) containing 50 µg of estrogen, 1.8 for combined OCs with 30–40 µg estrogen, and 0.9 for progestin-only products.</td>
<td>Related to both the estrogen and progestin components. Minimal risk with 35 µg/day and progestin-only preparations.</td>
</tr>
<tr>
<td>Cervical Cancer</td>
<td>Increased risk of cervical erosions, eversions, dysplasias, and conversion to cancer in situ. Relative risk is 1.8–2.1 times that of nonusers and increased with duration of use &gt;5 yr; other risk factors include multiple sexual partners and early sexual activity.</td>
<td>May increase risk of herpes or papillomavirus infection, which accelerate progression of preinvasive lesions.</td>
</tr>
<tr>
<td>Gallbladder Disease</td>
<td>Relative risk of 1.36 for gallstones in users compared to nonusers only during the first 4 yr of use, then risk returns to baseline.</td>
<td>Estrogens increase cholesterol saturation.</td>
</tr>
<tr>
<td>Hepatic Tumors</td>
<td>Both benign and malignant tumors reported. Relative risk is 2.6 for users; 9.6 with duration of use &gt;5 yr. Shock can result from rupture of mass. Surgical intervention may be needed, because tumors are not always reversible after discontinuation. Risk is greater in smokers and those with a history of hepatitis B infection or diabetes.</td>
<td>Unknown, although mestranol and higher-dosage formulations are implicated. Progestin-only contraceptives not implicated.</td>
</tr>
</tbody>
</table>

(continued)
<table>
<thead>
<tr>
<th>CONDITION</th>
<th>CLINICAL INFORMATION</th>
<th>COMMENTS</th>
</tr>
</thead>
<tbody>
<tr>
<td>Hyperglycemia</td>
<td>Abnormal glucose tolerance found in predisposed individuals (e.g., subclinical or gestational diabetes) and rare cases of diabetic ketoacidosis reported. These effects are minimal with combinations containing ≤35 µg/day of ethinyl estradiol or newer progestins. Norgestrel has greatest insulin-antagonizing activity.</td>
<td>Hyperinsulinemia with relative insulin resistance caused by progestins with minimal effect from estrogens.</td>
</tr>
<tr>
<td>Hyperlipidemia</td>
<td>Elevated triglycerides; can precipitate pancreatitis in patients with underlying hyperlipidemia; adverse effects on lipids are greatest with progestin-dominant products, especially levonorgestrel and ethynodiol diacetate, and lowest with norgestimate and desogestrel.</td>
<td>Estrogens increase triglycerides and HDL; progestins increase LDL and decrease HDL. Minimal effect with progestin-only products.</td>
</tr>
<tr>
<td>Hypertension</td>
<td>Mild BP elevations of 4 mm Hg systolic and 1 mm Hg diastolic, usually reversible upon drug discontinuation, occur in 1–5% of users. Rare with low-dose products. More common in older women and in those with a family history of hypertension.</td>
<td>Related to both estrogen and progestin components. Consider progestin-only contraceptives.</td>
</tr>
<tr>
<td>Infertility</td>
<td>Little risk of permanent sterility. Conception rate after discontinuation may temporarily lag behind that of nonusers for a few months.</td>
<td>Risk concentrated in older women with a long history of contraceptive use.</td>
</tr>
<tr>
<td>Myocardial Infarction</td>
<td>No increased risk in healthy nonsmokers; risk is increased 2.8 times that of nonusers in smokers ≥35 yr with presence of other predisposing factors (e.g., hyperlipidemia, diabetes, hypertension). Relative risk of 1.9 for current and past users of low-dose products.</td>
<td>Questionably thromboembolic because risk reverses after drug discontinuation.</td>
</tr>
</tbody>
</table>
## RISKS AND BENEFITS OF ORAL CONTRACEPTIVES COMPARISON CHART (continued)

<table>
<thead>
<tr>
<th>CONDITION</th>
<th>CLINICAL INFORMATION</th>
<th>COMMENTS</th>
</tr>
</thead>
<tbody>
<tr>
<td>Postpill Amenorrhea</td>
<td>Prevalence is 0.2–2.6% after use; check for pituitary tumor in presence of galactorrhea.</td>
<td>Risk is increased if menses were irregular prior to starting. Unrelated to duration or dose.</td>
</tr>
<tr>
<td>Pulmonary Embolism</td>
<td>Risk or fatal pulmonary embolism is 9.6-fold that of nonusers.</td>
<td>Risk appears related to progestin. Cyproterone, desogestrel and gestodene carry a 2- to 3-fold greater risk than levonorgestrel.</td>
</tr>
<tr>
<td>Thromboembolism and Thrombophlebitis</td>
<td>Risk is increased 2.8-fold that of nonuser; risk is greatest in smokers, sedentary females &gt;50 yr, those with hypertension, and duration of use &gt;5 yr. Desogestrel-containing products have a 2-fold risk compared with other progestins and 4- to 5-fold that of nonusers. Minimal risk with progestin-only products.</td>
<td>Related to desogestrel and to estrogen dose. Estrogens decrease antithrombin III and increase coagulation factors and platelet aggregation. A history of venous thrombosis might be a reason to avoid combination products. Factor V Leiden is also a risk factor.</td>
</tr>
<tr>
<td>Teratogenesis</td>
<td>No increased risk of congenital cardiac, limb, or other malformations if oral or progestin-only contraceptives taken during pregnancy. Reports of masculinization of female genitalia reported when high doses of progestin were used for threatened abortion.</td>
<td>Exhaustive review of 18 prospective studies and meta-analysis of 12 prospective cohorts show relative risk of 0.99–1.04.</td>
</tr>
</tbody>
</table>

### BENEFITSa

<table>
<thead>
<tr>
<th>Condition</th>
<th>Benefits</th>
<th>Comments</th>
</tr>
</thead>
<tbody>
<tr>
<td>CONDITION</td>
<td>CLINICAL INFORMATION</td>
<td>COMMENTS</td>
</tr>
<tr>
<td>---------------------------------</td>
<td>---------------------------------------------------------------------------------------</td>
<td>-----------------------------------------------</td>
</tr>
<tr>
<td>Ovarian Cancer</td>
<td>A 30% risk reduction with duration of use ≤4 yr, 60% risk reduction with &gt;5 yr, 80% risk reduction with &gt;12 yr of use compared to nonusers. Protection persists for 10 yr after drug discontinuation.</td>
<td>Mechanism unknown.</td>
</tr>
<tr>
<td>Pelvic Inflammatory Disease/Ectopic Pregnancy</td>
<td>Risk reduction of 50–70% with &gt;1 yr of use and beneficial reduction of ectopic pregnancy rate.</td>
<td>Does not protect against gonorrhea or chlamydial cervicitis.</td>
</tr>
<tr>
<td>Menstrual Cycle Effects</td>
<td>A 90% improvement in dysmenorrhea and 50% reduced risk of iron deficiency anemia. Reduction in premenstrual symptoms (eg, anxiety, depression, and headache).</td>
<td>Decrease in menstrual flow and menstrual fluid prostaglandins.</td>
</tr>
<tr>
<td>Acne</td>
<td>Combined oral contraceptives lower serum testosterone levels with improvement of acne.</td>
<td>Use least androgenic progestins (eg, desogestrel, norgestimate) or antiandrogen (ie, drospirenone) for greatest effect.</td>
</tr>
<tr>
<td>Rheumatoid Arthritis</td>
<td>A 50% reduction in frequency.</td>
<td>Progesterone attenuates immune response.</td>
</tr>
</tbody>
</table>

*Most risks and benefits have been documented with the higher-dose estrogen products (>50 µg/day). From references 76, 78, 95–100, and 112–118.*
**HORMONE EXCESS AND DEFICIENCY SYMPTOMATOLOGY COMPARISON CHART**

<table>
<thead>
<tr>
<th>CONDITION</th>
<th>SYMPTOMATOLOGY</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Estrogen</strong></td>
<td><strong>Excess</strong>&lt;sup&gt;a&lt;/sup&gt; Estrogen excess also can be a result of progestin deficiency. Symptoms include nausea, vomiting, vertigo, leukorrhea, increase in leiomyoma size, uterine cramps, breast tenderness with fluid retention, cystic breast changes, cholasma, edema, and fluid retention resulting in abdominal or leg pain with cyclic weight gain, headaches on pill days, and hypertension.</td>
</tr>
<tr>
<td><strong>Deficiency</strong></td>
<td>Estrogen deficiency also can be a result of progestin excess. Symptoms include irritability, nervousness, decreased libido, hot flashes, early and midcycle breakthrough bleeding and spotting (days 1–7), atrophic vaginitis, dyspareunia, no withdrawal bleeding with continued contraceptive use, and decreased amount of withdrawal bleeding.</td>
</tr>
<tr>
<td><strong>Progestin</strong></td>
<td><strong>Excess</strong> Progestin excess also can be a result of estrogen deficiency. Symptoms include increased appetite and weight gain on nonpill days, tiredness, fatigue, weakness, depression, decreased libido, decreased length of menstrual flow, Candida vaginitis, headaches on nonpill days, and breast tenderness on nonpill days.</td>
</tr>
<tr>
<td><strong>Deficiency</strong></td>
<td>Progestin deficiency also can be a result of estrogen excess. Symptoms include late breakthrough bleeding (days 8–21), heavy menstrual flow and clots, dysmenorrhea, and delayed onset of menses following last pill.</td>
</tr>
<tr>
<td><strong>Androgen</strong></td>
<td>Excess Symptoms include increased appetite and weight gain, oily scalp, acne, and hirsutism.</td>
</tr>
</tbody>
</table>

<sup>a</sup>Less likely with preparations containing <50 µg/day ethinyl estradiol. From references 76 and 96–98.

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**Female Sex Hormones**

**ESTRADIOL AND ITS ESTERS** Alora, Climara, Combi Patch, Delestrogen, Estinyl, Estrace, Estraderm, Estring, Vagifem, Vivelle, Various

**Pharmacology.** Estradiol (17β-estradiol; E2) is the most potent of the naturally occurring estrogens and the major estrogen secreted during the reproductive years. Estradiol and other estrogens produce characteristic effects on specific tissues (such as breast), cause proliferation of vaginal and uterine mucosa, increase calcium deposition in bone, and accelerate epiphyseal closure after initial growth stimulation. Addition of the ethinyl radical results in an orally active compound that is 200 times more potent than estradiol. (See Notes.)

**Administration and Adult Dosage.** For patients with an intact uterus, continuous daily or monthly (at least 10–12 days) administration of a progestin is recommended to induce endometrial sloughing and decrease the risk of endometrial cancer; administration of progestin quarterly (14 days of progestin q 3 months) also might be effective. PO for postmenopausal symptoms and atrophic...
**vaginitis** administer daily or, if uterus is present, continuous daily or cyclic regimen of 3 weeks on followed by 1 week off, using the smallest effective dosage; (micronized estradiol) 0.5–2 mg/day initially, adjusted as necessary to control symptoms; (ethinyl estradiol) 0.02 mg/day or every other day, to a maximum of 0.05 mg/day; severe cases may require 0.05 mg tid initially until improvement, then decrease to 0.05 mg/day; administer as with micronized estradiol; (micronized estradiol plus norgestimate) 1 tablet daily per packaging (see Dosage Forms) (Ortho-Prefest); (ethinyl estradiol plus norethindrone) 1 tablet daily and re-evaluate at 3–6 months (femhrt 1/5). **Top patch for postmenopausal symptoms or osteoporosis** initiate with a 25 or 50 μg/day patch; patch is changed once (Climara) or twice (Estraderm, Vivelle) weekly and administered continuously or cyclically (eg, for 3 weeks followed by 1 week without patch). Dosage can be increased if symptoms are not controlled. Combi Patch can be used continuously or sequentially, in which a 50 μg/day estradiol-only patch is used for the first 14 days and Combi Patch is used for the second 14 days of a 28-day cycle. Start either method with the 0.14 mg norethindrone patch and change the patch twice weekly. (See Notes.) **Vag for postmenopausal vasomotor symptoms and atrophic vaginitis** (micronized estradiol cream) 200–400 μg/day for 1–2 weeks, then reduce to 100–200 μg/day for 1–2 weeks, then to maintenance of 100 μg 1–3 times/week; (estradiol hemihydrate vaginal tablet) 1 tablet vaginally daily for 2 weeks, then 1 tablet vaginally twice weekly (Vagifem). **Vag for symptoms of postmenopausal urogenital atrophy** (Estring) insert one 2 mg ring into the upper vagina q 3 months. **PO for prevention of osteoporosis** use minimum effective dosage of 2 mg/day micronized estradiol; 20 μg/day of ethinyl estradiol or equivalent; or ethinyl estradiol 5 μg/day plus 1 mg norethindrone acetate (femhrt 1/5). **PO for dysfunctional uterine bleeding** 0.05–0.1 mg/day of micronized estradiol or 10–20 μg/day of ethinyl estradiol for 10–20 days with addition of progestin the third week. **PO for palliation of breast cancer in postmenopausal women** (ethinyl estradiol) 1 mg tid, or (micronized estradiol) 10 mg tid for at least 3 months. **PO for palliation of advanced inoperable prostatic cancer** (ethinyl estradiol) 0.15–2 mg/day, or (micronized estradiol) 1–2 mg tid. **IM for postmenopausal symptoms and prevention of osteoporosis** when oral or vaginal therapy does not provide expected response, is poorly tolerated, or when noncompliance occurs (estradiol cypionate) 1–5 mg q 3–4 weeks; (estradiol valerate) 10–20 mg q 4 weeks. **IM for dysfunctional uterine bleeding** (estradiol valerate) 20 mg initially, then 5 mg q 2 weeks with addition of progestin. **IM for palliation of advanced inoperable prostatic cancer** (polyestradiol phosphate) 40 mg q 2–4 weeks; (estradiol valerate) 30 mg or more q 1–2 weeks depending on patient response.

**Special Populations. Geriatric Dosage.** Same as adult dosage.

**Other Conditions.** Because estrogens can increase the risk of postsurgery thromboembolic complications, discontinue estrogens at least 4 weeks before surgery, if feasible.

**Dosage Forms. Tab** (micronized estradiol) 0.5, 1, 1.5, 2 mg; (ethinyl estradiol) 0.02, 0.05, 0.5 mg; **Tab** micronized estradiol 1 mg plus norethindrone acetate 0.5 mg (Activella); micronized estradiol 1 mg 3 tablets followed by micronized
estradiol 1 mg plus norgestimate 1/9262; ethinyl estradiol 5/9262 plus norethindrone acetate 1 mg (femhrt 1/5); SR Patch (estradiol) 25, 37.5, 50, 75, 100/9262; SR Patch (estradiol) 50/9262 plus norethindrone acetate 140 or 250/9262; Vag Crm (estradiol) 100/9262; Vag Ring (estradiol) 2 mg (Estring); Inj (estradiol cypionate in oil) 5 mg/mL; 2 mg/mL with testosterone cypionate 50 mg/mL (DepoTestadiol, various); (estradiol valerate in oil) 10, 20, 40 mg/mL; 2 mg/mL with testosterone enanthate 90 mg/mL.

Patient Instructions. Report immediately if any of the following occur: new severe or persistent headache or vomiting; blurred or lost vision; speech impairment; calf, chest, or abdominal pain; weakness or numbness of extremities; or any abnormal vaginal bleeding. This (oral) drug may be taken with food, milk, or an antacid to minimize stomach upset. (Patch) discard the protective liner and apply the patch to a clean, dry, and intact area of skin, preferably on the abdomen. Avoid excessively hairy, oily, or irritated areas. Apply immediately after opening and press the patch firmly in place with the palm of your hand for about 10 seconds to ensure good contact, particularly around the edges. Do not apply to the breasts or the waistline. To minimize irritation, rotate sites with an interval of at least 1 week between applications to a particular site.

Pharmacokinetics. Onset and Duration. (Menopausal symptoms) onset of therapeutic E2 levels after oral or vaginal administration is 0.5–1 hr, with peak levels at 5 hr and progressive decline toward baseline by 12–24 hr. Onset of relief of menopausal symptoms occurs within days of the first cycle of therapy. Reductions of LH and FSH levels occur within 3 hr and 6 hr, respectively, with a duration of 24 hr. Peak E2 levels after IM products are (valerate) 2.2 days, (cypionate) 4 days. Duration of depot products is variable after IM injection; (valerate) 14–21 days, (cypionate) 14–28 days, (polyestradiol phosphate) 14–28 days. (Cancer) response to estradiol therapy should be apparent within 3 months after initiation of oral therapy.

Serum Levels. (Relief of menopausal symptoms) E2 levels: apparent at >40 ng/L (147 pmol/L); 80% relief with 68 ng/L (250 pmol/L); 100% relief with 112 ng/L (411 pmol/L). (Prevention of osteoporosis) 60 ng/L (220 pmol/L).

Fate. (Ethinyl estradiol) PO administration of 20/9262 g yields ethinyl estradiol levels of 25 ng/L (84 pmol/L); 30/9262 g yields 60 ng/L (202 pmol/L). (See Combination Oral Contraceptives.)

(Estradiol) oral bioavailability of micronized estradiol (E2) is 4.9 ± 5% because of extensive and rapid first-pass metabolism. Topical absorption is affected by skin thickness and site of patch application: 100% (abdomen) and 85% (thigh). Oral or vaginal administration results in unphysiologic levels of estrone (E1 > E2; E1 is less after Vag than PO administration). Patch yields levels of E2 > E1 (minor E1 elevations). Steady-state E2 level after PO administration of 1 mg estradiol is 35 ± 5 ng/L (128 ± 18 pmol/L) or an increase of 25 ng/L (92 pmol/L) over baseline; after 2 mg, 63 ± 11 ng/L (231 ± 40 pmol/L) or 40 ng/L (147 pmol/L) over baseline; after 4 mg, 121 ± 15 ng/L (444 pmol/L) over baseline.
± 55 pmol/L) or 50 ng/L (183 pmol/L) over baseline; after 6 mg, 207 ± 200 ng/L (760 ± 734 pmol/L), (Vag) 0.2 mg estradiol yields 80 ± 19 ng/L (293 ± 7 pmol/L) of E₂; (Patch) 25 µg yields 25 ng/L (92 pmol/L); 50 µg yields 38 ± 10 ng/L (138 ± 36 pmol/L); 100 µg yields 89 ± 82 ng/L (327 ± 302 pmol/L) of E₂. Estradiol is about 60% bound to albumin, 38% to sex hormone-binding globulin, and 3% unbound. It is widely distributed and concentrated in fat. Vₐ is 10.9 ± 2.9 L; Cl is 24.2 ± 7 L/hr/m² or 0.77 L/hr/kg. Estradiol and its esters are converted in the liver, endometrium, and intestine, 15% to estrone (active), 65% to estrone sulfate and its conjugates (primarily sulfates and glucuronides with reconversions of 5% estrone and 1.4% estrone sulfate back to E₂). E₂ is excreted 50% in urine and 10% in feces, with some enterohepatic circulation. Less than 1% is excreted unchanged in urine and 50–80% as conjugates: estrone 20%, estradiol 20%, estradiol glucuronide 7%. (Estradiol valerate and cypionate) these are slowly hydrolyzed to E₂ and their respective free acids. (Polyestradiol phosphate) slowly hydrolyzed to E₂. t₁/₂ (Estradiol) 1 hr; (ethinyl estradiol) 15 ± 3 to 33 ± 10 hr.

**Adverse Reactions.** (See Postmenopausal Hormone Replacement Risks and Benefits Comparison Chart.) Nausea, vomiting, bloating, breast tenderness, and spotting occur frequently. (See Hormone Excess and Deficiency Symptomatology Comparison Chart.) Hypercalcemia occurs occasionally in patients with breast cancer. Thromboembolism, thrombophlebitis, diabetes, hypertension, and gallbladder disease are less likely to occur with hormone replacement dosages than with oral contraceptive dosages. Pain at injection site occurs frequently. Occasional redness and irritation at application site with patch; rash rarely.

**Contraindications.** Pregnancy; history or presence of estrogen-dependent cancer (except in appropriate patients treated for metastatic disease); undiagnosed abnormal genital bleeding; history or presence of thromboembolism or severe thrombophlebitis. A history of breast cancer might not be an absolute contraindication to estrogen therapy in women with severe menopausal symptoms. Active or severe chronic liver disease is a contraindication for combinations with testosterone.

**Precautions.** Use with caution in patients with disease states that could be exacerbated by increased fluid retention (eg, asthma; epilepsy; migraine; and cardiac, hepatic, or renal dysfunction); in women with strong family histories of breast cancer or presence of fibrocystic disease, fibroadenoma, or abnormal mammogram; in women with fibromyomata, cardiovascular disease, diabetes, hypertriglyceridemia, severe liver disease, or history of jaundice during pregnancy; and in young patients in whom bone growth is not complete. Oral estrogen can increase thyroid-binding globulin and cause false elevations in total T₄ and T₃ and false depression of resin T₃ uptake while the thyroid index, thyroid-stimulating hormone, and the patient remain euthyroid. Estrace 2 mg and Estinyl 0.02 mg contain tartrazine, which may cause allergic reactions, including bronchospasm, in susceptible individuals.

**Drug Interactions.** Estrogens can reduce the effects of tricyclic antidepressants and warfarin and increase the effects of corticosteroids by increasing their half-
lives. Barbiturates, rifampin, and other cytochrome P450 inducers can decrease estrogen levels.

**Parameters to Monitor.** Signs and symptoms of side effects, especially abnormal bleeding. Pretreatment and physical examination with reference to blood pressure, breasts, abdomen, pelvic organs, and Pap smear. Baseline laboratory tests should include glucose, triglycerides, cholesterol, LFTs, and calcium. Repeat physical examination annually; repeat laboratory tests only if abnormal at baseline.

**Notes.** Estradiol has been advocated as the estrogen replacement of choice because it is the principal estrogen of the reproductive years; however, advantages over other estrogens have not been established. Synthetic 17α-alkylated estrogens (eg, ethinyl estradiol) are generally not recommended in menopausal replacement therapy because of their potent hepatic effects. The combination of an androgen with estrogen is indicated for moderate to severe vasomotor symptoms in patients not improved by estrogen alone. Potential benefits include increased libido and psychological well-being. An alternative to estrogens for hot flashes is mestrol acetate 20 mg bid, which reduced hot flashes by 50% during 4 weeks of use in one study. The combination of norethindrone acetate and estradiol in a single patch (Combi Patch) results in less endometrial hyperplasia than an estradiol-only patch.

Nonoral estradiol administration (eg, patch, vaginal, implant, injection), avoids first-pass effect and theoretically results in a preferable premenopausal physiologic serum level ratio of $E_2 > E_1$. Oral administration results in an unphysiologic ratio of $E_2 < E_1$ ($E_3$ levels are not directly related to efficacy). Avoiding the first-pass effect allows a smaller dosage to be used and prevents undesirable changes from liver stimulation (ie, increases in renin substrate, sex hormone-binding globulin, thyroxine-binding globulin, coagulation factors, transferrin, growth hormone levels, and cortisol-binding globulin and a reduction in insulin-like growth factor) and their sequelae (ie, gallbladder disease, hypertension, and hypercoagulable states in some women). Hepatic stimulation varies with oral preparations, with ethinyl estradiol > conjugated estrogens > $E_2$. Enhanced liver action is also responsible for the cardioprotective effects on lipids and occurs even with vaginal estrogens. Transdermal administration appears to exert favorable effects on serum lipoproteins (ie, elevation of HDLs and depression of LDLs) after >4 months of use and protects against bone loss and fractures similarly to oral estrogens.

Postmenopausal women most likely to develop osteoporosis are whites and Asians; blacks are at less risk. Numerous estrogens and other drugs are available for the prevention and treatment of postmenopausal osteoporosis. In women in whom estrogen replacement therapy is intolerable or contraindicated, oral bisphosphonates have increased bone mass and reduced vertebral fractures, vertebral deformities, and loss of height. (See Alendronate). **Calcitonin salmon** (Miacalcin) 200 IU/day intranasally has increased bone mass in women >5 yr postmenopausal with low bone mass who cannot take estrogens. Slow-release fluoride (Slow Fluoride) appears to be useful in a dosage of 25 mg/day for up to 4 yr, but immediate-release products are not useful because the drug is irritating to the GI tract and the new bone formed is brittle and subject to fracture.
Pharmacology. Conjugated estrogens contain a mixture of 50–65% sodium estrone sulfate, 20–35% sodium equilin sulfate, and other estrogenic substances obtained from the urine of pregnant mares. Esterified estrogens are a combination of 75–85% sodium estrone sulfate and 6.5–15% sodium equilin sulfate prepared from Mexican yams. (See Estradiol and Its Esters.)

Administration and Adult Dosage. For patients with intact uteri, continuous daily or monthly (for at least 10–12 days) administration of a progestin is recommended to induce endometrial sloughing and decrease the risk of endometrial cancer; administration of progestin quarterly (14 days of progestin q 3 months) also might be effective.119–121 PO for postmenopausal symptoms and atrophic vaginitis use smallest effective dosage in the range of 0.3–1.25 mg/day continuously or, if uterus is present, in cycles of 21–25 days/month. PO for prevention of postmenopausal osteoporosis use minimum effective dosage of 0.625 mg/day continuously, or cyclically if uterus is present, or 0.3 mg/day if 1.5 g/day of elemental calcium is also used; higher dosages of 1.25 mg/day may be necessary after fractures caused by osteoporosis.119,133 For women experiencing migraine or other symptoms during the withdrawal period, a 5-day/week regimen or a shorter withdrawal period may be used. Vag for postmenopausal symptoms and/or atrophic vaginitis 1.25–2.5 mg/day; (atrophic vaginitis) 0.3 mg 3 times/week might be effective.124 IV (preferred) or IM for rapid cessation of dysfunctional uterine bleeding 25 mg of conjugated estrogens, may repeat in 6–12 hr prn, to a maximum of 3 doses.79 IV for bleeding from uremia 0.6 mg/kg/day diluted in 50 mL of NS and infused over 30–40 min for 5 days; dosages as high as 60 mg/day IV have been used.136,137 PO for palliation of breast cancer (patients should be ≥5 yr postmenopausal) 10 mg tid. PO for palliation of prostatic cancer 1.25–2.5 mg tid.

Special Populations. Geriatric Dosage. Same as adult dosage.

Dosage Forms. Tab (conjugated) 0.3, 0.625, 0.9, 1.25, 2.5 mg (Cenestin, Premarin, various); 0.625 mg with medroxyprogesterone acetate 2.5, 5 mg (Prempro); 0.625 mg with medroxyprogesterone acetate 5 mg (Premphase); (esterified) 0.3, 0.625, 1.25, 2.5 mg; 0.625 mg with testosterone 1.25 mg (Estratest H.S.); 1.25 mg with testosterone 2.5 mg (Estratest); Inj (conjugated) 25 mg; Vag Crm (conjugated) 0.625 mg/g.

Patient Instructions. Report immediately if any of the following occur: new severe or persistent headache or vomiting; blurred or loss of vision; speech impairment; calf, chest, or abdominal pain; weakness or numbness of extremities; or any abnormal vaginal bleeding. This (oral) drug may be taken with food, milk, or an antacid to minimize stomach upset.

Pharmacokinetics. Onset and Duration. (Menopausal symptoms) PO peak onset of equilin sulfate is 4 hr; onset of estrone is 3 hr, with a peak at 5 hr; duration is >24 hr. After vaginal administration, onset of therapeutic estradiol levels is 3 hr.
and peak occurs in 6 hr, with decline over 24 hr to baseline values. Gonadotropin suppression occurs within 1 month of therapy, although suppression to premenopausal levels might not occur. Uremia improvement in bleeding time occurs within 6 hr after starting estrogens; maximum improvement occurs within 2–5 days after initiation of estrogens; effects last 3–10 days after drug discontinuation.126

**Serum Levels.** (See Estradiol and Its Esters.)

**Fate.** Conjugated equilin and estrone sulfate are rapidly absorbed and hydrolyzed to unconjugated forms when given orally or vaginally. Oral administration of 0.3 mg yields steady-state estradiol (E2) levels of 48 ± 12 ng/L (175 ± 45 pmol/L) or an increase of 20 ng/L (73 pmol/L) over baseline; 0.625 mg yields 103 ± 33 ng/L (378 ± 120 pmol/L) or 50 ng/L (184 pmol/L) over baseline; 1.25 mg yields 125 ± 66 ng/L (460 ± 243 pmol/L) or 70 ng/L (257 pmol/L) over baseline. Vaginal administration of 0.3 mg yields steady-state E2 levels of 7 ± 22 ng/L (26 ± 81 pmol/L); 0.625 mg yields 36 ± 16 ng/L (131 ± 57 pmol/L); 1.25 mg yields 94 ± 44 ng/L (344 ± 161 pmol/L).82,122,123 (Estrone sulfate) Vd is 38 ± 13 L; Cl is 3.9 ± 1.2 L/hr/m².138,139 Estrone sulfate is rapidly converted to estrone and estradiol. (Equilin sulfate) Cl is 7.3 ± 4 L/hr/m². Approximately 30% of equilin sulfate is metabolized to active 17α-dihydroequilin sulfate and 2% to active 17α-dihydroequilin.138 Inactivation of estrogens occurs mainly in the liver, with degradation to less active estrogenic products (eg, estrone). Metabolites are conjugated with sulfate and glucuronic acid; urinary recovery is 70–88% within 5 days after oral administration. (See Estradiol and Its Esters.)

\[ t_{1/2} \] (Estrone sulfate) 4–5 hr. (Equilin) 19–27 min. (Equilin sulfate) 190 min. (17α-dihydroequilin) 45 ± 5 min. (17α-dihydroequilin sulfate) 2.5 ± 0.6 hr.138

**Adverse Reactions, Contraindications, Precautions, Drug Interactions, Parameters to Monitor.** (See Estradiol and Its Esters.)

**Notes.** Oral and vaginal administrations result in an unphysiologic E1 > E2 ratio, although higher E2 levels occur orally than vaginally.122–125 (See Estradiol Notes, Postmenopausal Hormone Replacement Risks and Benefits Comparison Chart.)

**ESTROPIPATE**

**Pharmacology.** Estropipate is estrone sulfate stabilized with inert piperazine. Estrone (E1) is the major estrogen produced in the postmenopausal period. It is one-half as potent as estradiol (E2) and shares the actions of other estrogens. (See Estradiol and Its Esters.)

**Administration and Adult Dosage.** For patients with intact uteri, continuous daily or monthly administration (minimum of 10–12 days) of progestin is recommended to induce endometrial sloughing and decrease the risk of endometrial cancer; administration of progestin quarterly (14 days of progestin q 3 months) also might be effective.119–121 PO for postmenopausal symptoms and prevention of osteoporosis use the smallest effective dosage in the range of 0.625–5 mg/day continuously or in cycles of 21–25 days/month; administer as with conjugated estrogens. Vag for postmenopausal symptoms and/or atrophic vaginitis 3–6 mg/day. PO for palliation of inoperable advanced prostatic cancer 3–6 mg tid.
Special Populations. Geriatric Dosage. Same as adult dosage.

Dosage Forms. Tab (as conjugated estrogens equivalent) 0.625, 1.25, 2.5, 5 mg; Vag Crm 1.5 mg/g.

Patient Instructions. Report immediately if any of the following occur: new severe or persistent headache or vomiting; blurred or loss of vision; speech impairment; calf, chest, or abdominal pain; weakness or numbness of extremities; or any abnormal vaginal bleeding. This (oral) drug may be taken with food, milk, or an antacid to minimize stomach upset.

Pharmacokinetics. Estrone is not orally active because of enzymatic degradation in the gut and liver. Addition of a piperazine moiety increases oral absorption such that $E_2$ levels are similar to those after administration of estradiol. Oral administration of 0.6 mg estropipate yields $E_2$ serum levels of 34 ng/L (124 pmol/L); 1.2 mg yields 42 ng/L (154 pmol/L). Estrone is hydroxylated to $\alpha$-hydroxyestrone, estriol, and 2-hydroxyestrone. The half-life of estrone is estimated to be 12 hr in serum; however, this does not reflect events in peripheral tissues. The half-life of estrone sulfate is 4–5 hr. (See Estradiol and Its Esters.)

Adverse Reactions, Contraindications, Precautions, Drug Interactions, Parameters to Monitor, Notes. (See Estradiol and Its Esters.)
### ESTROGENS COMPARISON CHART

<table>
<thead>
<tr>
<th>DRUG</th>
<th>DOSAGE FORMS</th>
<th>EQUIPOTENT PHYSIOLOGIC DOSEa,b</th>
<th>COMMENTS</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>STEROIDAL AGENTS</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Conjugated Estrogens</strong></td>
<td>Tab 0.3, 0.625, 0.9, 1.25, 2.5 mg</td>
<td>0.625 mg.</td>
<td>Mixture of 50–65% sodium estrone sulfate, 20–35% equilin sulfate, and other estrogenic substances from the urine of pregnant mares. Expensive; nausea is rare.</td>
</tr>
<tr>
<td>Premarin</td>
<td>Vag Crm 0.625 mg/g</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Various</td>
<td>Inj 25 mg.</td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Esterified Estrogens</strong></td>
<td>Tab 0.3, 0.625, 1.25, 2.5 mg</td>
<td>0.625 mg.</td>
<td>Similar to conjugated estrogens. Mixture of 75–85% sodium estrone sulfate and 6.5–15% sodium equilin sulfate obtained from Mexican yams.</td>
</tr>
<tr>
<td>Estratab</td>
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<td>Menest</td>
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<td></td>
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<tr>
<td>Various</td>
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<tr>
<td><strong>Estradiol, Micronized</strong></td>
<td>Tab 0.5, 1, 1.5, 2 mg</td>
<td>1 mg.</td>
<td>Moderate cost; some nausea with oral; estradiol is the major estrogen secreted during the reproductive years.</td>
</tr>
<tr>
<td>Estrace</td>
<td>Vag Crm 100 mg/g</td>
<td></td>
<td></td>
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<tr>
<td></td>
<td>Vag Ring 2 mg.</td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Estradiol</strong></td>
<td>SR Patch 25, 37.5, 50, 75, 100 µg/day</td>
<td>50 µg/day.</td>
<td>Estraderm contains alcohol; Climara and Vivelle do not contain alcohol and may be less irritating to the skin.</td>
</tr>
<tr>
<td>Climara</td>
<td></td>
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<tr>
<td>Estraderm</td>
<td></td>
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<tr>
<td>Vivelle</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Estradiol Cypionate</strong></td>
<td>Inj (in oil) 5 mg/mL.</td>
<td>—</td>
<td>Pain at injection site; variable onset with a duration of 14–28 days.</td>
</tr>
<tr>
<td>Depo-Estradiol</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Various</td>
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(continued)
### ESTROGENS COMPARISON CHART (continued)

<table>
<thead>
<tr>
<th>DRUG</th>
<th>DOSAGE FORMS</th>
<th>EQUIPOTENT PHYSIOLOGIC DOSE</th>
<th>COMMENTS</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Estradiol Valerate</strong></td>
<td>Inj (in oil) 10, 20, 40 mg/mL.</td>
<td>1 mg.</td>
<td>Pain at injection site; variable onset with a duration of 14–21 days.</td>
</tr>
<tr>
<td>Delestrogen</td>
<td>Various</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Estradiol</td>
<td>Inj (in oil) 10, 20, 40 mg/mL.</td>
<td>1 mg.</td>
<td></td>
</tr>
<tr>
<td><strong>Ethinyl Estradiol</strong></td>
<td>Tab 0.02, 0.05, 0.5 mg.</td>
<td>5 µg.</td>
<td></td>
</tr>
<tr>
<td>Estroline</td>
<td>Various</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Estrone</td>
<td>Inj 2, 5 mg/mL.</td>
<td>0.9 mg.</td>
<td>No advantage over conjugated/esterified estrogens; estrone is the major estrogen of the postmenopausal years.</td>
</tr>
<tr>
<td>Feminone</td>
<td>Various</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Estratropipate</td>
<td>Tab 0.625, 1.25, 2.5, 5 mg</td>
<td>0.625 mg.</td>
<td></td>
</tr>
<tr>
<td>Ogen</td>
<td>Vag Crm 1.5 mg/g.</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Ogen</td>
<td>Ogen 1.5 mg/g.</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Ogen</td>
<td>Ogen 2.5 mg = 3 mg estropipate.</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Ogen</td>
<td>Ogen 5 mg = 6 mg estropipate.</td>
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</tbody>
</table>

### NONSTEROIDAL AGENTS

<table>
<thead>
<tr>
<th>DRUG</th>
<th>DOSAGE FORMS</th>
<th>COMMENTS</th>
</tr>
</thead>
<tbody>
<tr>
<td>Dienestrol</td>
<td>Vag Crm 0.01%</td>
<td>—</td>
</tr>
<tr>
<td>Various</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

*a Potency of estrogens: estradiol > estrone. Potency is based on the effects on the liver.

*b See monographs or product information for exact dosage regimens for various uses.
## POSTMENOPAUSAL HORMONE REPLACEMENT RISKS AND BENEFITS COMPARISON CHART

<table>
<thead>
<tr>
<th>RISKS/BENEFITS</th>
<th>CLINICAL INFORMATION</th>
<th>COMMENTS</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Cancer, Breast</strong></td>
<td>Controversial; no association with &lt;5 yr duration of use to relative risk of 1.25–1.45 among current users with &gt;5 yr duration of use; highest risk of 1.7 reported among long-term users &gt;60 yr. No risk found in past users, regardless of duration of use. Two meta-analyses show minimal risk with &gt;15 yr of use.</td>
<td>Addition of progestin does not reduce risk. Regular mammography is recommended. Consider limiting duration of treatment to &lt;5 yr if risks of cancer outweigh cardioprotective benefits.</td>
</tr>
<tr>
<td><strong>Cancer, Colon</strong></td>
<td>A 46% decrease in colon cancer risk; no effect on rectal cancer. Relative risk of 8.2 with unopposed estrogen use; risk increases with higher dosage and duration &gt;5 yr; 34% risk after 3 yr; 20% lifetime probability of needing a hysterectomy with unopposed estrogen therapy.</td>
<td>In slender women, risk is reduced by up to 75%.</td>
</tr>
<tr>
<td><strong>Cancer, Endometrial</strong></td>
<td>Relative risk of 8.2 with unopposed estrogen use; risk increases with higher dosage and duration &gt;5 yr; 34% risk after 3 yr; 20% lifetime probability of needing a hysterectomy with unopposed estrogen therapy.</td>
<td>Relative risk of 1 with the concurrent addition of a minimum of 10–14 days of progestin. No increased risk of estrogen hyperplasia or need for hysterectomy with concurrent progestin therapy.</td>
</tr>
<tr>
<td><strong>Cardiovascular Disease</strong></td>
<td>Three meta-analyses and a cohort study suggest a 40–50% reduction in the risk of coronary and fatal heart disease with unopposed estrogens; benefits may be greater in those with heart disease and &gt;15 yr duration of use. Decreased lifetime probability of developing coronary artery disease. Hormone replacement for ≥1 yr associated with a 52% decreased risk of peripheral arterial disease. Unknown protection against stroke.</td>
<td>Combination with progestin may be protective, but data are insufficient. May be related to estrogen’s effects on lipids or direct effect of relaxing blood vessel walls.</td>
</tr>
<tr>
<td><strong>Hypertension</strong></td>
<td>Estrogens can reduce BP.</td>
<td>Hormone replacement is not contraindicated in hypertension.</td>
</tr>
<tr>
<td><strong>Lipids</strong></td>
<td>Unopposed oral estrogens reduce LDL and increase HDL by 10–15%; however, estrogens can increase triglyceride levels. Progesterone antagonizes beneficial estrogen lipid effects less than medroxyprogesterone. Most favorable effects on lipids occur with estrogen alone. Nonoral estrogens (eg, patch, vaginal) produce less HDL beneficial effects.</td>
<td>(continued)</td>
</tr>
</tbody>
</table>
### POSTMENOPAUSAL HORMONE REPLACEMENT RISKS AND BENEFITS COMPARISON CHART (continued)

<table>
<thead>
<tr>
<th>RISKS/BENEFITS</th>
<th>CLINICAL INFORMATION</th>
<th>COMMENTS</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Gallbladder Disease</strong></td>
<td>Estrogen treatment is associated with a 2.1 relative risk (RR). RR of 2.6 with &gt;10 yr of use; RR of 2.4 for users of 1.25 mg or more of conjugated/estrified estrogen.</td>
<td>Mortality unaffected; may require cholecystectomy.</td>
</tr>
<tr>
<td><strong>Osteoporosis</strong></td>
<td>Inhibits bone resorption and prevents bone loss; 15–50% increase in bone density if begun within 3 yr of menopause. Osteoporosis risk increased in Caucasian and Asian ethnic groups, in sedentary lifestyle, in smokers, with low calcium and vitamin D intake, and excessive alcohol or thyroxine intake.</td>
<td>Alendronate (Fosamax) orally, intranasal calcitonin (Miacalcin), etidronate (Didronel), and slow-release fluoride also may be effective. (See Estradiol Notes.)</td>
</tr>
<tr>
<td><strong>Fractures</strong></td>
<td>One-half as many fractures of spinal and hip bones with &gt;5 yr of use; 28% reduction with 10 yr use; 40% with 15 yr use; and 55% with 20 yr. Risk returns near baseline 6 yr or more after cessation of therapy. Decreased lifetime probability of osteoporotic fracture. Risk increases 4-fold for each 1 SD decrease in bone density at the hip; 66% of femoral neck fractures occur when bone density is below the lowest quartile.</td>
<td>Bone densitometry can identify women at highest risk.</td>
</tr>
<tr>
<td><strong>Vaginal Bleeding</strong></td>
<td>Unpredictable bleeding occurs in 35–40% of women with uteruses yearly.</td>
<td>Amenorrhea usually occurs after 6–8 months of combination estrogen/progestin therapy.</td>
</tr>
</tbody>
</table>

*From references 119, 121, 127, 130, 132, 139, and 157–164.*
Pharmacology. Medroxyprogesterone is a 17α-acetoxyprogesterone derivative with greater progestational effects and oral efficacy than progesterone. Progesterone transforms an estrogen-primed proliferative endometrium into a secretory endometrium.

Administration and Adult Dosage. PO for secondary amenorrhea, or abnormal uterine bleeding, or to induce withdrawal bleeding after postmenopausal estrogen replacement therapy 5–10 mg/day for 5–10 days, depending on the degree of endometrial stimulation desired, beginning on the presumed 16th or 21st day of the cycle for abnormal uterine bleeding. In secondary amenorrhea, therapy can be started at any time. PO for postmenopausal symptoms and osteoporosis (combined with continuous estrogen) 2.5–5 mg/day.140,141 (See Notes.) PO for relief of vasomotor symptoms 20 mg/day; IM for relief of vasomotor symptoms 150 mg/day.142 (See Notes.) IM for endometrial or renal carcinoma 400 mg–1 g/week initially for a few weeks, then, if improvement occurs, reduce to maintenance dosage of 400 mg/month. (See also Progestin-Only Contraceptives.)

Special Populations. Geriatric Dosage. Same as adult dosage.

Dosage Forms. Tab 2.5, 5, 10 mg; Inj 150, 400 mg/mL.

Patient Instructions. Report immediately if any of the following occur: new severe or persistent headache; blurred vision; calf, chest, or abdominal pain; or any abnormal vaginal bleeding. This (oral) drug may be taken with food, milk, or an antacid to minimize stomach upset. (Dysfunctional uterine bleeding) expect heavy and severely cramping flow 2–4 days after stopping therapy; expect a normal period after a few days.

Pharmacokinetics. Onset and Duration. Withdrawal bleeding (in estrogen-primed endometrium) occurs 3–7 days after the last dose.103,104 Onset of symptomatic relief of hot flashes within 4–7 days; maximum relief after 1 month; duration 8–20 weeks after discontinuation.142

Serum Levels. Inhibition of ovulation and tumor response occurs with medroxyprogesterone levels >0.1 μg/L (0.25 nmol/L).101,103,104,143

Fate. Medroxyprogesterone acetate (MPA) is rapidly absorbed orally with no first-pass metabolism; oral bioavailability is 5.7 ± 3.8%; IM bioavailability is 2.5 ± 1.7%, with a large interpatient variation in serum levels after oral or IM administration.82,144 Higher concentration depot formulation is associated with lower serum concentrations but equivalent bioavailability.144 Peak concentrations occur in 2–7 hr and are 2–10 times higher after oral than after IM depot injection. PO 10 mg yields peak levels of 3–4 μg/L (7.5–10 nmol/L), declining to 0.3–0.6 μg/L (0.8–1.5 nmol/L) by 24 hr; PO 100 mg yields 13 ± 7 μg/L (34 ± 18 nmol/L), declining to 2 μg/L (5 nmol/L) by 24 hr; PO 500 mg yields 13 ± 8 μg/L (34 ± 21 nmol/L). After 150 mg IM of the 150 mg/mL formulation, peak levels of 8.3 ± 3.2 μg/L (21 ± 8 nmol/L) occur within a few days, declining to levels of 0.8 ± 0.7 μg/L (2 ± 1.8 nmol/L) for 92 ± 44 days. After 400 mg IM of the 400 mg/mL formulation, peak serum levels of 6.2 ± 2.3 μg/L (16 ± 6 nmol/L) are achieved after 16.3 ± 15.6 days.82,94,103,104,144 The drug is stored in fat; >90% is
protein bound to albumin; 83% of a dose is present in serum as the parent drug and conjugated medroxyprogesterone; it is hydroxylated to 6α-hydroxy-MPA and 21-hydroxy-MPA, which have unknown activities. From 15% to 20% of a dose is excreted in urine as glucuronide and sulfate conjugates; 45–80% is excreted in feces.94

\[ t_{1/2} \approx 50 \text{ days}, \] reflecting slow IM absorption from depot.

**Adverse Reactions.** Frequent breast tenderness, weight gain, and depression occur. Adverse lipid effects (increased LDL, decreased HDL) occur with dosages \( \geq 10 \text{ mg/day} \); dosages of 2.5–5 mg/day have negligible effects.121,140,144

(See also Progestin-Only Contraceptives, Postmenopausal Hormone Replacement Risks and Benefits Comparison Chart, and Hormone Excess and Deficiency Symptomatology Comparison Chart.)

**Contraindications.** Known or suspected pregnancy or as a diagnostic test for pregnancy. Thrombophlebitis, history of deep vein thrombophlebitis, or thromboembolic disorders; known or suspected carcinoma of the breast or endometrium, or other estrogen-dependent tumors; undiagnosed abnormal genital bleeding. Although acute liver disease, benign or malignant liver tumors, and history of cholestatic jaundice of pregnancy or jaundice with prior hormonal contraceptive use are listed as contraindications by manufacturers, liver disease is not considered by others to be a contraindication to progestin-only contraceptives.96

**Precautions.** Use with caution in patients with histories of depression, diabetes, gestational diabetes, coronary artery disease, cerebrovascular disease, hyperlipidemia, liver disease, or hypertension. Although progestins are not harmful to the fetus during the first 4 months of pregnancy; confirm a negative pregnancy test before reinjecting a woman >2 weeks late for her IM injection.96,105 Progestin-only contraceptives used during breastfeeding pose no risk to the infant,96,103–107 and they usually do not decrease breastmilk production if begun after 6 weeks postpartum.

**Drug Interactions.** Rifampin and cytochrome P450–inducing anticonvulsants can increase progestin metabolism. Long-term use of griseofulvin can increase menstrual irregularities.76,96,103,104,108

**Parameters to Monitor.** Complete pretreatment physical examination with special reference to blood pressure, breasts, abdomen, pelvic organs, and Pap smear yearly.

**Notes.** Continuous administration of low-dose progestin and estrogen combinations in postmenopausal syndrome causes amenorrhea in >50% of women and does not appear to negatively influence blood lipids when compared with cyclic therapy.121,140,141,145 Concurrent administration of estrogen with progestin for amenorrhea might be associated with less breakthrough bleeding than with progestin alone. There is no evidence that progestins are effective in preventing habitual abortion or treating threatened abortion.

**MIFEPRISTONE**

*Pharmacology.* Mifepristone (RU-486) is a synthetic steroid with antiprogestational effects.
Adult Dosage. PO for pregnancy termination through day 49 of pregnancy 600 mg as a single dose, followed in 2 days by misoprostol 200 mg PO. Patients should return on day 14 to assess efficacy of the procedure and bleeding.

Dosage Forms. Tab 200 mg.

Pharmacokinetics. Oral bioavailability is 69% with a 20 mg dose. It is 98% bound to albumin and α- acid glycoprotein. It is metabolized primarily by CYP3A4 to three major metabolites. Most of drug is eliminated in feces, with 9% of the drug and metabolites eliminated in urine. Clearance is dose dependent, with 50% eliminated between 12 and 72 hr; the remaining drug is eliminated with a half-life of 18 hr.

Adverse Reactions. Vaginal bleeding and cramping are expected effects of the drug (plus misoprostol) and occur mostly on day 3. Bleeding is generally heavier than a normal menstrual period. Other frequent effects are nausea, vomiting, diarrhea, headache, dizziness, and fatigue. Drugs that affect CYP3A4 can alter mifepristone metabolism. The metabolism of drugs metabolized by CYP3A4 might be affected.

Contraindications. Confirmed or suspected ectopic pregnancy or undiagnosed abdominal mass; IUD in place; chronic adrenal failure; concurrent long-term corticosteroid use; allergy to mifepristone, misoprostol or other prostaglandin; hemorrhagic disorder; anticoagulant therapy; inherited porphyria.

Notes. Pregnancy termination should be conducted only in a setting where a qualified physician can assess the gestational age of the fetus, diagnose ectopic pregnancies, and provide surgical intervention in case of incomplete abortion or severe bleeding (or have made plans to provide such care through others).

**NORETHINDRONE ACETATE**

**Pharmacology.** Norethindrone acetate is a 19-nortestosterone derivative that shares the actions of progestins. It has oral efficacy, greater progestational activity than progesterone, and less androgenic activity than androgens. (See also Medroxyprogesterone Acetate, Progesterone.)

**Administration and Adult Dosage.** PO for withdrawal bleeding after postmenopausal estrogen replacement therapy or combined for estrogen replacement therapy 2.5–10 mg/day starting on days 15–20 of the cycle and continuing for 5–10 days, or 0.5–1 mg/day continuously combined with estrogen. PO for amenorrhea or abnormal uterine bleeding 2.5–10 mg/day starting on day 5 and ending on day 25 of menses. In cases of secondary amenorrhea, therapy can be started at any time. PO for endometriosis 5 mg/day for 2 weeks, increasing in 2.5 mg/day increments q 2 weeks until a maintenance dosage of 15 mg/day is reached.

**Special Populations.** Geriatric Dosage. Same as adult dosage.

**Dosage Forms.** Tab 5 mg.

**Patient Instructions.** Report immediately if any of the following occur: new severe or persistent headache; blurred vision; calf, chest, or abdominal pain; or any abnormal vaginal bleeding. This (oral) drug may be taken with food, milk, or an
antacid to minimize stomach upset. (Dysfunctional uterine bleeding) expect heavy and severely cramping flow 2 to 4 days after stopping therapy; expect a normal period after a few days.

**Pharmacokinetics.** **Onset and Duration.** (Uterine bleeding) after oral administration, acute bleeding should decrease in 1–2 days and stop in 3–4 days. (Withdrawal bleeding) onset 3–7 days after last oral dose.81

**Fate.** Norethindrone acetate is rapidly and completely absorbed, with a mean bioavailability of 64 ± 16% because of first-pass metabolism.81–85,94 Norethindrone acetate is rapidly converted to norethindrone in vivo.81,85,87,90 Norethindrone is 36% bound to sex hormone-binding globulin and 61% bound to albumin. It is concentrated in body fat and endometrium; breast milk levels are 10% of maternal serum levels. Vd is 4.3 ± 9 L/kg; Cl is 0.5 ± 1.5 L/hr/kg. Over 50% is eliminated in urine and 20–40% in feces as conjugated glucuronides and sulfates; <5% of norethindrone acetate is excreted as unchanged norethindrone.81–85,90,94  

\[ t_{1/2} \text{ (Norethindrone)} = 6.4 ± 3 \text{ hr}.81–85,90,94 \]

**Adverse Reactions.** (See Medroxyprogesterone Acetate, Postmenopausal Hormone Replacement Risks and Benefits Comparison Chart, and Hormone Excess and Deficiency Symptomatology Comparison Chart.)

**Contraindications.** (See Medroxyprogesterone Acetate, Postmenopausal Hormone Replacement Risks and Benefits Comparison Chart.)

**Precautions.** (See Medroxyprogesterone Acetate, Postmenopausal Hormone Replacement Risks and Benefits Comparison Chart, and Hormone Excess and Deficiency Symptomatology Comparison Chart.)

**Drug Interactions, Parameters to Monitor, Notes.** (See Medroxyprogesterone Acetate.)

**PROGESTERONE**  
Crinone, Progestasert, Prometrium, Various

**HYDROXYPROGESTERONE CAPROATE**  
Duralutin, Various

**Pharmacology.** Progesterone is the natural hormone that induces secretory changes in the endometrium, relaxes uterine smooth muscle, and maintains pregnancy. Hydroxyprogesterone is a natural progestin with minimal progestational activity; esterification with caproic acid produces a progestational compound more potent than progesterone with a prolonged duration of activity.

**Administration and Adult Dosage.** PO to prevent endometrial hyperplasia during postmenopausal estrogen replacement therapy (micronized progesterone) 200 mg/day for 12 days of cycle.118 IM for secondary amenorrhea or dysfunctional uterine bleeding (progesterone) 5–10 mg/day for 6–8 days or (only for amenorrhea) 100–150 mg as a single dose; (hydroxyprogesterone caproate) 375 mg, may repeat in 4 weeks prn. IM for palliation of metastatic endometrial cancer (hydroxyprogesterone caproate) 500 mg–1 g 2–3 times/week. Intrauterine for contraception (progesterone) 38 mg q 12 months, releases 68 μg/day; insert at any time during menstrual cycle or within 7 days of onset of menses, immediately postabortion, or no earlier than 6 weeks postpartum if
breastfeeding; insertion and removal are done by trained personnel. **Vag for progestosterone supplementation** (progesterone) 90 mg daily. (See Notes.)

**Special Populations.** **Geriatric Dosage.** Same as adult dosage.

**Dosage Forms.** **Cap** (micronized progesterone) 100, 200 mg (Prometrium); **Inj** (progesterone in oil) 50 mg/mL; (hydroxyprogesterone caproate in oil) 125, 250 mg/mL; **Intrauterine** (progesterone) 38 mg (Progestasert); **Vag Gel** (progesterone 8%) 90 mg/applicatorful (Crinone).

**Patient Instructions.** Report immediately if any of the following occur: new severe or persistent headache; blurred vision; calf, chest, or abdominal pain; or any abnormal vaginal bleeding. (Dysfunctional uterine bleeding) expect heavy flow and severe cramping 2 to 4 days after injection; expect a normal period after a few days. (Progestasert only) you might experience increased menstrual flow, cramping, and spotting. Check the position of the strings monthly after each period or after abnormal cramping to ensure proper placement of the IUD. Contact your prescriber immediately if the strings are missing, if you miss a menstrual period, or if you have fever, pelvic pain, severe cramping, unusual vaginal bleeding, or any signs of infection.

**Pharmacokinetics.** **Onset and Duration.** (Amenorrhea) onset of withdrawal bleeding occurs 48–72 hr after last dose of IM progesterone and 2 weeks after IM hydroxyprogesterone caproate; (dysfunctional uterine bleeding) onset within 6 days of IM progesterone. Duration is 12–24 hr with oral progesterone, 9–17 days with IM hydroxyprogesterone caproate. (Contraception) onset within 24 hr after insertion of Progestasert.

**Serum Levels.** (Endometrial progestational activity [luteal phase]) 15 µg/L (48 nmol/L) of progesterone.

**Fate.** (Progesterone) bioavailability of oral progesterone is incomplete because of first-pass metabolism, with wide interpatient variations; micronized forms are somewhat better absorbed. Higher levels of progesterone and active metabolites occur after IM, vaginal, or rectal administration because first-pass effect is avoided. Serum levels of progesterone after oral and IM increase rapidly to reach luteal-phase values within 2.4 ± 1.1 hr and remain elevated for <12 hr after oral administration and 48 hr after IM administration. (PO) 100 mg micronized progesterone yields peak progesterone levels of 7 ± 3.4 µg/L (23 ± 11 nmol/L); 200 mg yields 28 ± 19 µg/L (89 ± 59 nmol/L); (IM) 100 mg yields 60 µg/L (192 nmol/L); (Vag) 200 mg bid yields 19 ± 2 µg/L (61 ± 7 nmol/L); (Vag) 400 mg once daily yields 29 ± 53 µg/L (93 ± 188 nmol/L). Oral progesterone Vd is 850 ± 265 L/kg; Cl is 19 ± 38 L/hr/kg. Progestosterone circulates 80% bound to albumin and 17% to corticosteroid-binding globulin and distributes into fat. It undergoes rapid gut and hepatic metabolism, with formation of active metabolites: 20α-dihydroprogesterone (25–50% of the progestational activity of progesterone), 17-hydroxyprogesterone, and 11-deoxycorticosterone (a potent mineralocorticoid). Hydroxyprogesterone caproate is cleaved to form 17-hydroxyprogesterone in the body; 17-hydroxyprogesterone, whether formed from progesterone or exogenously administered, is further metabolized to 11-deoxycortisol and then cortisol. Urinary excretion of progesterone is 50–60%
as 5α-pregnanediol glucuronide and other conjugated glucuronic acid or sulfate metabolites; 5–10% excreted in feces.

\[ t_{1/2} \] (Progesterone) 32.6 ± 9.3 hr.145

**Adverse Reactions.** Local reactions and swelling at the site of progesterone injection. The beneficial effects of estrogen-increased HDL levels are not reversed by progesterone.121 (See Postmenopausal Hormone Replacement Risks and Benefits Comparison Chart, Hormone Excess and Deficiency Symptomatology Comparison Chart.) (Progestasert) intermenstrual spotting and menstrual bleeding irregularities, expulsion, ectopic pregnancy, uterine perforation, pelvic inflammatory disease, cramping, and pain. Intrauterine administration of contraceptive doses of progesterone has no systemic effects.76

**Contraindications.** (See Medroxyprogesterone Acetate.) Progestasert pregnancy; active, recent, or recurrent pelvic infections, including gonorrhea or *Chlamydia* infection.

**Precautions.** (See Medroxyprogesterone Acetate, Postmenopausal Hormone Replacement Risks and Benefits Comparison Chart, and Hormone Excess and Deficiency Symptomatology Comparison Chart.) Patients allergic to peanuts should not use Prometrium.

**Drug Interactions.** (See Medroxyprogesterone Acetate.)

**Parameters to Monitor.** Complete pretreatment and annual physical examinations with special reference to blood pressure, breasts, abdomen, pelvic organs, and Pap smear.

**Notes.** Progesterone is widely used in the treatment of premenstrual syndrome; however, in double-blind, controlled trials, oral micronized and vaginal progesterone were no better than placebo.150,151

**RALOXIFENE**

**Pharmacology.** Raloxifene is a selective estrogen receptor modulator similar to tamoxifen. It acts like an estrogen in the bone and like an estrogen antagonist on the breast and uterus. Raloxifene increases bone mineral density and decreases serum LDL cholesterol levels but does not stimulate endometrial growth.152,153

**Administration and Adult Dosage.** PO for prevention of postmenopausal osteoporosis 60 mg once daily with supplemental calcium.

**Dosage Forms.** Tab 60 mg.

**Pharmacokinetics.** Oral bioavailability is 2% because of an extensive first-pass effect. It is highly bound to albumin and α1-acid glycoprotein and has a \( V_d \) of 2348 L/kg. Cl is 40–60 L/hr/kg. The drug is metabolized to glucuronide metabolites, some of which undergo enterohepatic recycling, and can be converted back to the parent drug. Metabolites are excreted primarily in feces. The half-life is about 28 hr.

**Adverse Reactions.** Hot flashes occur in 25–30% of women; leg cramps also are frequent. It increases the risk of venous thrombosis and is a teratogen.
Contraindications. Women who might become pregnant or who have a history of venous thrombotic events.

Drug Interactions. Cholestyramine (and presumably colestipol) binds raloxifene and reduces its absorption and enterohepatic recirculation. The drugs should not be coadministered. Raloxifene decreases the effect of warfarin, and PT should be monitored carefully when they are given together.

Notes. Raloxifene increases bone mineral density, decreases the risk of vertebral fracture,154 and decreases the risk of invasive breast cancer.155 It also favorably alters cardiovascular risk factors (eg, LDL-c, lipoprotein-a, HDL-c), but protection against cardiovascular disease is not established.156

Thyroid and Antithyroid Drugs

IODIDES Various

Pharmacology. Iodide inhibits the synthesis and release of thyroid hormone and preoperatively decreases the size and vascularity of the hyperplastic thyroid gland. Large doses block the uptake of radioactive iodine by the thyroid gland.

Administration and Adult Dosage. PO for hyperthyroidism, as an adjunct to antithyroid agents or for preoperative thyroidectomy preparation 100–200 mg (5 drops of saturated solution of potassium iodide [SSKI] or 10–15 drops of Lugol’s solution) q 8 hr diluted in a glass of water, milk or juice; dosages as high as 500 mg/day have been used. However, administration of smaller doses of 30–50 mg iodine and continued suppression with doses of 15–50 mg/day also may be effective in patients with mild disease.165 Use for 7–10 days before surgery. PO for thyroid storm 200 mg q 6 hr. PO for prophylaxis in radiation emergency 100 mg iodine immediately before or within 1–2 hr after exposure and daily for 3–7 days, to a maximum of 10 days after exposure.

Special Populations. Pediatric Dosage. PO for thyrotoxicosis 300 mg (6 drops SSKI) q 8 hr diluted as above. PO for prophylaxis in a radiation emergency (<1 yr) 50 mg iodine immediately before or after exposure and daily for 3–7 days, to a maximum of 10 days after exposure; (>1 yr) same as adult dosage.

Geriatric Dosage. Same as adult dosage.

Dosage Forms. Soln (SSKI) 50 mg/drop iodide (1 g/mL); (Lugol’s or strong iodine) 8 mg/drop iodide (50 mg/mL iodine plus 100 mg/mL potassium iodide); Tab 100 mg iodide (130 mg potassium iodide); EC Tab not recommended.

Patient Instructions. Dilute solution in a glass (8 fluid ounces) of liquid before taking; it may be taken with food, milk, or an antacid to minimize stomach upset. Do not use if solution turns brownish-yellow. If crystals form in the solution, they can be dissolved by warming the closed container in warm water. Dissolve tablets in one-half glass of water or milk before taking. Do not use if you are breastfeeding; advise your physician if you are pregnant. Discontinue use and report if fever, skin rash, epigastric pain, or joint swelling occur.

Pharmacokinetics. Onset and Duration. Onset 24–48 hr in hyperthyroidism; maximum effect in 10–15 days. (See Notes.)
Serum Levels. (Iodide) >50 μg/L (0.4 mmol/L) inhibits iodide binding by thyroid in hyperthyroidism; >200 μg/L (1.6 mmol/L) inhibit iodide uptake by normal thyroid.166

Fate. Iodide is well absorbed throughout the GI tract and concentrated in the thyroid, stomach, salivary glands, and breastmilk. Renal clearance is 1.8 L/hr/kg; approximately 100 μg of iodine is excreted in urine daily; fecal excretion of iodine is negligible.166

Adverse Reactions. Any adverse reaction warrants drug discontinuation. Goiter, hypothyroidism, and hyperthyroidism occur frequently in euthyroid patients with a history of a thyroid disorder.167–171 Iodism occurs with prolonged use and is indicated by metallic taste, GI upset, soreness of teeth and gums, coryza, frontal headaches, painful swelling of salivary glands, diarrhea, acneiform skin eruptions, and erythema of face and chest. Rarely, hypersensitivity occurs and is manifested by angioedema, cutaneous hemorrhages, and symptoms resembling serum sickness. (See Precautions.)

Contraindications. Pulmonary tuberculosis; pulmonary edema; multinodular goiters. 165,167–170

Precautions. Pregnancy, because fetal goiter, asphyxiation, and death can occur; lactation. Use iodides with caution in patients with untreated Hashimoto’s thyroiditis, in iodide-deficient patients, in children with cystic fibrosis, and in euthyroid patients with histories of postpartum thyroiditis, subacute thyroiditis, amiodarone or lithium-induced thyroid disease, or previously treated Graves’ disease because they can be particularly sensitive to iodide-induced hypothyroidism.167–171 Patients with nontoxic multinodular goiters might be prone to development of hyperthyroidism. Avoid iodides entirely in patients with toxic nodular goiter or toxic nodules because thyrotoxicosis can be further aggravated.165,167–170 Iodides are not recommended for use as expectorants because of their potential to induce acneiform eruptions, exacerbate existing lesions, and adversely affect the thyroid. Small-bowel lesions are associated with enteric-coated potassium-containing tablets, which can cause obstruction, hemorrhage, perforation, and possible death. This dosage form is not recommended.

Drug Interactions. Iodide prevents uptake of 131I for several weeks and delays onset of thioamide action if given before the thioamide. Lithium can potentiate the antithyroid action of iodide. Serum iodine can be elevated if potassium-sparing diuretics are taken with potassium iodide.

Parameters to Monitor. Monitor for signs of iodism (see Adverse Reactions), hypothyroidism, hyperthyroidism, and parotitis occasionally during long-term use. Monitor thyroid function tests at least q 6–12 months during long-term use in patients with family histories of thyroid disease or goiter. Monitor serum potassium frequently in patients who are taking other drugs that might affect serum potassium (eg, diuretics).

Notes. Iodide has the most rapid onset of any treatment for hyperthyroidism. In thyroid storm, iodide theoretically should be given 1 hr after the thioamide dose but should not be withheld if oral thioamides cannot be given. The therapeutic ef-
fects of iodide are variable and transient, with “escape” occurring after 10–14 days; do not use iodide alone in the therapy of hyperthyroidism. Pharmacologic amounts of iodide can be present in serum from radiographic contrast agents and vaginal douches such as povidone-iodine.

**LEVOTHYROXINE SODIUM**

**Levothroid, Levoxyl, Synthroid, Unithroid, Various**

**Pharmacology.** Levothyroxine is a synthetic hormone identical to the thyroid hormone T4. Thyroid hormones are responsible for normal growth, development, and energy metabolism.

**Administration and Adult Dosage.** PO for replacement in hypothyroidism <6 months duration full replacement dosage of 1.6–1.7 μg/kg/day initially, increasing if needed and tolerated in 25–50 μg/day increments at 6–8 week intervals to a maintenance dosage that normalizes thyroid-stimulating hormone (TSH).\(^{172-174}\) **Usual maintenance dosages** 75–100 μg/day for women and 100–150 μg/day for men. Higher mean replacement dosages are required in patients with spontaneous hypothyroidism (1.7–1.8 μg/kg/day) than in those with iatrogenic hypothyroidism after radioiodine therapy for Graves’ disease (1.5–1.6 μg/kg/day).\(^{172-175}\) Once-weekly replacement therapy can be effective.\(^{176}\) **PO for replacement of subclinical hypothyroidism** same as adult dosage. The desirability of treatment is controversial; benefits are greatest in those with TSH >10 μIU/mL, hypercholesterolemia, and subtle symptoms of hypothyroidism. Replacement reduces levels of homocysteine and may lower the risk of clinical cardiovascular disease (eg, MI, aortic atherosclerosis).\(^{177-179}\) **PO for suppression therapy of nodules** 100–150 μg/day initially, increasing, if necessary and tolerated, in 25–50 μg/day increments at 6–8 week intervals to suppress TSH to below normal, detectable limits to prevent further thyroid growth. Dosages are usually higher than those required for replacement therapy and risks must be assessed, especially in patients with cardiac disease. If no improvement after 1 yr, consider stopping therapy.\(^{180,181}\) **PO for suppression therapy of thyroid cancer after thyroidectomy** 2.11 μg/kg/day initially, increasing, if needed and tolerated, in 25–50 μg/day increments at 6–8 week intervals to a dosage of 150–250 μg/day to suppress the TSH level to undetectable levels.\(^{172-175}\) **IV for myxedema coma** 400–500 μg or 300 μg/m² to increase serum T4 levels by 3–5 μg/dL (39–65 nmol/L), then 50–100 μg/day until oral administration is possible; use smaller dosages in cardiovascular disease.\(^{182,183}\) **IM** indicated only for replacement therapy if the patient cannot take oral medication; parenteral dosage is about 80% of the oral dosage because of bioavailability differences.\(^{172,173}\)

**Special Populations. Pediatric Dosage.** PO for hypothyroidism (preterm infants and full-term neonates to 1 yr) 10–15 μg/kg/day to normalize T4 to >10 μg/dL (129 nmol/L) within 3–4 weeks; (>1 yr) 3–5 μg/kg/day (average 3.5).\(^{172,184}\) Adjust maintenance dosage on the basis of growth, development, and T4 and TSH values. Replacement by at least 24 months of age corrects short stature by age 5 yr. Syrup can be formulated from tablets, with a stability of 15 days.\(^{185}\)

**Geriatric Dosage.** PO for hypothyroidism (>50 yr) start with 25–50 μg/day initially, then increase if tolerated in 12.2–25 μg/day increments at 6–8 week inter-
vals to a maintenance dosage necessary to normalize TSH; (>65 yr) <1 µg/kg/day may be required.172–175 IV for myxedema coma (>55 yr) <500 µg initially to improve outcome, then same as adult dosage.182 Poorly compliant elderly patients (mean age 86 yr) have been maintained on a twice-weekly dosing regimen; however, this regimen might be dangerous in cardiac patients.186

Other Conditions. In patients with cardiovascular disease or severe, long-standing (>6 months) myxedema, PO 12.5–25 µg/day initially, increasing, if tolerated, at 6–8 week intervals by 12.5–25 µg/day increments to a maintenance dosage necessary to normalize TSH.172–175 In patients with cardiovascular disease, particularly angina, dosage increments should be balanced between exacerbation of angina and maintenance of euthyroidism. In some patients with severe coronary disease, incomplete control of hypothyroidism might be necessary to prevent further exacerbation of angina. During pregnancy, a 20–50% increase in dosage might be required to maintain a normal TSH level.172–175 In those with continued mood disturbances on sole T4 therapy, see Liothyronine.

Dosage Forms. Tab 25, 50, 75, 88, 100, 112, 125, 137, 150, 175, 200, 300 µg; Inj 200, 500 µg.

Patient Instructions. This medication must be taken regularly to maintain proper hormone levels in the body. Report immediately if chest pain (especially in elderly patients), palpitations, sweating, nervousness, or other signs of overactivity occur.

Missed Doses. Take any missed dose as soon as it is remembered, but if more than 1 dose is missed, do not double dosage.

Pharmacokinetics. Onset and Duration. PO onset 3–5 days; peak effect 3–4 weeks; duration after cessation of therapy 7–10 days. IV onset in myxedema coma 6–8 hr, maximum effect in 1 day. Onset of hypothyroidism in serum T4 levels 3–5 µg/dL (39–65 nmol/L).172,183 Many drugs and pathologic and physiologic states affect binding and hence can affect results of some serum level determinations.166,167,169,171,174

Fate. Oral bioavailability ranges from 74 ± 11% to 93 ± 25% and can be decreased by many factors (eg, malabsorption, concurrent food, and drugs; see Drug Interactions).167,189 A dose of 500 µg IV increases serum T4 levels by 3–5 µg/dL (39–65 nmol/L).172,183 Only 0.03% is unbound in plasma. Vd is (hypothyroid) 0.17 ± 0.22 L/kg; (euthyroid) 0.16 ± 0.09 L/kg; (hyperthyroid) 0.23 ± 0.44 L/kg. Turnover is (hypothyroid) 9.2 ± 1.7%/day; (euthyroid) 11.2 ± 1.7%/day; (hyperthyroid) 21 ± 4.9%/day. Cl is (hypothyroid) 0.0008 ± 0.0003 L/hr/kg; (euthyroid) 0.00074 ± 0.0017 L/hr/kg; (hyperthyroid) 0.002 ± 0.0007 L/hr/kg.150 About 80% is deiodinated in the body; 35% is peripherally converted to the more active T3 and 45% to inactive reverse T3.166 Another 15–20% is conjugated in the liver to form glucuronides and sulfates, which undergo enterohepatic recirculation with reabsorption or excretion in the feces.
Protein binding affects half-life (increased binding retards elimination and decreased binding increases elimination).

**Adverse Reactions.** Most are dose related and can be avoided by increasing the initial dosage slowly to the minimum effective maintenance dosage. Signs of overdosage are headache, palpitations, chest pain, heat intolerance, sweating, leg cramps, weight loss, diarrhea, vomiting, nervousness, and other symptoms of hyperthyroidism. Long-term thyroid administration that results in TSH suppression can predispose to ventricular hypertrophy, atrial fibrillation, osteoporosis, and increased fracture risk by increasing bone resorption in postmenopausal women with a history of hyperthyroidism.\(^{172-175,191}\)

**Contraindications.** Thyrotoxicosis; uncorrected adrenal insufficiency.

**Precautions.** Initiate and increase dosage with caution in patients with cardiovascular disease, the elderly, and in long-standing hypothyroidism. In myxedema coma, give a corticosteroid concurrently.\(^{183}\) The status of other metabolic diseases, including diabetes, adrenal insufficiency, hyperadrenalism, and panhypopituitarism, can be affected by changes in thyroid status.

**Drug Interactions.** Bran, fiber, cholesterol-binding resins, sodium polystyrene sulfonate, iron, aluminum-containing products, and calcium carbonate can decrease oral absorption. Phenytoin, carbamazepine, and other enzyme inducers; sertraline and possibly other serotonin reuptake inhibitors, and ritonavir can increase levothyroxine requirements.\(^{167,169,189,192}\) The action of some drugs (eg, digoxin, warfarin, insulin, sympathomimetics, theophylline) can be altered by changing thyroid status.\(^ {169,174}\)

**Parameters to Monitor.** (Adults) TSH, free T\(_4\) or free T\(_4\) index, and clinical status of the patient q 6–8 weeks initially. Monitor trough levels or obtain levels at least 10 hr after tablet ingestion to avoid transient peak effects.\(^ {187,188}\) After stabilization, monitor free T\(_4\) or free T\(_4\) index, TSH, and clinical status at 6–12 month intervals. (Children) Monitor the parameters above q 4 weeks initially and q 3–4 months after stabilization. In congenital hypothyroidism, monitor T\(_4\) because TSH can remain elevated despite adequate replacement doses.\(^ {184}\) (>50 yr) Evaluate the replacement dosage annually and adjust downward as necessary because dosage requirements decrease with age.\(^ {172-175}\)

**Notes.** Levothyroxine is the drug of choice for thyroid replacement because of purity, long half-life, and close simulation to normal physiologic hormone levels. Protect from light and moisture. Concerns about tablet potency prompted the FDA to require that all manufacturers submit a new drug application for levothyroxine by August 2001. Unithroid is the first levothyroxine tablet to obtain FDA approval. Bioequivalence is reported between Synthroid, Levothroid, and Levoxyl.\(^ {173,188,193}\) Use of adjunctive thyroid hormones for depression may be effective; T\(_3\) is used instead of T\(_4\) (see Liothyronine).\(^ {194}\) Physiologic dosages of thyroid hormones in euthyroid patients are ineffective for weight reduction, obesity, or premenstrual tension; larger dosages might result in toxicity.\(^ {175}\) (See Thyroid Replacement Products Comparison Chart.)
Liothyronine is a synthetic hormone identical to the thyroid hormone T3, which is 4 times as potent by weight as T4. (See Levothyroxine.)

**Administration and Adult Dosage.** PO for replacement in hypothyroidism <6 months in duration 25 µg/day initially, increasing, if needed and tolerated, in 12.5–25 µg/day increments at 1–2 week intervals to a maintenance dosage of 25–100 µg/day to normalize TSH. PO for severe hypothyroidism 5 µg/day initially, increasing in 5–10 µg/day increments at 1–2 week intervals until 25 µg/day is reached, then increase in 12.5–25 µg/day increments at 1–2 week intervals until euthyroid. Dividing daily dosage into 2–3 doses can prevent wide serum level fluctuations. PO for augmentation of tricyclic therapy for depression 25–50 µg daily. No data are available in combination with serotonin reuptake inhibitors. IV for myxedema coma 25–50 µg initially, then 10–12.5 µg q 4–6 hr to a minimum of 10–15 µg q 12 hr until PO administration is possible; use smaller dosages of 10–20 µg IV initially in cardiovascular disease. Some suggest that T3 is preferable in myxedema coma when impairment of T4 to T3 conversion is suspected or in cardiac disease because adverse effects will dissipate faster. Limited experience exists with IV dosages >100 µg/day. PO for T3 suppression test 75–100 µg/day in 2–3 divided doses for 7 days, then repeat 131I thyroid uptake test.

**Special Populations. Pediatric Dosage.** PO for congenital hypothyroidism 5 µg/day initially, increasing in 5 µg/day increments at 3–4 day intervals until the desired effect is obtained. Usual maintenance dosage (<1 yr) 20 µg/day; (1–3 yr) 50 µg/day; (>3 yr) 25–100 µg/day. Levothyroxine is the drug of choice in congenital hypothyroidism.

**Geriatric Dosage.** Not recommended because of greater potential for cardiotoxicity. PO if used, start at PO 5 µg/day and increase in 5 µg/day increments at 2-week intervals, if tolerated, until desired response is obtained. (See Levothyroxine.)

**Other Conditions.** Not recommended in those with cardiovascular disease but, if used, start at PO 5 µg/day and increase in 5 µg/day increments at 2-week intervals, if tolerated, until desired response is obtained. (See Levothyroxine.) For those with continued mood disturbances on sole T4 therapy, substitution of T3 5 µg bid for 50 µg of levothyroxine of the total daily T4 replacement dosage has been advocated.

**Dosage Forms.** Tab 5, 25, 50 µg; Inj 10 µg/mL.

**Patient Instructions.** This medication must be taken regularly to maintain proper hormone levels in the body. Report immediately if chest pain (especially in elderly patients), palpitations, sweating, nervousness, or other signs of overactivity occur.

**Missed Doses.** Take any missed dose as soon as it is remembered, but if more than 1 dose is missed, do not double dosage.

**Pharmacokinetics. Onset and Duration.** PO onset 1–3 days; duration after cessation of therapy 3–5 days.
Serum Levels. During T₃ replacement, T₄ is maintained at ≤10 μg/L (13 nmol/L).¹⁹⁷

Fate. Oral absorption is usually complete but can decrease in CHF. With a typical replacement dosage, T₃ has a peak of 4.5–7 μg/L (7–11 nmol/L) 1–2 hr postdose, returning to 0.88–1.6 μg/L (1.4–2.5 nmol/L) before the next dose 24 hr later.¹⁹⁷ Vₐ is (hypothyroid) 0.53 ± 0.04 L/kg; (euthyroid) 0.52 ± 0.03 L/kg; (hyperthyroid) 0.94 ± 0.07 L/kg. Turnover is (hypothyroid) 50 ± 5%/day; (euthyroid) 68 ± 11%/day; (hyperthyroid) 110 ± 22%/day. Cl is (hypothyroid) 0.012 ± 0.002 L/hr/kg; (euthyroid) 0.02 ± 0.003 L/hr/kg; (hyperthyroid) 0.043 ± 0.013 L/hr/kg.¹⁹⁰ Excreted in urine as deiodinated metabolites and their conjugates. t₁/₂. (Hypothyroid) 38 ± 6 hr; (euthyroid) 25 ± 3 hr; (hyperthyroid) 17 ± 4.7 hr.¹⁹⁰

Adverse Reactions. (See Levothyroxine.) Dose-related adverse effects are more likely and appear more rapidly than with levothyroxine because regulation of dosage is more difficult. Liothyronine and its mixtures (eg, desiccated thyroid, lio-trix) cause “unphysiologic” toxic peaks in serum T₃ levels not found during levothyroxine replacement therapy.¹⁷²,¹⁷⁴,¹⁷⁵

Contraindications. (See Levothyroxine)

Precautions. (See Levothyroxine.)

Drug Interactions. Normal serum T₃ levels are age related and can be decreased by a wide variety of pharmacologic agents (eg, amiodarone, iodinated contrast dyes, corticosteroids, propylthiouracil) or clinical circumstances (eg, malnutrition, chronic renal, hepatic, pulmonary, or cardiac disease; or acute sepsis) which impair peripheral or pituitary T₄ to T₃ conversion.¹⁶⁶,¹⁶⁷,¹⁷⁴ (See also Levothyroxine Drug Interactions.)

Parameters to Monitor. Serum TSH and T₃ levels. (See Levothyroxine.)

Notes. Liothyronine is not considered the drug of choice for replacement therapy in hypothyroidism because of its shorter half-life (necessitating more frequent administration), greater potential for cardiotoxicity, the greater difficulty of monitoring, and its greater expense.¹⁷²–¹⁷⁴ Liothyronine is the preparation of choice when thyroid supplements must be stopped before isotope scanning. After scanning, maintenance therapy with levothyroxine is recommended. The use of IV T₃ after cardiopulmonary bypass might improve postoperative recovery and cardiac function in adults, children, and infants.¹⁹⁹–²⁰¹ (See Thyroid Replacement Products Comparison Chart.)
<table>
<thead>
<tr>
<th>DRUG</th>
<th>DOSAGE FORMS</th>
<th>EQUIVALENT DOSAGE</th>
<th>CONTENTS</th>
<th>RELATIVE ONSET AND DURATION(^a)</th>
<th>COMMENTS</th>
</tr>
</thead>
<tbody>
<tr>
<td>Levothyroxine</td>
<td>Tab 25, 50, 75, 88, 100, 112, 125, 137, 150, 175, 200, 300 µg</td>
<td>60 µg</td>
<td>T(_4)</td>
<td>Long</td>
<td>Preparation of choice. T(_4) content is now standardized using HPLC, and bioequivalence among products is likely.</td>
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<tr>
<td>Liothyronine</td>
<td>Tab 5, 25, 50 µg</td>
<td>25 µg</td>
<td>T(_3)</td>
<td>Short</td>
<td>Expensive; difficult to monitor. Preparation of choice if thyroid supplements are to be stopped for isotope scanning.</td>
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<tr>
<td></td>
<td>Inj 10 µg/mL</td>
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<tr>
<td>Liotrix</td>
<td>Tab 1/4, 1/2, 1, 2, 3, 4</td>
<td>#1 Tab(^b)</td>
<td>T(_4) and T(_3) in 4:1 ratio</td>
<td>Intermediate</td>
<td>No advantages; more costly and suffers from T(_3) content. (See Thyroid, Desiccated.)</td>
</tr>
<tr>
<td></td>
<td>Thyrolar</td>
<td></td>
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<td></td>
</tr>
<tr>
<td>Thyroid, Desiccated</td>
<td>Tab 15, 30, 60, 90, 120, 180, 240, 300 mg</td>
<td>60 mg</td>
<td>T(_4) and T(_3) in variable ratio</td>
<td>Intermediate</td>
<td>Inexpensive; allergy to animal protein rarely occurs; supra-physiologic elevations in T(_3) and T(_3) toxicosis may occur.</td>
</tr>
<tr>
<td>Various</td>
<td></td>
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</table>

\(^a\)With equivalent dosages.

\(^b\)Numbers represent equivalent dosage of thyroid in grains (ie, 15, 30, 60, 120, 180 mg, respectively).

\(^c\)Thyrolar-1 contains T\(_4\) 50 µg and T\(_3\) 12.5 µg; other strengths are in the same proportion.

From references 172–174.
Methimazole is a thioamide antithyroid drug that interferes with the synthesis of thyroid hormones by inhibiting iodide organification. Unlike propylthiouracil (PTU), methimazole does not block peripheral conversion of T₄ to T₃. Titers of thyroid receptor–stimulating antibody (TRab) decline during therapy, suggesting an immunosuppressive effect. Methimazole is 10 times more potent than PTU on a weight basis.

Administration and Adult Dosage. PO for hyperthyroidism 30–40 mg/day as a single dose. If GI intolerance occurs, divide dosage q 8 hr initially until euthyroid (usually 6–8 weeks), then decrease by 33–50% over several weeks to a maintenance dosage of 5–15 mg/day in a single dose. Severe disease might require 2 divided doses. The addition of levothyroxine is not recommended because remission rates have not shown improvement.²⁰² PO for thyroid storm 40–120 mg/day, divided q 8 hr until euthyroid. Traditional treatment duration for hyperthyroidism is 1–2 yr, although shorter courses of 8 months might be effective in mild disease.¹⁶⁵ Treatment may be continued indefinitely, if necessary, to control the disease and if no toxicity occurs. PR methimazole can be formulated for rectal administration.²⁰³

Special Populations. Pediatric Dosage. PO 0.5-0.7 mg/kg/day or 15–20 mg/m²/day, to a maximum of 30–60 mg/day given in 1–2 divided doses, with a maintenance dosage of 50% of the initial dosage.²⁰⁴

Geriatric Dosage. Same as adult dosage.

Other Conditions. In pregnancy, dosages should be as low as possible to maintain maternal T₄ levels in approximately the upper normal to mildly thyrotoxic range. Initially give a maximum of 20–30 mg/day orally in single or 3 divided doses for 4–6 weeks, then decrease to 5–15 mg/day in a single dose. The intellectual development and growth of children exposed to methimazole in utero appear to be similar to unexposed siblings.²⁰⁵

Dosage Forms. Tab 5, 10 mg.

Patient Instructions. Report sore throat, fever, or oral lesions immediately because they might be early signs of a rare, but severe, blood disorder. Also report any skin rashes, itching, or yellowing of eyes and skin. Be sure to take at prescribed dosage intervals.

Missed Doses. If you miss a dose, take it as soon as possible. If it is time for the next dose, take both doses.

Pharmacokinetics. Onset and Duration. PO onset about 2–3 weeks, which is consistent with the elimination of existing T₄ stores. Duration intrathyroidally 40 hr.¹⁶⁵,²⁰⁶

Serum Levels. <0.2 mg/L (1.8 μmol/L) inhibits iodide organification.²⁰⁶

Fate. Well absorbed orally. Considerable interindividual variations in pharmacokinetic parameters. Peak serum levels occur at 2.3 ± 0.8 hr; the peak after 30 mg orally is 0.8 ± 0.2 mg/L (6.8 ± 1.9 μmol/L); after 60 mg orally, 1.5 ± 0.5 mg/L (14 ± 4 μmol/L); after 60 mg rectally, 1.1 ± 0.5 mg/L (10 ± 5 μmol/L).²⁰³,²⁰⁶,²⁰⁷ The drug is actively concentrated in the thyroid gland, with peak intrathyroidal
levels of 0.11–1.1 mg/L (1–10 μmol/L) within 1 hr; there is minimal plasma protein binding; it is distributed into breast milk 10 times greater than PTU. Vd is 1.4 ± 0.6 L/kg; Cl is 0.072 ± 0.018 L/hr/kg. There are no active metabolites; 7–12% is excreted unchanged in urine, 6% excreted as inorganic sulfate, 1.5% as sulfur metabolites, and 50% as unknown metabolites.

$\frac{t_1}{2}$ phase 3 ± 1.4 hr; β phase 18.5 ± 13 hr in normal and hyperthyroid patients, increased to 21 hr in cirrhosis. Intrathyroidal half-life is 20 hr.

**Adverse Reactions.** Maculopapular skin rashes and itching occur frequently and can disappear spontaneously with continued treatment; urticaria requires drug discontinuation. Methimazole can be given to patients who develop only a nonurticarial maculopapular rash on PTU. Mild transient leukopenia occurs frequently in untreated Graves’ disease, does not predispose to agranulocytosis, and is not an indication to discontinue the drug. Agranulocytosis occurs occasionally, usually in the first 3 months of therapy. Risk increases with dosages >40 mg/day in patients >40 yr; granulocyte colony-stimulating factors (eg, filgrastim) can hasten recovery. Rarely, fever, arthralgias, cholestatic or hepatocellular toxicity, vasculitis, lupus-like syndrome, hypoprothrombinemia, aplastic anemia, thrombocytopenia, nephrotic syndrome, loss of taste, and spontaneous appearance of circulating antibodies to insulin or glucagon occur. Rare teratogenic risk of scalp defects.

**Contraindications.** Manufacturer states that breastfeeding is a contraindication, but most experts feel that breastfeeding can be performed with dosages of ≤10 mg/day and careful monitoring of infant thyroid function.

**Precautions.** Although methimazole crosses the placenta at rates 4 times greater than propylthiouracil and has been associated with scalp defects (aplasia cutis), recent reports indicate methimazole can be given in pregnant patients intolerant to PTU. Use with caution during lactation and in patients with severe allergic reactions to other thioamides. A low prevalence of cross-sensitivity occurs between thioamide compounds for nonurticarial skin rashes, so if these occur, another thioamide can be substituted. However, a 50% chance of cross-sensitivity exists for severe reactions (eg, agranulocytosis, hepatitis), so do not substitute another thioamide.

**Drug Interactions.** Iodide given before a thioamide delays the response to the thioamide, especially in thyroid storm. Changes in thyroid status can alter pharmacodynamics and pharmacokinetics of digoxin, warfarin, theophylline, β-blockers, and insulin.

**Parameters to Monitor.** Monitor clinical status; serum free T4 or T4 index, and TSH monthly initially until euthyroid, then q 3–6 months. Obtain occasional LFTs and CBC with differential (but these are not recommended routinely because they are not predictive of toxicity, and transient leukopenia and elevations in LFTs can occur). Obtain AST, ALT, total bilirubin, and alkaline phosphatase if patient reports signs of hepatitis; WBC and differential counts if patient reports signs of agranulocytosis such as fever, sore throat, or malaise.

**Notes.** Methimazole is the drug of choice for treatment of uncomplicated hyperthyroidism because it is better tolerated and fewer tablets can be given once daily.
improving patient compliance. Remission rates of 20–40% are common after cessation of therapy. Favorable remission rates correlate with longer duration of therapy, higher dosages, mild disease, shrinkage of goiter size with therapy, disappearance of thyroid receptor–stimulating antibodies, and initial presentation with $T_3$ toxicosis. Most patients eventually require surgery or radioiodine; however, a trial of a thioamide is worthwhile in patients with minimal thyroid enlargement or very mild hyperthyroidism. Adjunctive therapy with cholestyramine 4 g tid can lower thyroid hormone levels more rapidly. Methimazole rather than PTU may be preferred during radioactive iodine therapy because it does not interfere with the thyroid uptake of iodine like PTU. In thyroid storm, PTU is the drug of choice.

**Pharmacology.** Propylthiouracil (PTU) is a thioamide antithyroid drug that blocks the synthesis of thyroid hormones and, at dosages >450 mg/day, decreases the peripheral conversion of $T_4$ to $T_3$. Titers of thyroid receptor stimulating antibody decline during therapy, consistent with an immunosuppressive effect.

**Administration and Adult Dosage.** PO for hyperthyroidism 100–200 mg (depending on the severity of hyperthyroidism) q 6–8 hr initially until euthyroid (usually 6–8 weeks), then decrease by 33–50% over several weeks to a maintenance dosage of 50–150 mg/day in a single dose. Rarely, initial dosages of 1–1.2 g/day (maximum dosage) in 3–6 doses might be necessary. PO for thyroid storm 200–250 mg q 6 hr until euthyroid; maintenance dosage is determined by patient response. Traditional treatment duration for hyperthyroidism is 1–2 yr, although shorter courses of 8 months might be effective in mild disease. Treatment may be continued indefinitely, if necessary, to control the disease and if no toxicity occurs. The addition of levothyroxine is not recommended because remission rates have not shown improvement. PR PTU can be formulated for rectal administration.

**Special Populations.** Pediatric Dosage. Give orally in 3 divided doses. PO 150–300 mg/m²/day. Alternatively, (6–10 yr) 5–10 mg/kg/day or 50–150 mg/day initially; (≥10 yr) 150–300 mg/day initially. Maintenance dosage is determined by patient response.

**Geriatric Dosage.** Same as adult dosage.

**Other Conditions.** In pregnancy, the dosage should be as small as possible to maintain a mildly hyperthyroid maternal state; initially 300 mg/day orally in 3 divided doses for 4–6 weeks, then decrease to 50–150 mg/day in a single dose. The intellectual development and growth of children exposed to PTU in utero appear to be similar to unexposed siblings.

**Dosage Forms.** Tab 50 mg.

**Patient Instructions.** Report sore throat, fever, or oral lesions immediately because they may be an early sign of a severe, but rare, blood disorder. Also report any skin rashes, itching, or yellowing of eyes and skin. Be sure to take at prescribed dosage intervals.
Missed Doses. If you miss a dose, take it as soon as possible. If it is time for the next dose, take both doses.

Pharmacokinetics. Onset and Duration. PO onset of therapeutic effect 2–3 weeks, consistent with the elimination of existing thyroxine stores.

Serum Levels. Peak PTU levels >4 mg/L (24 μmol/L) produce antithyroid activity; 3 mg/L (18 μmol/L) reduces organification by 50%; 0.8 mg/L (5 μmol/L) reduces peripheral conversion activity by 50%.207,213

Fate. Oral bioavailability is 77 ± 13%. Peak levels occur 2 ± 0.3 hr after oral administration and 4.7 ± 1 hr after rectal administration. Peak serum level after an oral dose of 50 mg is 1 ± 0.2 mg/L (6 ± 1.2 μmol/L); after 200 mg, 4.5 ± 0.7 mg/L (26 ± 4 μmol/L); after 300 mg, 7 ± 0.8 mg/L (42 ± 5 μmol/L); after 400 mg rectally, 3 ± 0.8 mg/L (18 ± 5 μmol/L).211,212 PTU is actively concentrated in the thyroid gland, 40% as unknown metabolite, 32% as sulfate, and 20% as unchanged PTU; peak intrathyroidal levels of 0.17 ± 1.7 mg/L (1–10 μmol/L) occur within 1 hr.207 The drug is 80% plasma protein bound; it distributes poorly into breast milk.165 Vd is 0.29 ± 0.06 L/kg; Cl is 0.23 ± 0.04 L/hr/kg. About 85% is excreted in 24 hr, 61% as glucuronides, 8–9% as inorganic sulfates, 8–10% as unknown sulfur metabolites, and <10% excreted unchanged in urine.207,213

t½. 1.3 ± 0.6 hr.207,213

Adverse Reactions. (See Methimazole.) Agranulocytosis is not more prevalent at higher doses as it is with methimazole. Rarely, hepatitis occurs; hepatocellular toxicity is more frequent than cholestatic jaundice.165,175,214 Transient transaminase elevations can occur in asymptomatic individuals, which normalize within 3 months with continued drug administration.

Contraindications. Manufacturer states that breastfeeding is a contraindication, but it can be used with infant thyroid monitoring because of low milk levels and lack of effect on infants.165,205

Precautions. (See Methimazole.) Although it crosses the placenta poorly (25% that of methimazole), it can cause fetal hypothyroidism and goiter.205 Thyroid dysfunction can diminish as pregnancy progresses, allowing a reduction in dosage and, in some cases, a withdrawal of therapy 2–3 weeks before delivery. Adjunctive thyroid hormone therapy prevents maternal hypothyroidism but, because of minimal placental transfer, has little effect on the fetus.205 Use with caution before surgery or during treatment with anticoagulants because of hypoprothrombinemic effect.175

Drug Interactions. (See Methimazole.)

Parameters to Monitor. (See Methimazole.) INR monitoring is advisable, particularly before surgery.

Notes. Because propylthiouracil decreases peripheral conversion of T4 to T3, it is considered the thioamide of choice in treating thyroid storm. Some prefer PTU rather than methimazole in pregnancy and breastfeeding, although either can be used.165,205 Patients pretreated with PTU might require a 25% higher dosage of radioactive iodine for efficacy.210
REFERENCES


HORMONAL DRUGS


**Diuretics**

**Class Instructions.** Diuretics. If you are taking more than one dose a day, take the last dose in the afternoon or early evening to avoid having to void urine during the night. Avoid heavily salted foods, but rigid salt restriction is not necessary. Avoid excessive water intake. Report any dizziness or lightheadedness (especially when arising from sitting or lying), muscle cramps, weakness, lethargy, dry mouth, thirst, or low urine output.

**Missed Doses.** Take this drug at regular intervals. If you miss a dose, take it as soon as you remember. If it is about time for the next dose, take that dose only. Do not double the dose or take extra.

**AMILORIDE HYDROCHLORIDE**

**Pharmacology.** Amiloride is a potassium-sparing diuretic with a mechanism and site of action resembling triamterene. It has mild antihypertensive activity and a longer duration of action than triamterene.\(^1\)-\(^3\)

**Adult Dosage.** PO 10 mg/day in 1–2 doses, to a maximum of 20 mg/day, although a dosage >10 mg/day is seldom necessary.

**Dosage Forms.** Tab 5 mg; Tab 5 mg with hydrochlorothiazide 50 mg (Moduretic 5-50, various).

**Pharmacokinetics.** Onset within 2 hr; maximum effect 6–10 hr after an oral dose; duration about 24 hr. The drug is about 50% orally absorbed, decreasing to 30% when taken with food. Half the absorbed drug is excreted unchanged in urine. Half-life is 6–9 hr in normal renal function, increasing up to 144 hr in renal failure.

**Adverse Reactions.** Adverse reactions are generally similar to triamterene; however, in contrast to triamterene, renal stone formation has not been reported with amiloride.

**BUMETANIDE**

**Pharmacology.** Bumetanide is a loop diuretic with renal pharmacology similar to furosemide. Bumetanide is estimated to be approximately 40 times as potent as furosemide on a weight basis.\(^1\)-\(^6\)

**Administration and Adult Dosage.** PO for diuresis 0.5–2 mg as a single dose and repeat q 4–5 hr as needed, to a maximum of 10 mg/day. IV or IM dose is 0.5–1 mg, IV given over 1–2 min. Repeat doses may be administered as needed q 2–3 hr, to a maximum of 10 mg/day. For long-term control of edema, intermittent
regimens are recommended as alternate daily doses or daily doses for 3–4 days with 1–2 day drug holidays. **IV infusion** 1 mg IV bolus, followed by \((Cl_r > 75 \text{ mL/min})\) 0.5 mg/hr; \((Cl_r \ 25–75 \text{ mL/min})\) 0.5–1 mg/hr; or \((Cl_r < 25 \text{ mL/min})\) 1–2 mg/hr.\(^3\)

**Special Populations.** **Pediatric Dosage.** 0.015–0.1 mg/kg/dose q 6–24 hr, to a maximum of 10 mg/day.

**Geriatric Dosage.** Start with a low initial dose and titrate to response.

**Other Conditions.** No adjustment is necessary for renal impairment, hemodialysis, or chronic ambulatory peritoneal dialysis.

**Dosage Forms.** Tab 0.5, 1, 2 mg; **Inj** 0.25 mg/mL.

**Patient Instructions.** (See Class Instructions: Diuretics.)

**Pharmacokinetics.** **Onset and Duration.** Onset is within 30–60 min after oral administration and within minutes after IV administration. Durations of diuresis are 4–6 hr orally and 2–3 hr IV.\(^5,6\)

**Serum Levels.** Site of action is within the renal tubule and not the serum; therefore, serum concentrations do not reflect diuretic activity.

**Fate.** Bioavailability is 80–96%.\(^5,6\) \(V_{\text{dss}}\) is 0.16–0.24 L/kg in normal subjects. Protein binding to albumin is 94–97%. Renal excretion of unchanged drug accounts for 50% of administered drug, with hepatic metabolism and biliary excretion accounting for the remainder. The metabolites are inactive.

\(t_{1/2}\) 0.3–1.5 hr; 1.9 ± 0.1 hr in renal insufficiency; 2.3 ± 0.4 hr in cirrhosis.\(^5,6\)

**Adverse Reactions.** Hypokalemia, hyponatremia, and hyperuricemia occur frequently. Muscle cramps, dizziness, hypotension, headache, and nausea occur occasionally. The ototoxic potential of bumetanide is believed to be less than that of furosemide and most likely associated with rapid IV administration, high-dose therapy, or use in renal impairment.

**Contraindications.** Anuria; hepatic coma; coexisting severe electrolyte depletion.

**Precautions.** (See Furosemide.)

**Drug Interactions.** Aminoglycoside-related ototoxicity risk can be increased with concomitant bumetanide therapy. Cardiac glycoside toxicity is enhanced with diuretic-induced hypokalemia and hypomagnesemia. Concomitant use with other loop or thiazide diuretics enhances diuresis.

**Parameters to Monitor.** (See Furosemide.)

**FUROSEMIDE** Lasix, Various

**Pharmacology.** Furosemide is a loop diuretic that is actively secreted via the non-specific organic acid transport system into the lumen of the thick ascending limb of Henle’s loop, where it decreases sodium reabsorption by competing for the chloride site on the \(\text{Na}^+\text{-K}^+\text{-2Cl}^-\) cotransporter.\(^1,2\) Medullary hypertonicity is diminished, thereby decreasing the kidney’s ability to reabsorb water. Excretion of sodium, chloride, potassium, hydrogen ion, calcium, magnesium, ammonium, bicarbonate, and possibly phosphate is enhanced. IV furosemide increases venous
capacitance independent of diuretic effect, producing rapid improvement in pulmonary edema.\(^1\)

**Administration and Adult Dosage.** PO for edema 20–80 mg as a single dose initially; double successive doses q 6–8 hr until response is obtained. The maximum single oral dose depends on the disease state: 80 mg for hepatic cirrhosis with preserved renal function,\(^3\) 240 mg for nephrotic syndrome,\(^3\) 80–160 mg for CHF (with normal kidney function);\(^3\) however, dosages up to 2500 mg/day have been recommended in refractory CHF.\(^7\) After response, effective dosage is given in 1–3 doses daily; usual daily maintenance dosage depends on the single dose to which the patient responded. PO for chronic renal failure 80 mg initially, increasing in 80 mg/day increments until response is obtained, to a maximum of 160 mg and 400 mg for Cl\(_r\) <20 mL/min.\(^5\) (See Special Populations.) PO for hypertension 40 mg bid. IV should be used only when oral administration is not feasible. IV doses may be given over 1–2 min, except the rate should not exceed 4 mg/min when large doses are given. IM or IV for edema use one-half the dose of PO furosemide (as outlined above);\(^6\) may double the dose q 2 hr or more until desired response is obtained. This dosage is then given in 1–2 doses daily for maintenance. Dosages up to 4 g/day IV have been used in refractory CHF.\(^7\) IV for acute pulmonary edema 40 mg initially, may repeat in 30–60 min with 80 mg, if necessary (assuming relatively normal kidney function). For patients with renal impairment, the initial and subsequent doses must be adjusted based on renal function. For example, if Cl\(_r\) is 50 mL/min (about one-half normal), the dose must be doubled; if Cl\(_r\) is 25 mL/min, the dose must be quadrupled. Continuous IV infusion for edema 40 mg loading dose followed by (Cl\(_r\) 75 mL/min) 10 mg/hr; (Cl\(_r\) 25–75 mL/min) 10–20 mg/hr; (Cl\(_r\) <25 mL/min) 20–40 mg/hr.\(^3\)

**Special Populations.** Pediatric Dosage. PO for edema 2 mg/kg in 1 dose initially, increasing by 1–2 mg/kg in 6–8 hr, if necessary, to a maximum of 6 mg/kg/day. IM or IV 1 mg/kg in 1 dose initially, increasing by 1 mg/kg q 2 hr or more until desired response is obtained, or to a maximum of 6 mg/kg/day. Maximum single dose depends on renal function. (See Notes.) Geriatric Dosage. Start with a low initial dose and titrate to response.

**Other Conditions.** For Cl\(_r\) <20 mL/min, maximal response is attained with single IV doses of 200 mg (400 mg PO). Hence, there appears to be no need to administer larger single doses to such patients.\(^8\) A diminished response can occur in severe decompensated CHF, caused in part by alterations in oral absorption\(^9\) and decreased renal blood flow (despite relatively normal GFR), resulting in decreased delivery of furosemide to the renal tubule; IV administration circumvents absorption problems. However, decompensated CHF usually affects only the rate rather than the extent of oral furosemide absorption.\(^10\) In addition, high-dose therapy might be beneficial in severe, refractory CHF.\(^9\) (See Administration and Adult Dosage.) For patients with cirrhosis, dosage is based on renal function.

**Dosage Forms.** Soln 8, 10 mg/mL; Tab 20, 40, 80 mg; Inj 10 mg/mL.

**Patient Instructions.** (See Class Instructions: Diuretics.)

**Pharmacokinetics.** Onset and Duration. (Venous capacitance) IV onset 5 min, duration >1 hr. (Diuresis) PO onset 30–60 min, peak 1–2 hr, duration 6 hr; IV
onset 15 min, peak 30–60 min, duration 1–2 hr; duration might be prolonged in severe renal impairment. (Hypertension) maximum effect on BP might not occur for several days.

**Serum Levels.** Site of action is within the renal tubules and not the serum; therefore, serum concentrations do not reflect diuretic activity. High serum levels can be associated with ototoxicity.11

**Fate.** Pharmacokinetics are variable and absorption is erratic; 61 ± 17% (range 20–100) is bioavailable in normals, 30–100% in renal failure.11,12 The rate, but not the extent, of absorption might be decreased in patients with edematous bowel caused by decompensated CHF;10 96–99% is plasma protein bound, reduced in CHF, renal disease, or cirrhosis.12 \( V_d \) is 0.11 L/kg; Cl is 0.12 ± 0.24 L/hr/kg; \( V_d \) and Cl depend on protein binding.11,12 Two possible inactive metabolites exist: a glucuronide, which is excreted primarily in urine (and to a lesser extent in the feces by passive diffusion into the GI lumen), and saluamine, which could be a true metabolite or an analytical artifact. Renal clearance is primarily by active secretion; 50–80% (IV) and 20–55% (PO) are excreted unchanged in urine. Renal clearance is decreased in renal failure, consistent with decreased renal blood flow, a reduction of functioning nephrons, and the presence of competitive inhibitors for secretion.12

\( t_{1/2} \). 92 ± 7 (range 30–120) min in normals, can be extended in cirrhosis to 81 ± 8 min or CHF to 122 min, and markedly prolonged in end-stage renal disease to 9.7 hr, and in multiorgan failure to 20–24 hr. Mean residence time has been proposed as a more appropriate estimate of duration: 51.4 min (IV); 135–195 min (PO).11,12

**Adverse Reactions.** Dehydration, hypotension, hypochloremic alkalosis, and hypokalemia are frequent. Hyperglycemia and glucose intolerance occur as with thiazides. (See Hydrochlorothiazide.) With high-dose therapy (>250 mg/day), hyperuricemia occurs frequently. Tinnitus and hearing loss, occasionally permanent, occur frequently in association with rapid IV injection of large doses in patients with renal impairment.8,12,13 Rarely, thrombocytopenia, neutropenia, jaundice, pancreatitis, and a variety of skin reactions occur.

**Contraindications.** Anuria (except for single dose in acute anuria).

**Precautions.** Use with caution in patients with severe or progressive renal disease; discontinue if renal function worsens. Use with caution in liver disease (can precipitate hepatic encephalopathy), history of diabetes mellitus or gout, and in patients allergic to other sulfonamide derivatives. Use with caution in patients with hypokalemia, hypomagnesemia, or hypocalcemia.

**Drug Interactions.** Cholestyramine and colestipol decrease furosemide absorption, and NSAIDs can decrease the diuretic effect of furosemide. Aminoglycoside ototoxicity can be enhanced in renally impaired patients. IV furosemide can produce flushing, sweating, and BP variations in patients taking chloral hydrate.

**Parameters to Monitor.** Monitor serum potassium closely, other electrolytes periodically, and serum glucose, uric acid, BUN, and Cr, occasionally. Observe for clinical signs of fluid or electrolyte depletion such as dry mouth, thirst, weakness, lethargy, muscle pains or cramps, hypotension, oliguria, tachycardia, and GI upset.
Notes. Furosemide is light sensitive; oral solution should be stored at 15–30°C and protected from light. In severe proteinuria (>3.5 g/day), urinary albumin binds furosemide and reduces its effectiveness, explaining the higher dosage required to achieve adequate free drug concentrations. In general, clinical nonresponders tend to have a decreased fraction of loop diuretics excreted in the urine. For these patients and those with CHF and renal impairment, larger doses may force more drug into the tubule; however, the risk of ototoxicity must be considered. Alternatively, combined use with a thiazide or metolazone orally can be effective by blocking sodium reabsorption at multiple tubule sites; however, these agents (especially metolazone) have a slow onset of action. Alternative treatment regimens include IV acetazolamide and low-dose dopamine given with the loop diuretic. (See Loop Diuretics Comparison Chart.)
## Loop Diuretics Comparison Chart

<table>
<thead>
<tr>
<th>Drug</th>
<th>Dosage Forms</th>
<th>Adult Dosage&lt;sup&gt;a&lt;/sup&gt;</th>
<th>Pediatric Dosage&lt;sup&gt;b&lt;/sup&gt;</th>
<th>Dosage in Renal Impairment</th>
<th>Comments</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Bumetanide</strong>&lt;br&gt;Bumex**&lt;br&gt;Various**</td>
<td>Tab 0.5, 1, 2 mg&lt;br&gt;<strong>Inj 0.25 mg/mL</strong></td>
<td>PO for edema 0.5–2 mg/day, to a maximum of 10 mg/day, IM or IV over 1–2 min 0.5–1 mg, to a maximum of 10 mg/day IV continuous infusion 1 mg, then 1–2 mg/hr.</td>
<td>PO, IM, or IV 0.01–0.02 mg/kg, to a maximum of 0.4 mg/kg or 10 mg total daily dosage.</td>
<td>For Cl&lt;sub&gt;cr&lt;/sub&gt; &lt; 15 mL/min, maximal response is attained with single PO or IV doses of 8–10 mg; IV infusion of 12 mg over 12 hr may be more effective and less toxic.</td>
<td>1 mg PO or IV = 40 mg IV furosemide.</td>
</tr>
<tr>
<td><strong>Ethacrynic Acid</strong>&lt;br&gt;Edecrin</td>
<td>Tab 25, 50 mg&lt;br&gt;<strong>Inj 50 mg.</strong></td>
<td>PO minimal dose in the range of 50–200 mg/day initially, to a maximum of 200 mg bid IV 50 mg or 0.5–1 mg/kg, to a maximum of 100 mg.</td>
<td>PO (infants) not established; (children) 25 mg or 1 mg/kg initially, increased in 25 mg increments to desired effect; IV not established; 1 mg/kg has been used.</td>
<td>Not recommended with Cl&lt;sub&gt;cr&lt;/sub&gt; &lt; 10 mL/min; for Cl&lt;sub&gt;cr&lt;/sub&gt; of 10–50 mL/min, increase interval to q 8–12 hr.</td>
<td>Nonsulfonamide, Reliable potency data not available; however, 50 mg IV is about equal to furosemide 35 mg IV.</td>
</tr>
<tr>
<td><strong>Furosemide</strong>&lt;br&gt;Lasix**&lt;br&gt;Various</td>
<td>Tab 20, 40, 80 mg&lt;br&gt;<strong>Sol 8, 10 mg/mL</strong>&lt;br&gt;<strong>Inj 10 mg/mL.</strong></td>
<td>(See monograph.)</td>
<td>(See monograph.)</td>
<td>Maximum response occurs with 200 mg IV or an average of 400 mg PO, although quite variable.</td>
<td>IV dose averages 50% of PO dose, with great variability.</td>
</tr>
<tr>
<td><strong>Torsemide</strong>&lt;br&gt;Dernadex</td>
<td>Tab 5, 10, 20, 100 mg&lt;br&gt;<strong>Inj 10 mg/mL.</strong></td>
<td>(See monograph.)</td>
<td>(See monograph.)</td>
<td>Maximal response occurs with PO or IV dose of 50–100 mg.</td>
<td>5–10 mg PO or IV = 20 mg IV furosemide.</td>
</tr>
</tbody>
</table>

<sup>a</sup>Higher doses needed for patients with CHF, liver cirrhosis, and nephrotic syndrome. From references 5–7, 13, and 14, and product information.
**Pharmacology.** Thiazides increase sodium and chloride excretion by interfering with their reabsorption in the cortical diluting segment of the nephron; a mild diuresis of slightly concentrated urine results.\(^1,2\) Excretion of potassium, bicarbonate, magnesium, phosphate, and iodide excretion is increased; calcium excretion is decreased. Decreases in interstitial fluid volume, reductions in intracellular calcium secondary to a fall in smooth muscle sodium concentration, and a change in the affinity of cell surface receptors to vasoconstrictive hormones are thought to be among the mechanisms for the hypotensive effect of the thiazides. Urine output is paradoxically decreased in diabetes insipidus.\(^3\)

**Administration and Adult Dosage.** PO for edema 25–200 mg/day in 1–3 doses initially; 25–100 mg/day or intermittently for maintenance, to a maximum of 200 mg/day. PO for hypertension 12.5–50 mg/day. Maintenance dosages >50 mg/day provide little additional benefit in controlling essential hypertension and can increase the frequency of dose-related biochemical abnormalities.\(^15\)

**Special Populations.** Pediatric Dosage. PO (<6 months) up to 3.3 mg/kg/day in 2 divided doses; (>6 months) 2–2.2 mg/kg/day in 2 divided doses.

**Geriatric Dosage.** Start with a low initial dose and titrate to response.

**Other Conditions.** At a Cl\(_r\) <30 mL/min, usual dosages of thiazides and most related drugs are not very effective as diuretics but may be used in conjunction with loop diuretics.\(^16\)

**Dosage Forms.** Tab 25, 50, 100 mg; Cap 12.5 mg; Soln 10 mg/mL.

**Patient Instructions.** (See Class Instructions: Diuretics.) If stomach upset occurs, take drug with meals. Report persistent anorexia, nausea, or vomiting.

**Pharmacokinetics.** Onset and Duration. Onset of diuresis within 2 hr; peak in 4–6 hr; duration 6–12 hr. Onset of hypotensive effect in 3–4 days; duration \(\leq 1\) week after discontinuing therapy.

**Serum Levels.** The site of diuretic action is within the renal tubules and not the serum; therefore, serum concentrations do not reflect diuretic activity.

**Fate.** Oral bioavailability is 71 ± 15% in healthy individuals, increased when given with an anticholinergic, and decreased by one-half in CHF and after intestinal shunt surgery. There are no differences in absorption among single-entity formulations. The drug is 58 ± 17% plasma protein bound; \(V_d\) is 0.83 ± 0.31 L/kg; Cl is 0.29 ± 0.07 L/hr/kg. Over 95% is excreted unchanged in urine by filtration and secretion. In severe renal impairment, renal clearance is prolonged 5-fold, with nonrenal clearance (mechanism as yet unidentified) playing a larger role in elimination.\(^11,17\)

\[t_{1/2} = 2.5 ± 0.2\ hr;\] prolonged in uncompensated CHF or renal impairment.\(^11,17,18\)

**Adverse Reactions.** Hypokalemia is frequent; however, its treatment in otherwise healthy hypertensive patients is usually unnecessary. Potassium supplements or potassium-sparing diuretics (see Notes) may be indicated in patients with arrhythmias, MI, or severe ischemic heart disease; those with chronic liver disease; elderly eating poor diets; patients taking digoxin, a corticosteroid, or drugs that in-
interfere with ventricular repolarization such as phenothiazines and heterocyclic antidepressants; and those whose serum potassium level fall below 3 mEq/L. Hyperuricemia occurs frequently but is reversible, and treatment is unnecessary unless the patient has renal impairment or a history of gout. Hyperglycemia and alterations in glucose tolerance (usually reversible), loss of diabetic control, or precipitation of diabetes mellitus occur occasionally. Decreased glucose tolerance might increase in prevalence after several years of therapy. Thrombocytopenia and pancreatitis occur rarely. Elevation of serum cholesterol and triglycerides occurs; the clinical importance is unknown but can increase the risk of coronary heart disease.

**Contraindications.** Anuria; pregnancy, unless accompanied by severe edema; allergy to sulfonamide derivatives.

**Precautions.** Use with caution in patients with renal function impairment, liver disease (can precipitate hepatic encephalopathy), history of diabetes mellitus, or gout. Use with caution in patients with diabetes mellitus because thiazides might worsen glucose intolerance.

**Drug Interactions.** Cholestyramine and colestipol decrease oral absorption of thiazides, and NSAIDs can decrease the diuretic effect of thiazides. Anticholinergics can increase oral bioavailability. Dosage of potent hypotensive agents might have to be reduced if a thiazide is added to the regimen. Concurrent calcium-containing antacids can cause hypercalcemia. Long-term thiazides can reduce lithium excretion.

**Parameters to Monitor.** Monitor serum potassium weekly to monthly initially; q 3–6 months when stable. Monitor other serum electrolytes periodically. Monitor all electrolytes more closely when other losses occur (eg, vomiting, diarrhea). Observe for clinical signs of fluid or electrolyte depletion such as dry mouth, thirst, weakness, lethargy, muscle pains or cramps, hypotension, oliguria, tachycardia, and GI upset. Monitor BP periodically during antihypertensive therapy and serum glucose in patients with diabetes mellitus.

**Notes.** For the prevention of hypokalemia during thiazide therapy, a potassium-sparing diuretic may be preferred over potassium supplements in alkalotic patients because these agents decrease hydrogen ion loss, which can correct alkalosis and drive more potassium extracellularly. Potassium-sparing diuretics also may be preferred for patients predisposed to hypomagnesemia and for those with serum potassium <3 mEq/L because potassium supplements rarely correct hypokalemia of this severity. (See Thiazides and Related Diuretics Comparison Chart.) JNC-VI guidelines recommend diuretics and β-blockers as initial drugs of choice for patients with hypertension based on demonstrated reductions in morbidity and mortality.
<table>
<thead>
<tr>
<th>DRUG</th>
<th>DOSAGE FORMS</th>
<th>ORAL DIURETIC DOSAGE RANGE (MG/DAY)</th>
<th>EQUIVALENT DIURETIC DOSAGE (MG)</th>
<th>PEAK EFFECT (HR)</th>
<th>DURATION OF DIURESIS (HR)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Bendroflumethiazide</td>
<td>Tab 5, 10 mg.</td>
<td>2.5–15</td>
<td>5</td>
<td>4</td>
<td>6–12</td>
</tr>
<tr>
<td>Naturetin</td>
<td></td>
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</tr>
<tr>
<td>Benzthiazide</td>
<td>Tab 50 mg.</td>
<td>50–150</td>
<td>50</td>
<td>4–6</td>
<td>12–18</td>
</tr>
<tr>
<td>Exta</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Various</td>
<td></td>
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<td></td>
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</tr>
<tr>
<td>Chlorothiazide</td>
<td>Tab 250, 500 mg</td>
<td>500–2000</td>
<td>500</td>
<td>4 (PO)</td>
<td>6–12 (PO)</td>
</tr>
<tr>
<td>Diuril</td>
<td>Susp 50 mg/mL</td>
<td></td>
<td></td>
<td>0.5 (IV)</td>
<td>2 (IV)</td>
</tr>
<tr>
<td>Various</td>
<td>Inj 500 mg.</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Chlorthalidone^d</td>
<td>Tab (Thalitone)^e 15, 25 mg</td>
<td>100–200</td>
<td>50</td>
<td>2</td>
<td>24–72</td>
</tr>
<tr>
<td>Hygroton</td>
<td>Tab 25, 50, 100 mg.</td>
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<tr>
<td>Thalitone</td>
<td>Tab 25, 50, 100 mg.</td>
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<td></td>
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<tr>
<td>Various</td>
<td></td>
<td></td>
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<td></td>
<td></td>
</tr>
<tr>
<td>Hydrochlorothiazide</td>
<td>Cap 12.5 mg</td>
<td>25–100</td>
<td>50</td>
<td>4–6</td>
<td>6–12</td>
</tr>
<tr>
<td>Various</td>
<td>Tab 25, 50, 100 mg</td>
<td></td>
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<tr>
<td>Soln 10, mg/mL.</td>
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<tr>
<td>Hydroflumethiazide</td>
<td>Tab 50 mg.</td>
<td>25–200</td>
<td>50</td>
<td>3–4</td>
<td>12–24</td>
</tr>
<tr>
<td>Diucardin</td>
<td></td>
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<tr>
<td>Saluron</td>
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<tr>
<td>Various</td>
<td></td>
<td></td>
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</tr>
<tr>
<td>Indapamide^d</td>
<td>Tab 1.25, 2.5 mg.</td>
<td>2.5–5</td>
<td>2.5</td>
<td>2</td>
<td>up to 36</td>
</tr>
<tr>
<td>Lozol</td>
<td></td>
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<td></td>
<td></td>
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<tr>
<td>Various</td>
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</tbody>
</table>

^a Continued
<table>
<thead>
<tr>
<th>Drug</th>
<th>Dosage Forms</th>
<th>Oral Diuretic Dosage Range (mg/day)</th>
<th>Equivalent Diuretic Dosage (mg)</th>
<th>Peak Effect (hr)</th>
<th>Duration of Diuresis (hr)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Methyclothiazide</td>
<td>Tab 2.5, 5 mg.</td>
<td>2.5–10</td>
<td>5</td>
<td>6</td>
<td>24</td>
</tr>
<tr>
<td>Aquatensen</td>
<td></td>
<td></td>
<td></td>
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<td></td>
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<tr>
<td>Enduron</td>
<td></td>
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<td></td>
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<tr>
<td>Various</td>
<td></td>
<td></td>
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</tr>
<tr>
<td>Metolazone</td>
<td>Tab (Mykrox) 0.5 mg</td>
<td>5–20</td>
<td>5</td>
<td>2</td>
<td>12–24</td>
</tr>
<tr>
<td>Mykrox</td>
<td>Tab 2.5, 5, 10 mg.</td>
<td>0.5–1 (Mykrox)</td>
<td>(Zaroxolyn)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Zaroxolyn</td>
<td></td>
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</tr>
<tr>
<td>Polythiazide</td>
<td>Tab 1, 2, 4 mg.</td>
<td>1–4</td>
<td>2</td>
<td>6</td>
<td>24–48</td>
</tr>
<tr>
<td>Renese</td>
<td></td>
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</tr>
<tr>
<td>Quinethazone</td>
<td>Tab 50 mg.</td>
<td>50–200</td>
<td>50</td>
<td>6</td>
<td>18–24</td>
</tr>
<tr>
<td>Hydromox</td>
<td></td>
<td></td>
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<td></td>
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</tr>
<tr>
<td>Trichlormethiazide</td>
<td>Tab 2, 4 mg.</td>
<td>2–4</td>
<td>2</td>
<td>6</td>
<td>24</td>
</tr>
<tr>
<td>Metahydrin</td>
<td></td>
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<td></td>
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<tr>
<td>Naqua</td>
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<tr>
<td>Various</td>
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</tr>
</tbody>
</table>

*From USP-DI and product information; patients unresponsive to maximal dosage of one agent are unlikely to respond to another agent.

*Dosages are for edema.

*There is no therapeutic advantage in giving the drug parenterally.

*Not a thiazide, but similar in structure and mechanism of action.

*Thalitone and Mykrox are more bioavailable than other formulations of the respective drugs.
MANNITOL

Pharmacology. Mannitol and other osmotic diuretics do not act on specific receptors but rather on tubular fluid composition after filtration at the glomerulus. Mannitol inhibits sodium and chloride reabsorption in the proximal tubule and ascending loop of Henle predominantly. Excretion of sodium, potassium, calcium, and phosphate is increased. Renal blood flow is increased, the GFR of superficial nephrons is increased, and that of deep nephrons is decreased. Mannitol increases serum osmolality by expanding intravascular volume and decreasing intraocular and intracranial pressures.

Administration and Adult Dosage. Never administer IM or SC or add to whole blood for transfusion. IV as diagnostic evaluation of acute oliguria (if BP and CVP are normal and after cardiac output is maximized) give test dose of 12.5 g as a 15–20% solution over 3–5 min (often given with furosemide 80–120 mg IV), may repeat in 1 hr if urine output is <50 mL/hr. If no response after 2 doses, give no more mannitol and treat for acute tubular necrosis. If response occurs, look for underlying cause of oliguria (eg, hypovolemia). IV for prevention of acute renal failure give test dose as above to a total dose of ≥50 g in 1 hr as a loading dose, then maintain urine output at 50 mL/hr with continuous infusion of 5% solution, plus 20 mEq/L sodium chloride and 1 g/L calcium gluconate. IV for reduction of intracranial or intraocular pressure 1.5–2 g/kg over 30–60 min as a 15–20% solution. IV for reduction of nephrotoxicity of cisplatin 12.5 g IV push just before cisplatin, then 10 g/hr for 6 hr as a 20% solution. Replace fluids with 0.45% sodium chloride with 20–30 mEq/L potassium chloride at 250 mL/hr for 6 hr. IV for reduction of intracranial or intraocular pressure 1.5–2 g/kg over 30–60 min as a 15–25% solution.

Special Populations. Pediatric Dosage. IV for oliguria or anuria give test dose of 20 mg/kg as above; the therapeutic dose is 2 g/kg over 2–6 hr as a 15–20% solution. IV for reduction of intracranial or intraocular pressure 2 g/kg over 30–60 min as a 15–25% solution. IV for intoxications 2 g/kg as 5–10% solution as needed to maintain a high urinary output. (See Notes.)

Geriatric Dosage. Start with a low initial dose and titrate to response.

Dosage Forms. Inj 5, 10, 15, 20, 25%.

Pharmacokinetics. Onset and Duration. (Diuresis) onset 1–3 hr, duration depends on half-life. (Decrease in intraocular pressure) onset in 30–60 min, duration 4–6 hr. (Decrease in intracranial pressure) onset within 15 min, peak 60–90 min, duration 3–8 hr after stopping infusion.

Serum Levels. The site of diuretic action is within the renal tubules and not the serum; therefore, serum concentrations do not reflect diuretic activity.

Fate. About 17% is absorbed orally. IV doses of 1 and 2 g/kg increase serum osmolality by 11 and 32 mOsm/kg, decrease serum sodium by 8.7 and 20.7 mEq/L, and decrease hemoglobin by 2.2 and 2.5 g/dL, respectively. Vc is 0.074 L/kg; Vd is 0.23 L/kg; Cl is 0.086 L/hr/kg. Mannitol is eliminated almost completely unchanged in urine.

\[ t_{1/2} = \alpha \text{ phase } 0.11 \pm 0.12 \text{ hr; } \beta \text{ phase } 2.2 \pm 1.3 \text{ hr}. \]
**Adverse Reactions.** Most serious and frequent reactions are fluid and electrolyte imbalance, in particular symptoms of fluid overload (e.g., pulmonary edema, hypertension, water intoxication, and CHF). Acute renal failure has been reported occasionally with high doses, especially in patients with renal impairment.\(^{29,30}\) Dermal necrosis can occur if solution extravasates. Anaphylaxis has been reported rarely.

**Contraindications.** Patients with well-established anuria caused by severe renal disease or impaired renal function who do not respond to test dose; severe pulmonary congestion, frank pulmonary edema, or severe CHF; severe dehydration; edema not caused by renal, cardiac, or hepatic disease associated with abnormal capillary fragility or membrane permeability; active intracranial bleeding except during craniotomy.

**Precautions.** Pregnancy. Observe solution for crystals before administering. (*See Notes.*) Water intoxication can occur if fluid input exceeds urine output. Masking of inadequate hydration or hypovolemia can occur by drug-induced sustaining of diuresis. If extravasation occurs, aspirate any accessible extravasated solution, remove the IV catheter, and apply a cold compress to the area. Mannitol should not be added to whole blood for transfusion.

**Drug Interactions.** None known.

**Parameters to Monitor.** Monitor urine output closely and discontinue drug if output is low. Monitor serum electrolytes closely, taking care not to misinterpret low serum sodium as a sign of hypotonicity. (*See Fate.*) If serum sodium is low, measure serum osmolality. Observe for clinical signs of fluid or electrolyte depletion such as dry mouth, thirst, weakness, lethargy, muscle pains or cramps, hypotension, oliguria, tachycardia, and GI upset.

**Notes.** Mannitol can crystallize out of solution at concentrations >15%. The crystals can be redissolved by warming containers in hot water and shaking or by autoclaving; cool to body temperature before administration. Administer concentrated solutions through an inline filter. Addition of electrolytes to solutions of ≥20% concentration can cause precipitation.

**SPIRONOLACTONE**

**Pharmacology.** Spironolactone is a steroidal competitive aldosterone antagonist that acts from the interstitial side of the distal and collecting tubular epithelium to block sodium–potassium exchange, producing a delayed and mild diuresis. The diuretic effect is maximal in states of hyperaldosteronism. Excretion of sodium and chloride excretion is increased; excretion of potassium and magnesium is decreased.\(^{31-33}\) Spironolactone has mild antihypertensive activity and has demonstrated a beneficial effect in class III and IV CHF.\(^{34}\)

**Administration and Adult Dosage.** PO for edema 25–200 mg/day (usually 100 mg) in 2–4 divided doses initially, adjusting dosage after 5 days. If response is inadequate, add a thiazide or loop diuretic to the regimen. PO for essential hypertension 50–100 mg/day initially, adjusting dosage after 2 weeks. PO for ascites 100 mg/day initially, increasing to 200–400 mg/day in 2–4 divided doses. Restrict sodium to ≤2 g/day and, if necessary, fluid to 1 L/day. To eliminate delay
in onset, a loading dose of 2–3 times the daily dosage may be given on the first day of therapy. PO for Class III or IV CHF 12.5–25 mg/day. Special Populations. Pediatric Dosage. PO (neonates) 1–3 mg/kg/day q 12–24 hr; (older children) 1.5–3.3 mg/kg/day in divided doses q 6–24 hr. Geriatric Dosage. Start with a low initial dose and titrate to response. Dosage Forms. Tab 25, 50, 100 mg; Tab 25 mg with hydrochlorothiazide 25 mg (Aldactazide, various); Tab 50 mg with hydrochlorothiazide 50 mg (Aldactazide 50/50). Patient Instructions. (See Class Instructions: Diuretics.) Avoid excessive amounts of high-potassium foods or salt substitutes. Pharmacokinetics. Onset and Duration. Onset 1–2 days, peak 2–3 days with continued administration; onset can be hastened by giving loading dose; duration 2–3 days after cessation of therapy. Serum Levels. Not established and not used clinically. Fate. Bioavailability is about 90%; food promotes absorption and possibly decreases first-pass effect. Spironolactone undergoes rapid and extensive metabolism to canrenone (active metabolite), 7α-thiomethylspironolactone (major metabolite), and other sulfur-containing metabolites; together with the parent drug, these metabolites contribute to the overall antimineralocorticoid activity. Metabolites are eliminated primarily renally, with minimal biliary excretion. Little or no parent drug is excreted unchanged in urine. $t_\ell$ (Spironolactone) 1.4 ± 0.5 hr; (7α-thiomethylspironolactone) 13.8 ± 6.4 hr; (canrenone) 16.5 ± 6.3 hr. Adverse Reactions. Hyperkalemia can occur, most frequently in patients with renal function impairment (especially those with diabetes mellitus) and those receiving potassium supplements or concomitant ACE inhibitors. Dehydration and hyponatremia occur occasionally, especially when the drug is combined with other diuretics. In patients receiving high dosages, frequent estrogen-like side effects such as gynecomastia, decreased libido, and impotence in males occur; menstrual irregularities and breast tenderness occur in females. These effects are reversible after drug discontinuation. Contraindications. Anuria; acute renal insufficiency; rapidly deteriorating renal function; severe renal failure; serum potassium >5.5 mEq/L or development of hyperkalemia while taking the drug; hypermagnesemia. Precautions. Pregnancy. Patients with renal impairment, especially those with diabetes mellitus and/or receiving an ACE inhibitor, are at risk for developing hyperkalemia. Use with caution in patients with hepatic disease. Do not use with triamterene or amiloride. Give potassium supplements only to patients with demonstrated hypokalemia who are taking a proximally acting diuretic and a corticosteroid concurrently with spironolactone or only for very short periods in treating cirrhosis and ascites. Drug Interactions. Use with ACE inhibitors increases risk of hyperkalemia, especially in renal impairment. Spironolactone increases serum concentration of
digoxin by reducing renal clearance. In addition, spironolactone and its metabolites cross-react with digoxin-binding antibody in some digoxin immunoassays.

**Parameters to Monitor.** Monitor serum electrolytes, in particular potassium, periodically, especially early in the course of therapy. Monitor BUN and/or Cr, periodically. In ascites, also obtain daily weight and urinary electrolytes and maintain weight loss at no greater than 0.5–1 kg/day and urinary Na+/K+ ratio at >1. Observe for clinical signs of fluid or electrolyte depletion such as dry mouth, thirst, weakness, lethargy, muscle pains or cramps, hypotension, oliguria, tachycardia, and GI upset.

**Notes.** Spironolactone is used in the diagnosis of primary aldosteronism and may be useful in the management of the condition in patients unable to undergo surgery.

**TORSEMIDE**

**Pharmacology.** Torsemide is a loop diuretic similar to furosemide. Over the normal dosage range, its diuretic potency by weight is about 2–4 times that of furosemide. Onset of diuresis is similar but duration is longer (up to 8–12 hr orally).

**Adult Dosage.** PO for hypertension 5–10 mg/day orally. Initial PO or IV for edema or chronic renal failure 20 mg/day; dosage may be doubled until the desired response is obtained, to a usual maximum of 200 mg/day; or, IV by continuous infusion, give 20 mg loading dose, then 10–20 mg/hr. PO or IV for cirrhosis 5–10 mg/day initially with a potassium-sparing diuretic, to a usual maximum of 40 mg/day. (See Furosemide Notes and Loop Diuretics Comparison Chart.)

**Dosage Forms.** Tab 5, 10, 20, 100 mg; Inj 10 mg/mL.

**Pharmacokinetics.** Oral bioavailability is 79–91% (median 80); Vd is 0.14–0.19 L/kg. In healthy individuals, the elimination half-life of torsemide is dose dependent, ranging from 2.2 to 3.8 hr. Nonrenal Cl remains essentially constant over a dosage range of 5–20 mg, but renal Cl and fraction excreted decrease, suggesting saturable renal clearance. Further studies are needed to clarify whether torsemide undergoes dose-dependent renal elimination. Renal impairment (Clcr <60 mL/min) does not appreciably alter pharmacokinetic parameters; hemodialysis and hemofiltration do not markedly influence serum clearance.

**Adverse Reactions.** Although the potential for hypokalemia exists, torsemide’s kaliuretic potency is less than that of furosemide, suggesting that it is less potassium wasting during long-term therapy; the clinical relevance of this observation is unknown. Precautions and monitoring parameters are the same as those for furosemide.

**TRIAMTERENE**

**Pharmacology.** Triamterene acts directly from the distal tubular lumen on active sodium exchange for potassium and hydrogen, producing a mild diuresis that is independent of aldosterone concentration. Excretion of sodium, chloride, calcium, and possibly bicarbonate excretion is increased; excretion of potassium and possi-
bly magnesium excretion is decreased. Antihypertensive activity is inconsistent and less pronounced than with thiazides or spironolactone.32,33

Administration and Adult Dosage. PO initially 100 mg bid after meals if used alone, lower dosage if used with another diuretic. Adjust the maintenance dosage to the needs of the patient, which can range from 100 mg/day to 100 mg every other day, to a maximum of 300 mg/day.

Special Populations. Pediatric Dosage. PO 2–4 mg/kg/day initially, may increase to 6 mg/kg/day in 1–2 doses after meals, to a maximum of 300 mg/day. Decrease dosage if used with another diuretic.

Geriatric Dosage. Start with a low initial dose and titrate to response.

Dosage Forms. Cap 50, 100 mg; Cap 50 mg with hydrochlorothiazide 25 mg (Dyazide, various); Tab 75 mg with hydrochlorothiazide 50 mg (Maxzide, various); 37.5 mg with hydrochlorothiazide 25 mg (Maxzide-25, various).

Patient Instructions. (See Class Instructions: Diuretics.) This drug may be taken with food or milk to minimize stomach upset. Report persistent loss of appetite, nausea, or vomiting. Avoid eating excessive amounts of high-potassium foods or salt substitutes.

Pharmacokinetics. Onset and Duration. Onset 2–4 hr; full therapeutic effect might not occur for several days; duration 7–9 hr.

Serum Levels. The site of diuretic action is within the renal tubules and not the serum; therefore, serum concentrations do not reflect diuretic activity.

Fate. Variable absorption, depending on formulation,44,45 bioavailability is 52 ± 22%.55 When the total urinary excretion of triamterene and its pharmacologically active metabolite are considered, the bioavailability value of triamterene reaches 83.2 ± 25.9%.55 Triamterene undergoes marked first-pass metabolism with rapid hydroxylation followed by immediate conjugation to the sulfate ester, which is the predominant form in plasma and urine.45 The sulfate conjugate is nearly equipotent with the parent in causing sodium excretion and sparing of potassium.46,47 Triamterene is 50–55% plasma protein bound,34,35 and its sulfate conjugate is 91% protein bound.45 After oral administration, serum concentrations of triamterene and its sulfate conjugate undergo a rapid decline over the first 6–8 hr after administration, followed by a slower terminal phase.56 Both are eliminated renally by filtration and secretion. The fraction of a dose excreted as the parent is 3 ± 2%; that for the sulfate conjugate is 34 ± 8%.56 The sulfate conjugate can accumulate in renal impairment.47

\[ t_{1/2} \] phase (healthy adults) 4.3 ± 0.7 hr for triamterene and 3.1 ± 1.2 hr for sulfate;55 up to 12 hr in cirrhosis.48 Half-lives might be prolonged in the elderly.39

Adverse Reactions. Nausea, vomiting, diarrhea, and dizziness occur occasionally. Dehydration and hyponatremia with an increase in BUN occur occasionally, especially when the drug is combined with other diuretics. Triamterene renal stones occur occasionally. Hyperkalemia occurs occasionally, especially in diabetics and those with renal impairment; metabolic acidosis has been reported. Megaloblastic anemia can occur in alcoholic cirrhosis.
**Contraindications.** Severe or progressive renal disease or dysfunction (except possibly nephrosis); severe renal failure; severe hepatic disease; serum potassium $>5.5$ mEq/L or development of hyperkalemia while taking the drug; hypermagnesemia.

**Precautions.** Pregnancy. Patients with renal impairment, especially those with diabetes mellitus and/or receiving an ACE inhibitor, are at risk for developing hyperkalemia. Can elevate serum uric acid in patients predisposed to gout. Do not use with spironolactone or amiloride.

**Drug Interactions.** Use with ACE inhibitors increases risk of hyperkalemia, especially in renal impairment. Indomethacin (and probably other NSAIDs) can reduce renal function when combined with triamterene.

**Parameters to Monitor.** Monitor serum electrolytes, in particular potassium, periodically, especially early in the course of therapy. Monitor BUN and/or Cr, periodically. Observe for clinical signs of fluid or electrolyte depletion such as dry mouth, thirst, weakness, lethargy, muscle pains or cramps, hypotension, oliguria, tachycardia, and GI upset.
<table>
<thead>
<tr>
<th>CONDITION</th>
<th>LOOP DIURETICS</th>
<th>OSMOTIC DIURETICS</th>
<th>THIAZIDES</th>
<th>POTASSIUM-SPARING AGENTS</th>
<th>COMMENTS</th>
</tr>
</thead>
<tbody>
<tr>
<td>Relative potency</td>
<td>&gt;15%</td>
<td>10–15%</td>
<td>5–10%</td>
<td>&lt;5%</td>
<td>Values refer to maximum fraction of filtered sodium excreted after maximally effective dose of drug.</td>
</tr>
<tr>
<td>Hypertension</td>
<td>A</td>
<td>—</td>
<td>A</td>
<td>D</td>
<td>Sustained antihypertensive effect of thiazides exhibits a flat dose-response curve and occurs at doses below the threshold for diuresis. Loop diuretics are diuretics of choice with Cl&lt;sub&gt;cr&lt;/sub&gt; &lt; 30 mL/min.</td>
</tr>
<tr>
<td>Congestive heart failure</td>
<td>A (spironolactone)</td>
<td>—</td>
<td>A</td>
<td>A</td>
<td>Begin with thiazide with low dosage; if ineffective, substitute a loop diuretic. Loop diuretics are diuretics of choice with Cl&lt;sub&gt;cr&lt;/sub&gt; &lt; 30 mL/min. Spironolactone reduces morbidity and mortality in NYHA Class III and IV CHF.</td>
</tr>
<tr>
<td>Pulmonary edema</td>
<td>A (IV)</td>
<td>—</td>
<td>—</td>
<td>—</td>
<td>Prompt venodilation precedes diuretic effect.</td>
</tr>
<tr>
<td>Hepatic ascites</td>
<td>B</td>
<td>—</td>
<td>A</td>
<td></td>
<td>Spironolactone is the agent of choice; urine Na:K ratio &lt; 1 indicates need for higher dosage (200–1000 mg/day). Rate of diuresis should not exceed 750 mL/day (no peripheral edema), or up to 2 L/day (if edema is present).</td>
</tr>
<tr>
<td>Renal failure</td>
<td>A</td>
<td>C</td>
<td>—</td>
<td>—</td>
<td>A loop diuretic plus a thiazide (in a high dose) can evoke a clinically useful diuresis even when Cl&lt;sub&gt;cr&lt;/sub&gt; is &lt; 15 mL/min; however, provocative diuretic challenges in oliguric patients can be potentially hazardous, especially if the cause of renal failure is uncertain.</td>
</tr>
</tbody>
</table>

(continued)
## DIURETICS OF CHOICE COMPARISON CHART (continued)

<table>
<thead>
<tr>
<th>CONDITION</th>
<th>LOOP DIURETICS</th>
<th>OSMOTIC DIURETICS</th>
<th>THIAZIDES</th>
<th>POTASSIUM-SPARING AGENTS</th>
<th>COMMENTS</th>
</tr>
</thead>
<tbody>
<tr>
<td>Diabetes insipidus</td>
<td>—</td>
<td>—</td>
<td>A</td>
<td>—</td>
<td>Thiazides are most useful in the nephrogenic form; a long-acting agent is preferred. In pituitary form, oral diuretics may be a useful alternative for patients who prefer oral therapy to the use of intranasal or IV desmopressin.</td>
</tr>
<tr>
<td>Hypercalcemia</td>
<td>A</td>
<td>—</td>
<td>—</td>
<td>—</td>
<td>High-dose furosemide (IV 80–100 mg q 1–2 hr) with IV saline for forced diuresis to promote calcium excretion.</td>
</tr>
<tr>
<td>Hypercalciuria</td>
<td>—</td>
<td>—</td>
<td>A</td>
<td>—</td>
<td>Thiazides cause marked reduction in urinary calcium excretion; they also appear effective in preventing calcium stone formation irrespective of whether urinary calcium is abnormally elevated.</td>
</tr>
</tbody>
</table>

A = diuretic of choice; B = diuretic of second choice if patient is unresponsive to first choice; C = useful in some circumstances; D = useful as an adjunct to a more potent diuretic to reduce potassium loss and possibly enhance therapeutic effect.

*This table is a guide to the selection of the most appropriate diuretic for the condition listed but is not an all-inclusive guide to therapy. From references 32, 33, and 50–52.*
**Electrolytes**

**Class Instructions.** Oral Electrolytes. Take oral products with (tablets) or diluted in (liquids and powders) 6 to 8 fluid ounces of water or juice to avoid gastrointestinal injury or laxative effect. However, if you are undergoing hemodialysis, you may need to limit the volume of water you take. This medication may be taken with food or after meals if upset stomach occurs.

**Pharmacology.** Calcium plays an important role in neuromuscular activity, pancreatic insulin release, gastric hydrogen secretion, blood coagulation, and platelet aggregation; as a cofactor for some enzyme reactions; and in bone and tooth metabolism.\(^{53}\)

**Administration and Adult Dosage.** PO as dietary supplement (elemental calcium) recommended intake is (19–50 yr, including pregnant and lactating women) 1000 mg/day; (≥50 yr) 1200 mg/day.\(^{54,55}\) **PO to lower serum phosphate in end-stage renal disease (ESRD)** (calcium carbonate) 650 mg with each meal initially, adjust dosage to decrease serum phosphate to <6 mg/dL;\(^{56}\) (calcium acetate) 1334 mg with each meal initially, adjust dosage to decrease serum phosphate to <6 mg/dL. (See Notes.) **IV for emergency elevation of serum calcium** (calcium gluconate) 15 mg/kg in NS or D5W infused over 8–10 hr (typically raises serum calcium by 2–3 mg/dL),\(^{57}\) may repeat q 1–3 days depending on response; (calcium gluceptate) 1.1–1.4 g infused at a rate not to exceed 36 mg/min of elemental calcium. **IV for hypocalcemic tetany** 10–20 mL calcium gluconate infused over 10 min, may repeat until tetany is controlled. **Faster IV infusion rates can result in cardiac dysfunction.**\(^{58}\)

**Special Populations.** Pediatric Dosage. PO as dietary supplement (elemental calcium) adequate intake is (0–6 months) 210 mg/day; (7–12 months) 270 mg/day; (1–3 yr) 500 mg/day; (4–8 yr) 800 mg/day; (9–18 yr) 1300 mg/day.\(^{54}\) PO for hypocalcemia (elemental calcium) (neonates) 50–150 mg/kg/day in 4–6 divided doses, to a maximum of 1 g/day; (children) 20–65 mg/kg/day in 4 divided doses. **IV for emergency elevation of serum calcium** (infants) <1 mEq, may repeat q 1–3 days depending on response; (children) 1–7 mEq, may repeat q 1–3 days depending on response. **IV for hypocalcemic tetany** (infants) 2.4 mEq/day in divided doses; (children) 0.5–0.7 mEq/kg tid–qid, or more until tetany controlled.

**Geriatric Dosage.** Postmenopausal women have a requirement of 1200 mg/day, including those on estrogen replacement or a bisphosphonate.\(^{55}\) Lower dosage might be required in some patients because of the age-related decrease in renal function; conversely, requirements might increase with advanced renal insufficiency.

**Other Conditions.** Adolescence, renal impairment, and pregnancy might increase requirements; base maintenance dosage on serum calcium, serum phosphate, and diet.\(^{53,54}\)
**Dosage Forms.** (See Oral Calcium Products Comparison Chart.) **Inj** (chloride) 1 g/10 mL (contains 273 mg or 13.6 mEq Ca); (gluconate) 1 g/10 mL (contains 93 mg or 4.65 mEq Ca); (gluceptate) 1.1 g/5 mL (contains 90 mg or 4.5 mEq Ca).

**Patient Instructions.** (See Class Instructions: Oral Electrolytes.) Do not take within 2 hours of taking oral tetracycline or fluoroquinolone products. Take calcium tablets with food to maximize absorption. If used as a phosphate binder, calcium must be taken with food. Allow effervescent tablets to degas in a glass of water (about 4 minutes) before taking.

**Missed Doses.** Take this drug at regular intervals. If you miss a dose, take it as soon as you remember and then return to your normal dosage schedule.

**Pharmacokinetics. Serum Levels.** Normal serum total calcium is 8.4–10.2 mg/dL (2.1–2.6 mmol/L) for an adult with a serum albumin of 4 g/dL. Because a lesser fraction of calcium is protein bound in hypoalbuminemia, the patient’s value must be corrected based on serum albumin:

\[
\text{Corrected Serum Calcium} = \text{Serum Calcium in mg/dL} + (0.8 \times [4 – \text{Serum Albumin in g/dL}]).
\]

**Fate.** Oral calcium absorption is about 30% and depends on vitamin D and parathyroid hormone. Absorption decreases with age, high intake, achlorhydria, and estrogen loss at menopause; absorption increases when taken with food or in divided doses. Bioavailability from various salt forms does not appear to differ substantially in normals; however, differences in disintegration and dissolution among commercial formulations exist. About 99% of total body calcium is found in bone and teeth; of the 1% in extracellular fluid, 40–45% is plasma protein bound (mostly to albumin); 8–10% is complexed to citrate, phosphate, and other anions; and 45–50% is diffusible and physiologically active. About 135–155 mg/day are secreted into the GI tract, with 85% reabsorbed. Fecal loss of unabsorbed dietary calcium and endogenous excretion is 100–130 mg/day, urine loss is 150 mg/day, and sweat loss is 15 mg/day.

**Adverse Reactions.** IV calcium solutions, especially calcium chloride, are extremely irritating to the veins. Constipation or flatulence occurs frequently, especially with high dosages; the frequency probably does not differ markedly among salt forms. Calcium overload caused by oral calcium supplements is rare; immobilization, dosages in excess of 3–4 g/day, vitamin D therapy, and renal impairment can contribute to hypercalcemia, hypercalciuria, or nephrolithiasis during oral supplementation. Symptoms of calcium intolerance include nausea, intestinal bloating, excess gas, vomiting, constipation, abdominal pain, dry mouth, and polyuria.

**Contraindications.** Hypercalcemia; sarcoidosis; severe cardiac disease; digitalis glycoside therapy; calcium nephrolithiasis; calcium-phosphate product >60–70 in the setting of uremia is associated with calcification in extraosseous tissues and should be avoided. To determine calcium–phosphate product, multiply the serum phosphate value (in mg/dL) by the serum calcium value (in mg/dL).
Precautions. Avoid extravasation of parenteral calcium products. If extravasation occurs, aspirate any accessible extravasated solution, remove IV catheter, and apply a cold compress to the area.

Drug Interactions. Concomitant thiazide diuretic therapy and sodium depletion or metabolic acidosis can increase tubular reabsorption of calcium. Calcium reduces oral absorption of fluoroquinolones, tetracyclines, and iron salts. Concomitant use with sodium polystyrene sulfonate can lead to metabolic alkalosis and compromised activity of the binding resin.

Parameters to Monitor. Serum calcium regularly, with frequency determined by patient’s condition; BUN and/or Cr, serum phosphate, magnesium, and serum albumin (especially if low) periodically.

Notes. Calcium supplementation also can be achieved by dietary measures: skim milk provides 300 mg calcium/8 fluid ounces, 300 mg/8 fluid ounces of low-fat yogurt, 272 mg/ounce of Swiss cheese, and 200 mg/6 fluid ounces of calcium-fortified orange juice. Calcium carbonate is inexpensive and a good first-line agent. However, dissolution of calcium from phosphate and carbonate salts is pH dependent. These salts might not be optimal calcium sources for patients with elevated GI pH, such as the elderly or those with achlorhydria. Calcium carbonate as a chewable tablet or nougat or the use of an alternative calcium salt have been recommended. In ESRD, use calcium salts when serum phosphate is <8 mg/dL; when serum phosphate is >8 mg/dL, use aluminum hydroxide. Calcium acetate binds about twice the amount of phosphorus for the same quantity of calcium absorbed; however, the frequency of hypercalcemia does not seem to be diminished. (See Oral Calcium Products Comparison Chart.)
**ORAL CALCIUM PRODUCTS COMPARISON CHART**

<table>
<thead>
<tr>
<th>PRODUCT</th>
<th>PERCENTAGE ELEMENTAL CALCIUM</th>
<th>ELEMENTAL CALCIUM CONTENT</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Calcium Acetate</strong></td>
<td>25</td>
<td>667 mg Tab = 169 mg</td>
</tr>
<tr>
<td>Calphron</td>
<td></td>
<td>667 mg Tab = 169 mg</td>
</tr>
<tr>
<td>PhosLo</td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Calcium Carbonate</strong></td>
<td>40</td>
<td>6 mL Susp = 500 mg</td>
</tr>
<tr>
<td>Calciday-667</td>
<td></td>
<td>650 mg Tab = 260 mg</td>
</tr>
<tr>
<td>Cal-Sup</td>
<td></td>
<td>667 mg Tab = 267 mg</td>
</tr>
<tr>
<td>Caltrate 600</td>
<td></td>
<td>750 mg Tab = 300 mg</td>
</tr>
<tr>
<td>Os-Cal</td>
<td></td>
<td>1250 mg Tab = 500 mg</td>
</tr>
<tr>
<td>Tums</td>
<td></td>
<td>1500 mg Tab = 600 mg</td>
</tr>
<tr>
<td><strong>Calcium Citrate</strong></td>
<td>21.1</td>
<td>950 mg Tab = 200 mg</td>
</tr>
<tr>
<td>Citracal Tablets</td>
<td></td>
<td>2376 mg Tab = 500 mg</td>
</tr>
<tr>
<td>Citracal Liquitabs</td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Calcium Glubionate</strong></td>
<td>6.5</td>
<td>5 mL Syrup = 115 mg</td>
</tr>
<tr>
<td>Neo-Caliglucon</td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Calcium Gluconate</strong></td>
<td>9.3</td>
<td>500 mg Tab = 45 mg</td>
</tr>
<tr>
<td>Various</td>
<td></td>
<td>650 mg Tab = 58.5 mg</td>
</tr>
<tr>
<td></td>
<td></td>
<td>975 mg Tab = 87.8 mg</td>
</tr>
<tr>
<td></td>
<td></td>
<td>1000 mg Tab = 90 mg</td>
</tr>
<tr>
<td><strong>Calcium Lactate</strong></td>
<td>13</td>
<td>325 mg Tab = 42.3 mg</td>
</tr>
<tr>
<td>Various</td>
<td></td>
<td>650 mg Tab = 84.5 mg</td>
</tr>
<tr>
<td><strong>Calcium Phosphate, Tribasic</strong></td>
<td>39</td>
<td>1565 mg Tab = 600 mg</td>
</tr>
<tr>
<td>Posture</td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Dairy Products</strong></td>
<td></td>
<td>Cheese 28 g = 300–400 mg</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Skim milk 250 mL = 300 mg</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Yogurt 28 g = 43 mg</td>
</tr>
</tbody>
</table>

*From references 54–57 and product information.*

**MAGNESIUM SALTS**

**Pharmacology.** Magnesium is the second most abundant intracellular cation, with an essential role in neuromuscular function and protein and carbohydrate enzymatic systems; it functions as a cofactor for enzymes involved in transfer, storage, and utilization of intracellular energy. Magnesium also is an integral component of bone matrix.61

**Administration and Adult Dosage.** **PO as dietary supplement** (elemental magnesium) RDA is (≥11 yr) 410–420 mg/day for males and 320–360 mg for non-pregnant, nonlactating women.54 **PO for symptomatic chronic deficiency** (elemental magnesium) 12–24 mg/kg in divided doses.62 A renal threshold for magnesium excretion exists, so replacement is best accomplished slowly, usually over 5 days. **IV for prevention of negative balance** (elemental magnesium) 100–200 mg/day in parenteral nutrition solution.62 **IM for mild deficiency** 1 g MgSO₄ q 4–6 hr until serum magnesium is normalized or signs and symptoms
abate. IM for severe hypomagnesemia 2 g MgSO₄ as a 50% solution q 8 hr until serum magnesium is normalized or signs and symptoms abate; because IM injections are painful, continuous IV infusions might be preferred. IV infusion for severe hypomagnesemia 48 mEq/day (6 g MgSO₄) for 3–7 days by continuous infusion. IV for life-threatening hypomagnesemia (acute arrhythmias and seizures) 8–16 mEq (1–2 g MgSO₄) over 5–10 min, followed by continuous infusion of 48 mEq magnesium/day. IV for pre-eclampsia or eclampsia 4–6 g MgSO₄, then 1–2 g/hr by continuous infusion to maintain target serum level. (See Notes.)

Special Populations. Pediatric Dosage. IV for hypomagnesemia 25 mg/kg MgSO₄ as a 25% solution over 3–5 min q 6 hr for 3–4 doses. IM for seizures 20–40 mg/kg MgSO₄ as a 20% solution as needed. IV for severe seizures 100–200 mg/kg MgSO₄ as a 1–3% solution infused slowly with close monitoring of blood pressure. Administer one-half the dose during the initial 15–20 min and the total dose within 1 hr.

Geriatric Dosage. Lower dosage might be required in some patients because of the age-related decrease in renal function.

Other Conditions. Base maintenance dosage on serum magnesium and diet. Renal impairment decreases requirement. In severe renal failure, reduce dosage by at least 50% of the recommended amount and monitor serum magnesium after each dose. Concomitant administration of potassium and calcium may be necessary because many causes of hypomagnesemia also lead to hypocalcemia and hypokalemia.

Dosage Forms. (See Magnesium Products Comparison Chart.)

Patient Instructions. (See Oral Electrolytes Class Instructions.)

Missed Doses. Take this drug at regular intervals. If you miss a dose, take it as soon as you remember. Do not double the dose or take extra.

Pharmacokinetics. Onset and Duration. Peak levels are achieved immediately after IV, 1 hr after IM. Duration (anticonvulsant) is 30 min with IV, 3–4 hr post-onset with IM.

Serum Levels. (Normal) 1.3–2.1 mEq/L (0.65–1.1 mmol/L); (pre-eclampsia or eclampsia) 4–6 mEq/L (2–3 mmol/L). Intracellular and extracellular concentrations can vary independently; hence, serum magnesium levels might not be indicative of total body stores.

Fate. Oral absorption varies inversely with intake; in general, 24–76% is absorbed, principally in upper small intestine. Total body content is about 24 g, 60% of which is in bone, 39% in tissues, and 1% in extracellular fluid; 30% is plasma protein bound. Elimination is primarily by the kidneys, with only 1–2% in feces. Raising the serum concentration above normal exceeds the maximum tubular reabsorption capacity with subsequent excretion of excess.

Adverse Reactions. Serum concentration related: (3–5 mEq/L; 1.5–2.5 mmol/L) hypotension; (5–10 mEq/L; 2.5–5 mmol/L) PR interval changes, QRS prolongation, peaked T waves; (10 mEq/L; 5 mmol/L) areflexia; (15 mEq/L; 7.5 mmol/L)
respiratory paralysis; (25 mEq/L; 12.5 mmol/L) cardiac arrest. Pain on IM injection occurs very frequently. Pain on IM injection occurs very frequently.

**Contraindications.** Hypermagnesemia; heart block; myocardial damage; severe renal failure.

**Precautions.** Use with caution in patients with renal impairment (Clcr <30 mL/min) and those concurrently taking a digitalis glycoside. With bolus MgSO₄ administration, 1 g of 10% calcium gluconate IV should be available in case apnea or heart block occurs.

**Drug Interactions.** IV magnesium can potentiate neuromuscular blocking agents.

**Parameters to Monitor.** Serum magnesium regularly, frequency determined by condition of patient; BUN and/or Crs, serum potassium, and calcium periodically. Deep tendon reflexes, respiratory rate, BP, and ECG periodically.

**Notes.** For mild deficiencies, dietary supplementation may be sufficient to normalize magnesium stores; sources are cereals, nuts, green vegetables, meat, and fish. Magnesium gluconate is preferred for oral replacement and supplementation because it is possibly better absorbed and potentially causes less diarrhea. Patients on long-term diuretic therapy who are prone to hypomagnesemia may benefit from using the minimally effective dose of diuretic and 20–30 mEq/day of magnesium orally or changing to a magnesium-sparing agent (ie, amiloride, spironolactone, or triamterene). Drugs known to produce hypomagnesemia are aminoglycoside antibiotics, amphotericin B, diuretics, alcohol, and cisplatin. Coadministration of 3 g MgSO₄ IV with high-dose cisplatin chemotherapy has been recommended. The role of IV magnesium for acute MI remains unresolved. Correction of refractory hypocalcemia and hypokalemia with concurrent hypomagnesemia requires magnesium replacement to restore mineral balance. To avoid precipitation when MgSO₄ and calcium chloride are added to parenteral nutrition mixtures, use of calcium gluceptate has been recommended because it reacts more slowly than calcium chloride and a precipitate does not form. (See Magnesium Products Comparison Chart.)
<table>
<thead>
<tr>
<th>PRODUCT</th>
<th>MAGNESIUM CONTENT&lt;sup&gt;a&lt;/sup&gt; (MEQ/G)</th>
<th>DOSAGE FORMS&lt;sup&gt;b&lt;/sup&gt;</th>
<th>COMMENTS</th>
</tr>
</thead>
<tbody>
<tr>
<td>Magnesium, Chelated</td>
<td>8.3</td>
<td>Tab 500 mg = 100 mg Mg.</td>
<td>Amino acid chelate; sodium free; oral use only.</td>
</tr>
<tr>
<td>Chelated magnesium</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Magnesium Chloride</td>
<td>9.8</td>
<td>SR Tab 535 mg = 64 mg Mg</td>
<td>Alternative to parenteral MgSO&lt;sub&gt;4&lt;/sub&gt;.</td>
</tr>
<tr>
<td>Slo-Mag</td>
<td></td>
<td>Inj 200 mg/mL = 23.6 mg/mL Mg.</td>
<td></td>
</tr>
<tr>
<td>Various</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Magnesium Citrate</td>
<td>4.4</td>
<td>Soln 60 mg/mL = 3.2 mg/mL Mg.</td>
<td>Oral use only.</td>
</tr>
<tr>
<td>Various</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Magnesium Gluconate</td>
<td>4.5–4.8</td>
<td>Tab 500 mg = 27–29 mg Mg</td>
<td>Very soluble; well absorbed; produces no diarrhea.</td>
</tr>
<tr>
<td>Almora</td>
<td></td>
<td>Soln 11 mg/mL = 0.63 mg/mL Mg.</td>
<td></td>
</tr>
<tr>
<td>Magatrate</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Magonate</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Magnesium Hydroxide</td>
<td>34</td>
<td>Susp 40 mg/mL = 16.3 mg/mL Mg.</td>
<td>Readily available in combination antacid formulations.</td>
</tr>
<tr>
<td>Milk of Magnesia</td>
<td></td>
<td>Susp 80 mg/mL = 32.6 mg/mL Mg.</td>
<td>Start with 5 mL Susp or 1 Tab, increase as tolerated to qid. Requires gastric acid for absorption. Inexpensive.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Tab 300 mg = 122 mg Mg</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>Tab 600 mg = 244 mg Mg</td>
<td></td>
</tr>
</tbody>
</table>

(continued)
### MAGNESIUM PRODUCTS COMPARISON CHART (continued)

<table>
<thead>
<tr>
<th>PRODUCT</th>
<th>MAGNESIUM CONTENT(^a) (MEQ/G)</th>
<th>DOSAGE FORMS(^b)</th>
<th>COMMENTS</th>
</tr>
</thead>
<tbody>
<tr>
<td>Magnesium Oxide</td>
<td>49.6</td>
<td>Cap 140 mg = 84 mg Mg.</td>
<td>Poorly soluble; net absorption low, especially in malabsorptive states.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Tab 400 mg = 238 mg Mg.</td>
<td></td>
</tr>
<tr>
<td>Magnesium Sulfate</td>
<td>8.1</td>
<td>Inj 10% = 9.6 mg/mL Mg.</td>
<td></td>
</tr>
<tr>
<td>Epsom salt</td>
<td></td>
<td>Inj 12.5% = 12 mg/mL Mg.</td>
<td>Use IV, IM, or PO.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Inj 50% = 48 mg/mL Mg.</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>Pwdr 1 g = 97.2 mg Mg.</td>
<td></td>
</tr>
</tbody>
</table>

\(^a\) 1 mEq = 12 mg = 0.5 mmol Mg.

\(^b\) Magnesium products exhibit variable oral absorption; increase dosage incrementally until no further rise in serum magnesium occurs or until diarrhea occurs.
Pharmacology. Phosphate is a structural element of bone and is involved in carbohydrate metabolism, energy transfer, and muscle contraction, and as a buffer in the renal excretion of hydrogen ion. Many of the factors that influence serum calcium concentration also influence serum phosphate directly or indirectly.

Administration and Adult Dosage. The RDA is 700 mg/day. PO for phosphate replacement 250–500 mg (8–16 mmol) of phosphorus tid–qid; IV replacement (recent and uncomplicated hypophosphatemia) 0.08 mmol/kg, to a maximum of 0.2 mmol/kg; (prolonged and multiple causes) 0.16 mmol/kg, to a maximum of 0.24 mmol/kg. Infuse doses over 6 hr and additional dosage guided by serum concentrations. IV for symptomatic hypophosphatemia patients with phosphate levels of 1.6–1.9 mg/dL have received 15 mmol over 2 hr and those with phosphate <1.24 mg/dL have received 30 mmol over 3 hr with success (both without regard to weight). Reassess at completion of infusion to determine need for additional therapy. When serum concentration reaches 2 mg/dL and the patient can eat a normal diet, change to oral administration and a phosphate-rich diet. (See Phosphate Products Comparison Chart)

Special Populations. Pediatric Dosage. The RDAs are (0–6 months) 100 mg/day; (7–12 months) 275 mg/day; (1–8 yr) 460–500 mg/day; (9–18 yr) 1250 mg/day. PO for replacement (<4 yr) 250 mg (8 mmol) qid initially; (4 yr) same as adult dosage. IV replacement (serum phosphate 0.5–1 mg/dL) 0.05–0.08 mg/kg (0.15–0.25 mmol/kg) per dose over 4–6 hr; (serum phosphate <0.5 mg/dL) 0.08–0.12 mg/kg (0.25–0.35 mmol/kg) per dose over 6 hr. Repeat doses as needed to achieve desired serum concentration. Actual dosage depends on signs, symptoms, and serum phosphate concentration.

Geriatric Dosage. Lower dosage might be required in some patients because of the age-related decrease in renal function.

Other Conditions. Renal impairment decreases requirement. Choose the appropriate salt form based on the patient’s sodium and potassium requirements. Requirement is increased during alcohol withdrawal, diabetic ketoacidosis, respiratory alkalosis, aluminum antacid therapy, burns, postsurgical status, and nutritional repletion.

Dosage Forms. (See Phosphate Products Comparison Chart.)

Patient Instructions. (See Class Instructions: Oral Electrolytes.) Do not take capsules whole; instead, dissolve contents in 3/4 glass of water before taking. Powder in packets must be dissolved in 1 gallon of water before using. Chilling solution may improve palatability. Do not take with calcium-containing products.

Missed Doses. Take this drug at regular intervals. If you miss a dose, take it as soon as you remember. Do not double the dose or take extra.

Pharmacokinetics. Serum Levels. (As phosphorus) adults 2.7–4.5 mg/dL (0.9–1.5 mmol/L); children 4.5–5.5 mg/dL (1.5–1.8 mmol/L). Normal serum phosphorus concentrations can differ by as much as 0.6 mg/dL throughout the day because of changes in transcellular distribution. Concentrations <1.5 mg/dL indicate severe hypophosphatemia and require replacement therapy.
**Fate.** Normal adult dietary intake is 1–1.8 g/day, 60–70% of which is absorbed, primarily in the duodenum and jejunum. Most of the absorbed phosphorus is excreted in urine.

**Adverse Reactions.** Diarrhea and stomach upset occur frequently with oral administration. Dose-related hyperphosphatemia, metastatic calcium deposition, dehydration, hypotension, hypomagnesemia, and hyperkalemia or hypernatremia (depending on salt used) can occur.

**Contraindications.** Hyperphosphatemia; hypocalcemia; hyperkalemia (potassium salt); hypernatremia (sodium salt); severe renal failure.

**Precautions.** Use cautiously in patients with renal impairment and those with hypocalcemia. Dilute IV forms before use and administer slowly.

**Drug Interactions.** None known.

**Parameters to Monitor.** Serum phosphorus regularly, frequency determined by condition of patient; BUN and/or Cr, serum calcium, and magnesium periodically. Monitor serum sodium and/or potassium periodically, depending on salt form used.

**Notes.** Phosphate salts can precipitate in the presence of calcium salts in IV solutions; add no more than 40 mmol of phosphate and 5 mEq of calcium per liter. Calcium supplementation may be necessary to prevent hypocalcemic tetany during phosphate repletion. IV calcium gluconate or calcium chloride may be given until tetany subsides. Inorganic phosphorus exists in the body as mono- and dibasic forms, the relative proportions of which are pH dependent. It is therefore preferable to report concentrations as mg/dL or mmol/L rather than mEq/L. (See Phosphate Products Comparison Chart.)
# PHOSPHATE PRODUCTS COMPARISON CHART

<table>
<thead>
<tr>
<th>PRODUCT</th>
<th>DOSAGE FORMS</th>
<th>PHOSPHORUS CONTENT&lt;sup&gt;b&lt;/sup&gt;</th>
<th>CATION CONTENT</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>mg</td>
<td>mmol</td>
</tr>
<tr>
<td><strong>POTASSIUM SALTS</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>K-Phos Original</td>
<td>Tab.</td>
<td>114</td>
<td>3.6</td>
</tr>
<tr>
<td>Neutra-Phos K</td>
<td>Pwdr Packet.</td>
<td>250 (per packet)</td>
<td>8</td>
</tr>
<tr>
<td>Potassium Phosphate</td>
<td>Inj.</td>
<td>94  (per mL)</td>
<td>3</td>
</tr>
<tr>
<td><strong>SODIUM SALTS</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Fleet's Phospho-Soda</td>
<td>Soln.</td>
<td>128 (per mL)</td>
<td>4.1</td>
</tr>
<tr>
<td>Sodium Phosphate</td>
<td>Inj.</td>
<td>94  (per mL)</td>
<td>3</td>
</tr>
<tr>
<td><strong>SODIUM-POTASSIUM SALTS</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>K-Phos Neutral</td>
<td>Tab.</td>
<td>250 (per tablet)</td>
<td>8</td>
</tr>
<tr>
<td>Neutra-Phos Plain</td>
<td>Pwdr Packet.</td>
<td>250 (per packet)</td>
<td>8</td>
</tr>
<tr>
<td>Skim Milk</td>
<td>Liquid.</td>
<td>931  (per quart)</td>
<td>30</td>
</tr>
</tbody>
</table>

<sup>a</sup>Contents of capsules, tablets, and powders must be diluted in water before administration.

<sup>b</sup>31.25 mg = 1 mmol.

*From references 69, 72, and 73 and product information.*
Potassium is the major cation of the intracellular space, where its major role is regulating muscle and nerve excitability. Another role is controlling intracellular volume (similar to sodium's control of extracellular volume), protein synthesis, enzymatic reactions, and carbohydrate metabolism. The chloride salt is preferred for most uses because concomitant chloride loss and metabolic alkalosis frequently accompany hypokalemia. Nonchloride salts are preferred in acidosis (eg, secondary to amphotericin B or carbonic anhydrase inhibitor therapy and in chronic diarrhea with bicarbonate loss).

**Administration and Adult Dosage.** Variable, must be adjusted to needs of patient. PO for prophylaxis with diuretic therapy prevention of hypokalemia can generally be accomplished by giving 20 mmol/day of KCl, whereas treatment requires as much as 40–100 mmol/day. For nonedematous, ambulatory patients with uncomplicated hypertension, the goal should be to achieve a serum potassium of ≥4 mmol/L, and concentrations ≤3.4 mmol/L should be treated. For edematous patients (eg, with CHF), consider routine supplementation with KCl even if the potassium is normal (eg, 4 mmol/L). In those with mild potassium deficits, 40–80 mEq/day is recommended; with severe deficit, 100–120 mEq/day is indicated with careful monitoring of serum potassium. IV administration in peripheral vein (serum potassium >2.5 mEq/L) may be infused at 10–20 mEq/hr; reserve rates faster than 20 mEq/hr for emergency situations; may repeat q 2–3 hr as needed; do not exceed a maximum concentration of 40 mEq/L. IV administration in central vein (serum potassium <2.5 mEq/L) 30–60 mEq/hr may be administered; do not exceed a maximum concentration of 80 mEq/L. Infusion into a central vein requires use of a volume control device. Potassium concentration should not exceed 60 mEq/L unless the infusion site is through a large vein distal to the heart (eg, femoral vein) or more than one IV line is available, in which case the potassium dose may be delivered through two different ports; however, more concentrated solutions (200 mEq/L) infused at slow rates (20 mEq/hr) have been used with relative safety. (See Special Populations, Other Conditions.)

**Special Populations. Pediatric Dosage.** PO 1–2 mEq/kg/day during diuretic therapy.

**Geriatric Dosage.** Lower dosage might be required in some patients because of the age-related decrease in renal function.

**Other Conditions.** Base maintenance dosage on serum potassium; renal impairment decreases requirement. For patients with renal impairment or any form of heart block, decrease infusion rate by one-half and do not exceed 5–10 mEq/hr.

**Dosage Forms.** PO. (See Potassium Products Comparison Chart.) Inj (potassium chloride) 2 mEq/mL; (potassium acetate) 2, 4 mEq/mL; (potassium phosphate) 4.4 mEq/mL of potassium and 3 mmol/mL of phosphate. (See Potassium Products Comparison Chart.)

**Patient Instructions.** (See Class Instructions: Oral Electrolytes.) Do not chew or crush tablets. The expanded wax matrix of sustained-release forms may be found in the stool, but this does not imply a lack of absorption.
Missed Doses. Take this drug at regular intervals. If you miss a dose, take it as soon as you remember and then return to your normal dosage schedule. Do not double the dose or take extra.

Pharmacokinetics. Onset and Duration. Peak elevation of serum potassium concentrations after SR preparations is slightly delayed (median of 2 hr) compared with the liquid form (median of 1 hr). Effect on serum potassium is most pronounced in the first 3 hr after administration.83

Serum Levels. Differs depending on laboratory. Normal serum levels are (newborn) 5–7.5 mEq/L, (child) 3.4–4.7 mEq/L, and (adult) 3.5–5.1 mEq/L. Total body stores are about 50 mEq/kg or 3500 mEq. As a general rule, a decrease of 1 mEq/L in serum potassium reflects a 10–20% total body deficit; however, there is considerable variation;81 signs of hypokalemia appear <2.5 mEq/L; concentrations >7 mEq/L or <2.5 mEq/mL are dangerous. Clinical signs of hypokalemia or hyperkalemia are not reliable indicators of serum concentrations. Alkalosis decreases concentrations, and acidosis increases concentrations. Any hypokalemia-induced change in ECG must be treated as a medical emergency with IV potassium. Likewise, hyperkalemia-induced changes in ECG must be treated as a medical emergency.

Fate. When initially administered, the rates of absorption and excretion are more rapid with the liquid than with the SR forms; however, bioavailability is the same (78–90%) during long-term administration.83,84 About 10 mEq/day is eliminated in feces, 60–90 mEq/day in urine, and 7.5 mEq/L in sweat.

Adverse Reactions. Bad taste, nausea, vomiting, diarrhea, and abdominal discomfort occur frequently with oral liquids. Do not use enteric-coated tablets because they can cause small-bowel and occasionally gastric ulceration.81 Local tissue necrosis can occur if IV solution extravasates. Hyperkalemia can occur occasionally. Patients with diabetic nephropathy are at increased risk for hyperkalemia.85

Contraindications. Severe renal impairment; untreated Addison’s disease; adynamia episodica hereditaria; acute dehydration; heat cramps; hyperkalemia; concurrent ACE inhibitor or potassium-sparing diuretic in patients with renal impairment.81 In addition, all solid dosage forms (including SR products) are contraindicated in patients in whom delay or arrest of the tablet through the GI tract can occur.

Precautions. Use with caution (if at all) in patients receiving potassium-sparing diuretics or ACE inhibitors and those with digitalis-induced atrioventricular conduction disturbances or renal failure. Avoid extravasation of parenteral potassium products. If extravasation occurs, aspirate any accessible extravasated solution, remove IV catheter, and apply a cold compress to the area.

Drug Interactions. Use with an ACE inhibitor or potassium-sparing diuretic can result in hyperkalemia.

Parameters to Monitor. Serum potassium weekly to monthly initially, q 3–6 months when stable, BUN and/or Cr, periodically. For supplementation in patients on long-term diuretic therapy, obtain pretreatment serum levels of potassium and magnesium and reassess after 2–3 weeks and then monthly to determine pattern of
potassium loss. Once steady state or normokalemia is achieved, assess quarterly or as condition requires.81

Notes. Place the patient on a cardiac monitor before starting IV potassium.79 A potassium-sparing diuretic may be preferable to potassium supplementation when large supplements are needed, aldosterone concentrations are elevated, enhanced diuretic response is desired, or magnesium loss is of concern. If large doses of potassium fail to correct hypokalemia, suspect hypomagnesemia because potassium balance is strongly dependent on magnesium homeostasis.78,86 If a hypokalemic patient is also hypomagnesemic, as occurs with amphotericin B therapy, the patient might not respond to potassium replacement therapy unless magnesium balance is restored. (See Potassium Products Comparison Chart.)

<table>
<thead>
<tr>
<th>POTASSIUM PRODUCTS COMPARISON CHART</th>
</tr>
</thead>
<tbody>
<tr>
<td>PRODUCT</td>
</tr>
<tr>
<td>Potassium Acetate</td>
</tr>
<tr>
<td>Various</td>
</tr>
<tr>
<td>Inj 2, 4 mEq/mL</td>
</tr>
<tr>
<td>Useful in metabolic acidosis; avoid in metabolic alkalosis.</td>
</tr>
<tr>
<td>Potassium Acetate/Soln</td>
</tr>
<tr>
<td>3 mEq/mL.a</td>
</tr>
<tr>
<td>Preferred form in patients with delayed GI transit time or metabolic acidosis; avoid nonchloride salts in metabolic alkalosis.</td>
</tr>
<tr>
<td>Potassium Chloride</td>
</tr>
<tr>
<td>K-Lyte/Cl</td>
</tr>
<tr>
<td>Soln 20, 30, 40, 45</td>
</tr>
<tr>
<td>Ideal for hypochloremic metabolic alkalosis.</td>
</tr>
<tr>
<td>K-Lyte</td>
</tr>
<tr>
<td>Tab, Effervescent 25, 50 mEq.</td>
</tr>
<tr>
<td>Preferred form in patients with delayed GI transit time or metabolic acidosis; avoid nonchloride salts in metabolic alkalosis.</td>
</tr>
<tr>
<td>Potassium Bicarbonate/Citrate</td>
</tr>
<tr>
<td>TriKates</td>
</tr>
<tr>
<td>Tri-K</td>
</tr>
<tr>
<td>Soln 3 mEq/mL.a</td>
</tr>
<tr>
<td>Preferred form in patients with delayed GI transit time or metabolic acidosis; avoid nonchloride salts in metabolic alkalosis.</td>
</tr>
<tr>
<td>Potassium Gluconate</td>
</tr>
<tr>
<td>Kao</td>
</tr>
<tr>
<td>Soln 1.33 mEq/mL.a</td>
</tr>
<tr>
<td>Preferred form in patients with metabolic acidosis; avoid nonchloride salts in metabolic alkalosis.</td>
</tr>
</tbody>
</table>

aLiquids have rapid absorption, low frequency of GI ulceration, and unpleasant taste.
bBioequivalent to liquid forms; avoid in patients with delayed GI transit time.

From references 78, 79, 81 and product information.
ORAL REHYDRATION SOLUTIONS

Pharmacology. Oral rehydration solutions supply sodium, chloride, potassium, and water to prevent or replace mild to moderate fluid loss (5–10% dehydration) in diarrhea or postoperative states or when food and liquid intakes are temporarily discontinued. A carbohydrate (usually 2–2.5% glucose) is present to aid in sodium transport and subsequent water absorption.87,89

Administration and Adult Dosage. PO 1900–2850 mL (2–3 quarts)/day. Give only enough solution to supply the calculated water loss plus daily requirement.

Special Populations. Pediatric Dosage. Depends primarily on estimated fluid and electrolyte losses.90 PO for mild dehydration (3–5%) 50 mL/kg of oral rehydration plus replacement of ongoing losses (10 mL/kg for each diarrheal stool and replace estimated emesis) over 4 hr.90 Reassess patient q 2 hr. PO for moderate dehydration (6–9%) 100 mL/kg or oral rehydration fluid plus replacement of ongoing losses over 4 hr.90 Reassess hydration status of patient hourly. Severe dehydration (10% or greater) IV replacement fluids are indicated.

Geriatric Dosage. Lower dosage might be required because of the age-related decrease in renal function.

Other Conditions. Adjust intake based on fluid status and serum electrolytes. When electrolyte-containing foods are restarted, adjust solution intake accordingly.

Dosage Forms. (See Oral Rehydration Solutions Comparison Chart.)

Patient Instructions. These products are not for fluid replacement in prolonged or severe diarrhea. Reconstitute powdered products in tap water; do not mix with milk or fruit juices. If additional fluids are desired, drink water or other nonelectrolyte-containing fluids to quench thirst.

Adverse Reactions. Hypernatremia, hyperkalemia, and acid–base disturbances can occur occasionally, especially in renal insufficiency or if errors occur in reconstituting bulk powders.

Contraindications. Intractable vomiting; adynamic ileus; intestinal obstruction; perforated bowel; shock; renal dysfunction (anuria, oliguria); monosaccharide malabsorption.88

Precautions. Use parenteral replacement to correct electrolyte imbalances caused by severe fluid loss (10–15% of body weight), inability to take oral fluids, severe gastric distention, or severe vomiting. Errors in reconstituting or diluting commercial powders can have severe consequences.

Drug Interactions. None known.

Parameters to Monitor. Serum sodium, potassium, chloride, and bicarbonate regularly, with frequency determined by condition of patient; BUN and/or Cr, and urine-specific gravity periodically; input and output, weight, and signs and symptoms of dehydration daily.

Notes. To prevent dehydration early in the course of diarrhea or maintain hydration after parenteral replacement in adults and children, 90 mEq/L of sodium is acceptable. For infants who have higher insensible water losses, diluted solutions containing 50–60 mEq/L of sodium are suggested.92 Alternatively, solutions of
higher sodium concentration may be used in a ratio of 2:1 with additional free water. Vomiting does not preclude use of oral replacement solutions; spooning small quantities into the mouth of the child who is experiencing some vomiting usually results in the administration of sufficient fluid to correct dehydration. (See Oral Rehydration Solutions Comparison Chart.)
## Oral Rehydration Solutions Comparison Chart

<table>
<thead>
<tr>
<th>Solution</th>
<th>Dosage Forms</th>
<th>Electrolytes (mEq/L)&lt;sup&gt;a&lt;/sup&gt;</th>
<th>Carbohydrate</th>
</tr>
</thead>
<tbody>
<tr>
<td>Infalyte</td>
<td>Soln 1000 mL</td>
<td>Na&lt;sup&gt;+&lt;/sup&gt; 50 K&lt;sup&gt;+&lt;/sup&gt; 25 Cl&lt;sup&gt;−&lt;/sup&gt; 45 Base 34 Citrate Other —</td>
<td>Rice syrup solids 3%</td>
</tr>
<tr>
<td>Pedalyte</td>
<td>Soln 237, 946 mL</td>
<td>Na&lt;sup&gt;+&lt;/sup&gt; 45 K&lt;sup&gt;+&lt;/sup&gt; 20 Cl&lt;sup&gt;−&lt;/sup&gt; 35 Base 30 Citrate Other —</td>
<td>Dextrose 2.5%</td>
</tr>
<tr>
<td>Rehydralyte</td>
<td>Soln 237 mL</td>
<td>Na&lt;sup&gt;+&lt;/sup&gt; 75 K&lt;sup&gt;+&lt;/sup&gt; 20 Cl&lt;sup&gt;−&lt;/sup&gt; 65 Base 30 Citrate Other —</td>
<td>Dextrose 2.5%</td>
</tr>
<tr>
<td>Resol</td>
<td>Soln 960 mL</td>
<td>Na&lt;sup&gt;+&lt;/sup&gt; 50 K&lt;sup&gt;+&lt;/sup&gt; 20 Cl&lt;sup&gt;−&lt;/sup&gt; 50 Base 34 Citrate Other 4 Ca&lt;sup&gt;++&lt;/sup&gt; 4 Mg&lt;sup&gt;++&lt;/sup&gt; 5 HPO&lt;sub&gt;4&lt;/sub&gt;&lt;sup&gt;−&lt;/sup&gt;</td>
<td>Glucose 2%</td>
</tr>
<tr>
<td>WHO Oral Rehydration Salts&lt;sup&gt;b&lt;/sup&gt;</td>
<td>Powder&lt;sup&gt;c&lt;/sup&gt;</td>
<td>Na&lt;sup&gt;+&lt;/sup&gt; 90 K&lt;sup&gt;+&lt;/sup&gt; 20 Cl&lt;sup&gt;−&lt;/sup&gt; 80 Base 30 Bicarbonate Other —</td>
<td>Glucose 2%</td>
</tr>
</tbody>
</table>

<sup>a</sup>Optimal solution (mEq/L): Na<sup>+</sup> 75–100, K<sup>+</sup> 20–30, Cl<sup>−</sup> 65–100, base 20–30, carbohydrate 1.5–2%.

<sup>b</sup>Reconstitute powder in tap water; do not mix with milk or fruit juices.

<sup>c</sup>WHO = World Health Organization; available from Jianas Brothers Packaging, 2533 SW Boulevard, Kansas City, MO; tel (816) 421–2880.

From references 89–92 and product information.
SEVELAMER HYDROCHLORIDE

Pharmacology. Sevelamer is a polymer that binds phosphate in the GI tract. It is used in patients with end-stage renal disease to lower serum phosphate.

Adult Dosage. PO for hyperphosphatemia dosage is based on serum phosphate:
- For serum phosphate of (≥6 and ≤7.5 mg/dL) 800–806 mg tid;
- (≥7.5 and <9 mg/dL) 1200–1209 mg tid;
- (≥9 mg/dL) 1600–1612 mg tid.

Pediatric Dosage. Safety and efficacy not established.

Dosage Forms. Cap 403 mg; Tab 400, 800 mg.

Patient Instructions. Take the dosage form whole with meals. Do not chew tablet or capsule or take capsule apart. Take any other medications one hour or more before or 3 hours after taking this medication.

Pharmacokinetics. Sevelamer is not absorbed from the GI tract. It is eliminated in the feces.

Adverse Reactions. Well tolerated. Occasional nausea, dyspepsia, diarrhea, flatulence and constipation reported.

Contraindications. Hypophosphatemia; bowel obstruction.

Precautions. Use with caution in patients with dysphagia, swallowing disorders, GI motility disorders or major GI tract surgery.

Drug Interactions. Sevelamer might bind with concomitantly administered drugs and decrease their absorption.

SODIUM POLYSTYRENE SULFONATE

Pharmacology. A cation exchange resin that exchanges potassium for sodium. Each gram of resin binds up to 1 mEq of potassium and liberates 1–2 mEq of sodium.93 Sorbitol is present in some products to induce diarrhea and reduce the potential for fecal impaction. (See Notes.)

Administration and Adult Dosage. PO 15–20 g, may repeat as often as q 2 hr,79 although doses up to 40 g have been recommended;94 total dosage and duration of therapy depend on patient response. If suspension does not contain sorbitol, give powder with, or suspended in, a sorbitol solution (eg, 15 mL of 70% sorbitol). PR as enema 50 g retained for 30 min, if possible, may repeat as often as q 45 min.79 Follow enema by an irrigation of up to 2 L of nonsodium-containing fluid to remove resin from bowel.

Special Populations. Pediatric Dosage. For small children and infants, calculate dosage on the basis of 1 g of resin binding 1 mEq of potassium.

Geriatric Dosage. Lower dosage might be required in some patients because of the age-related decrease in renal function.

Other Conditions. In severe situations, such as ongoing tissue damage or rapidly rising serum potassium in renal failure, a dosage of 80–100 g for every mEq/L of potassium above 5 mEq/L has been recommended.94 However, under such circumstance, other forms of therapy may be considered.

Dosage Forms. Pwdr 454 g; Susp (containing sorbitol) 15 g/60 mL.
Pharmacokinetics. Onset and Duration. PO onset 1–2 hr; PR retention enema lowers potassium within 0.5–1 hr.79

Fate. Not absorbed from GI tract; binds potassium and liberates sodium as it passes through the intestine.

Adverse Reactions. Anorexia, nausea, and vomiting occur frequently with large doses; gastric irritation, constipation, and fecal impaction (especially in the elderly) occur occasionally. These effects can be avoided with enema. However, intestinal necrosis caused by enema has been reported. Use of sorbitol in the enema and failure to follow it with a cleansing enema can predispose uremic patients to potentially fatal intestinal necrosis.93

Precautions. Use with caution in patients who cannot tolerate any additional sodium load (eg, severe CHF, severe hypertension, marked edema). In addition to potassium, other cations (eg, magnesium, calcium) can bind to the resin, causing electrolyte imbalances. If rapid potassium lowering is required, give insulin with or without glucose.

Drug Interactions. None known.

Parameters to Monitor. Serum potassium at least daily and more frequently if indicated; serum magnesium and calcium periodically; ECG and patient signs and symptoms are useful in evaluating status.

Notes. On average, 50 g of resin will lower serum potassium by 0.5–1 mEq/L.95,96 Although sodium polystyrene sulfonate is used because of its ability to bind potassium, it also exchanges sodium for other di- and trivalent ions (eg, calcium, magnesium, iron). Rectal administration is less effective than oral use. Heating can alter the exchange properties of the resin. Sodium polystyrene sulfonate–induced constipation may be treated with 70% sorbitol in oral doses (ie, 10–20 mL/2 hr) sufficient to produce 1 or 2 watery stools/day.

Bisphosphonates

Aldronate Sodium

Pharmacology. Alendronate is a nitrogen-containing bisphosphonate that is 100–1000 times as potent as etidronate in inhibiting bone resorption in the rat.97 Bisphosphonates are cleared rapidly from the circulation and localized to hydroxyapatite bone mineral surfaces where they influence osteoclast function. Postulated cellular mechanisms of action include inhibition of osteoclast formation/recruitment, inhibition of osteoclast activation, inhibition of mature osteoclast activity, and induction of osteoclast apoptosis.98 Alendronate’s action on osteoclast function is hypothesized to be related to the inhibition of the intracellular mevalonate pathway.99

Administration and Adult Dosage. PO for prevention of osteoporosis in postmenopausal women 5 mg/day or 35 mg once weekly;100,101 PO for treatment of osteoporosis in postmenopausal women 10 mg/day or 70 mg once weekly;101–104 PO for osteoporosis in men 10 mg/day;105 PO for glucocorticoid-induced
osteoporosis in men and women 5 mg/day, except for postmenopausal women not receiving estrogen, for whom the recommended dosage is 10 mg/day;\textsuperscript{106} PO for Paget’s disease of bone in men and women 40 mg/day for 6 months.\textsuperscript{107}

**Special Populations. Pediatric Dosage.** (<18 yr) Safety and efficacy not established.

**Geriatric Dosage.** Same as adult dosage.

**Other Conditions.** Dosage adjustment is unnecessary in patients with hepatic impairment or Cl\(_r\) >35 mL/min.

**Dosage Forms.** Tab 5, 10, 35, 40, 70 mg.

**Patient Instructions.** Take alendronate with 180–240 mL (6 to 8 fluid ounces) of water on an empty stomach in the morning at least 30 minutes before any food, beverage, or other medicines. Food and beverages, including mineral water, coffee, tea, or juice, decrease alendronate absorption. Antacids or calcium or vitamin supplements also decrease the absorption of alendronate. Do not lie down for 30 minutes after taking alendronate.

**Missed Doses.** If you miss a dose of this medicine, resume your usual schedule the next morning. Do not double doses. If you are taking your dose once weekly and miss a scheduled dose, take your weekly dose the next day. Do not take two tablets on the same day.

**Pharmacokinetics. Fate.** Oral bioavailability is 0.9–1.8% in animals.\textsuperscript{108,109} Plasma protein binding to albumin is 70–80% in animals; \(V_d\) is 28 L exclusive of bone distribution. It is excreted renally.

\(t_{1/2}\). Plasma concentrations fall by >95% within 6 hr after IV administration. The terminal half-life in humans is estimated to exceed 10 yr, reflecting skeletal release of alendronate.

**Adverse Reactions.** Mild, transient falls in serum calcium and phosphate have been reported. Dose-related abdominal pain, dyspepsia, constipation, diarrhea, esophageal ulcer, dysphagia, and abdominal distention can occur. Postmarketing surveillance showed an increased risk of erosive esophagitis, some with ulcerations, primarily in patients who did not comply with recommended administration guidelines.\textsuperscript{110-112} Ulcerations are occasionally severe, necessitating hospitalization.\textsuperscript{110}

**Contraindications.** Abnormalities of the esophagus that delay esophageal emptying such as stricture or achalasia; inability to stand or sit upright for at least 30 min; hypocalcemia.

**Precautions.** The drug must be taken with 180–240 mL of water and patients must not lie down for at least 30 min after oral administration. Avoid use with Cl\(_r\) <35 mL/min.

**Drug Interactions.** Concomitant calcium-containing products interfere with alendronate absorption and should be administered no sooner than 30 min after a dose. Concomitant use with IV ranitidine results in a 2-fold increase in bioavailability. Avoid concomitant ingestion with food, orange juice, or caffeine.

Pamidronate is a nitrogen-containing bisphosphonate that is about 100 times as potent as etidronate in inhibiting bone resorption in the rat. 97 (See Alendronate.)

Administration and Adult Dosage. IV for hypercalcemia of malignancy (moderate hypercalcemia: corrected serum calcium of 12–13.5 mg/dL) 60–90 mg. 113 The 60 mg dose is given as an initial, single-dose infusion over at least 4 hr, and the 90 mg dose must be given by an initial, single-dose infusion over 24 hr; (severe hypercalcemia: corrected serum calcium >13.5 mg/dL) 90 mg as an initial, single-dose infusion over 24 hr.

Corrected Serum Calcium = Serum Calcium in mg/dL + (0.8 × [4 – Serum Albumin in g/dL]).

IV for Paget’s disease 30 mg/day as a 4-hr infusion on 3 consecutive days for a total of 90 mg. IV for osteolytic bone lesions of multiple myeloma 90 mg once monthly as a 4-hr infusion. 114 IV for osteolytic bone metastases of breast cancer 90 mg q 3–4 weeks as a 2-hr infusion. 115,116

Special Populations. Pediatric Dosage. Safety and efficacy not established.

Geriatric Dosage. Same as adult dosage.

Other Conditions. Although pharmacokinetic data are lacking, dosage adjustment appears unnecessary in patients with hepatic impairment. Renal clearance is correlated with Clcr and renally impaired patients excrete less unchanged drug. 117 In patients receiving intermittent therapy, dosage adjustment is probably unnecessary.

Dosage Forms. Inj 30, 90 mg.

Pharmacokinetics. Fate. Oral bioavailability is estimated to be 0.3%. 109 Pamidronate is not metabolized and eliminated exclusively by renal excretion. About 46 ± 16% of the drug is excreted unchanged in the urine within 120 hr. 

1/2 2.5 hr.

Adverse Reactions. Generalized malaise has occurred. Hypocalcemia has been reported in patients with hypercalcemia and Paget’s disease. Abdominal pain, anorexia, constipation, nausea, and vomiting have been reported in at least 15% of patients receiving pamidronate for hypercalcemia and 5% of patients with Paget’s disease. Transient mild temperature elevation (1°C) has occurred. Redness, swelling/induration, and pain on palpation can occur at the IV insertion site.

Precautions. Obtain laboratory tests at the start of therapy. (See Parameters to Monitor.) Use with caution in patients with Clcr >5 mg/dL.

Notes. Do not mix pamidronate with any calcium-containing products. (See Bisphosphonates Comparison Chart.)
<table>
<thead>
<tr>
<th>DRUG</th>
<th>DOSAGE FORMS</th>
<th>INDICATIONS</th>
<th>DOSAGE</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Alendronate Sodium</strong></td>
<td>Tab 5, 10, 35, 40, 70 mg.</td>
<td>Osteoporosis treatment and prevention; corticosteroid-induced osteoporosis; Paget’s disease.</td>
<td>(See monograph.)</td>
</tr>
<tr>
<td><em>Fosamax</em></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Etidronate Disodium</strong></td>
<td>Tab 200, 400 mg Inj 300 mg.</td>
<td>Hypercalcemia of malignancy; Paget’s disease; heterotropic ossification.</td>
<td>IV for hypercalcemia 7.5 mg/kg/day over ≥2 hr for 3 days, followed by PO 20 mg/kg/day for 30 days prn. PO for Paget’s disease 5–10 mg/kg/day for up to 6 months or 11–20 mg/kg/day for up to 3 months. PO for heterotropic ossification 20 mg/kg/day for 1 month before and 3 months after hip replacement or, if caused by spinal cord injury, 20 mg/kg/day for 2 weeks, then 10 mg/kg/day for 10 weeks.</td>
</tr>
<tr>
<td><em>Didronel</em></td>
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<tr>
<td><strong>Pamidronate Disodium</strong></td>
<td>Inj 30, 90 mg.</td>
<td>Hypercalcemia of malignancy; Paget’s disease; osteolytic bone lesions and metastases.</td>
<td>(See monograph.)</td>
</tr>
<tr>
<td><em>Aredia</em></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Risedronate</strong></td>
<td>Tab 5, 30 mg.</td>
<td>Treatment and prevention of osteoporosis; Paget’s disease.</td>
<td>PO for osteoporosis 5 mg/day. PO for Paget’s disease 30 mg/day for 2 months.</td>
</tr>
<tr>
<td><em>Actonel</em></td>
<td></td>
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<tr>
<td><strong>Tiludronate</strong></td>
<td>Tab 240 mg (200 mg of free acid).</td>
<td>Paget’s disease.</td>
<td>PO 400 mg qid for 3 months.</td>
</tr>
<tr>
<td><em>Skelid</em></td>
<td></td>
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</tr>
<tr>
<td><strong>Zolendronate</strong></td>
<td>Injection.</td>
<td>Hypercalcemia of malignancy; osteolytic bone lesions of metastatic breast cancer and multiple myeloma; Paget’s disease.</td>
<td>IV for hypercalcemia of malignancy 0.02–0.04 mg/kg; IV for osteolytic bone lesions 1–3 mg; IV for Paget’s disease 0.2–0.4 mg.</td>
</tr>
<tr>
<td><em>Zometa (Investigational—Novartis)</em></td>
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From references 101–107, 109, 113–116, and 118–127.
Gout Therapy

ALLOPURINOL  Zyloprim, Various

Pharmacology. Allopurinol, a structural analogue of the purine base hypoxanthine, competitively inhibits xanthine oxidase. This reduces serum and urinary uric acid levels by blocking the conversion of hypoxanthine and xanthine to uric acid and decreasing urine synthesis.128–130

Administration and Adult Dosage. PO for control of gout 100 mg/day initially, increasing in 100 mg/day increments at weekly intervals until a serum uric acid level of ≤6 mg/dL is attained. PO for maintenance of mild gout 200–300 mg/day in single or divided doses; PO for maintenance of moderately severe tophaceous gout 400–600 mg/day, to a maximum of 800 mg/day for resistant cases. Give dosages that exceed 300 mg/day in divided doses. Give prophylactic colchicine 0.5–1.2 mg/day and/or an NSAID starting before allopurinol and continuing for 1 to several months after initiation of therapy because of an initial increased risk of gouty attacks.128–131 A fluid intake sufficient to yield a daily urinary output of at least 2 L and the maintenance of a neutral or slightly alkaline urine are desirable. In transferring from a uricosuric agent to allopurinol, reduce the uricosuric dosage over several weeks while gradually increasing the dosage of allopurinol. PO or IV for secondary hyperuricemia associated with vigorous treatment of malignancies 600–800 mg/day for 2–3 days is advisable with a high fluid intake and then reduce to 300 mg/day. Start at least 2–3 days (preferably 5 days) before initiation of cancer therapy. Discontinue when the potential for uric acid overproduction is no longer present.132,133 IV should be used only in those who do not tolerate PO allopurinol. PO for recurrent calcium oxalate stones in hyperuricosuria 200–300 mg/day adjusted based on control of hyperuricosuria.

Special Populations. Pediatric Dosage. PO for secondary hyperuricemia associated with malignancies (<6 yr) 150 mg/day; (6–10 yr) 300 mg/day; alternatively, 2.5 mg/kg q 6 hr, to a maximum of 600 mg/day. Start at least 2–3 days (preferably 5 days) before cancer therapy. Evaluate response 48 hr after cancer therapy is started and adjust dosage as needed.

Geriatric Dosage. Lower dosage might be required in some patients because of the age-related decrease in renal function.

Other Conditions. In renal impairment, reduce initial dosage as follows: (Clcr 80 mL/min) 250 mg/day; (Clcr 60 mL/min) 200 mg/day; (Clcr 40 mL/min) 150 mg/day; (Clcr 20 mL/min) 100 mg/day; (Clcr 10 mL/min) 100 mg q 2 days; (Clcr <10 mL/min) 100 mg q 3 days.134 Base subsequent dosage adjustment on serum uric acid levels.

Dosage Forms. Tab 100, 300 mg; Inj 500 mg.

Patient Instructions. This drug may be taken with food, milk, or an antacid to minimize stomach upset. Adults should drink at least 10–12 full glasses (each containing 8 fluid ounces) of fluid each day. Avoid large amounts of alcohol (can increase uric acid in blood) or vitamin C (can increase the possibility of kidney stones by making the urine more acidic). Report any skin rash, painful urination,
blood in urine, eye irritation, swelling of lips or mouth, itching, chills, fever, sore throat, nausea, or vomiting while taking this drug. Allopurinol can cause drowsiness; use caution while driving or performing other tasks requiring alertness, coordination, or physical dexterity.

**Missed Doses.** Take this drug at regular intervals. If you miss a dose, take it as soon as you remember. Do not double the dose or take extra.

**Pharmacokinetics.**

**Onset and Duration.** A measurable decrease in uric acid occurs in 2–3 days; normal serum uric acid is achieved in 1–3 weeks.

**Fate.** Well absorbed orally (67–81%) but rectal absorption is poor (0–6% of oral bioavailability). Rapidly oxidized to oxypurinol, an active, but less potent, inhibitor of xanthine oxidase. Protein binding of allopurinol or oxypurinol is negligible.\(^\text{165}\) Allopurinol \(V_d\) is 1.5 ± 0.7 L/kg, \(Cl\) is 0.77 ± 0.22 L/hr/kg; oxypurinol \(V_d\) is about 1.6 L/kg.\(^\text{135,165}\) Oxypurinol and allopurinol are excreted unchanged in urine in a ratio of about 10:1.\(^\text{165}\)

\(t_\text{1/2}\). (Allopurinol) 1.4 ± 0.4 hr; (oxypurinol) 19.7 ± 7.3 hr with normal renal function, 5–10 days in renal failure.\(^\text{165}\)

**Adverse Reactions.** A mild maculopapular skin rash occurs in about 2% of patients, but the percentage increases to about 20% with concurrent ampicillin. These rashes might not recur if allopurinol is stopped and restarted at a lower dosage and oral desensitization to minor rashes from allopurinol in patients has been effective.\(^\text{130,136,137}\) Exfoliative, urticarial, purpuric, and erythema multiform lesions also are reported occasionally. These more severe reactions require drug discontinuation because severe hypersensitivity reactions such as vasculitis, toxic epidermal necrolysis, Stevens–Johnson syndrome, renal impairment, and hepatic damage can result. An occasional hypersensitivity syndrome (frequently marked by fever, rash, hepatitis, renal failure, and eosinophilia) has a mortality rate reportedly as high as 27%. It can begin 1 day–2 yr (average 6 weeks) after start of therapy and appears related to pre-existing renal dysfunction, elevated oxypurinol serum levels, or concurrent thiazide or other diuretic therapy.\(^\text{134,138,139}\) Occasionally, nausea, vomiting, abdominal pain, and drowsiness occur. Rarely, alopecia, cataract formation, hepatotoxicity, bone marrow depression, leukopenia, leukocytosis, or renal xanthine stones occur.\(^\text{140}\)

**Contraindications.** Children (except for hyperuricemia secondary to malignancy). Do not restart the drug in patients who have developed severe reactions. (See Adverse Reactions.)

**Precautions.** Pregnancy; lactation. Use with caution and in reduced dosage in renal impairment. Adjust dosage conservatively in patients with impaired renal function who are on a diuretic concomitantly.\(^\text{134,139}\)

**Drug Interactions.** Diuretics can contribute to allopurinol toxicity, although a cause-and-effect relationship has not been established. Allopurinol markedly increases the toxicity of oral azathioprine and mercaptopurine. Allopurinol can increase the risk of hypersensitivity reactions to captopril, ampicillin skin rashes, bone marrow suppression caused by cyclophosphamide, neurotoxicity of vidarabine, and nephrotoxicity of cyclosporine. Allopurinol also can increase the effect of some oral anticoagulants but probably not that of warfarin. Large doses (600
mg/day) of allopurinol can increase theophylline serum levels. Concurrent use of salicylate for its antirheumatic effect does not compromise the action of allopurinol. Uricosuric agents can increase the excretion and decrease the effect of allopurinol.

**Parameters to Monitor.** Monitor serum uric acid levels; pretreatment 24-hr urinary uric acid excretion. Periodically determine liver function (particularly in patients with pre-existing liver disease). Monitor renal function tests and CBC, especially during the first few months of therapy. Renal function is particularly important in patients on concurrent diuretic therapy.

**Notes.** Allopurinol is the drug of choice for patients with impaired renal function who respond poorly to uricosuric agents; however, these patients should be monitored closely because of increased frequency of adverse reactions. Current data do not support the routine treatment of asymptomatic hyperuricemia in patients other than those receiving vigorous treatment of malignancies and in marked overexcreters. Allopurinol has been used investigentially to reduce tissue damage during coronary artery bypass surgery, for organ transplantation storage solutions, and in the treatment of leishmaniasis. Because of limited studies showing very poor or no absorption of extemporaneously compounded allopurinol suppositories, this dosage form is not recommended. Although preliminary reports indicated that extemporaneously prepared allopurinol mouthwash might be effective in protecting against fluorouracil-induced mucositis, one well-controlled clinical trial found it ineffective for this indication, and it is not recommended.

**Pharmacology.** Colchicine is an anti-inflammatory agent relatively specific for gout, with activity probably because of the impairment of leukocyte chemotaxis, mobility, adhesion and phagocytosis, and a reduction of the lactic acid production resulting from a decrease in urate crystal deposition.

**Administration and Adult Dosage.** PO for acute gout 1–1.2 mg initially at the first warning of an attack, then 0.5–1.2 mg q 1–2 hr until pain is relieved or GI toxicity occurs (ie, nausea, vomiting, stomach pain, or diarrhea), to a maximum total dosage of 4–8 mg. Pain and swelling typically abate within 12 hr and usually are gone in 24–48 hr. An interval of 3 days is advised if a second course is required. PO for prophylaxis in chronic gout 0.5–1.8 mg/day or every other day depending on severity; divided doses are preferred with higher dosages. PO for surgical prophylaxis in patients with gout 0.5–0.6 mg tid, 3 days before and after surgery. Slow IV for acute gout (if patient cannot take oral preparation) 1–2 mg initially, diluted (if desired) in nonbacteriostatic NS, over 2–5 min, then 0.5 mg q 6–24 hr prn, to a maximum of 4 mg in 24 hr, or a maximum 4 mg for a single course of treatment. Some clinicians recommend a single IV dose of 3 mg over 5 min; others recommend an initial dose of ≤1 mg, then 0.5 mg 1–2 times daily prn. If pain recurs, give IV 1–2 mg/day for several days; however, no more colchicine should be given by any route for at least 7 days after a full course (4 mg) of IV therapy. IV colchicine is very irritating and extravasa-
tion must be avoided to prevent tissue and nerve damage; change to oral therapy as soon as possible. Do not administer by SC or IM routes. (See Notes.)

Special Populations. Geriatric Dosage. Reduce the maximum IV colchicine dosage to 2 mg, with at least 3 weeks between courses, and lower the dosage further if previously maintained on oral colchicine.142

Other Conditions. Reduce the total IV and PO dosage of colchicine in renal impairment in proportion to the remaining renal function.142,143 The dosage of prophylactic colchicine should not exceed 0.5 mg/day with Clr ≤ 50 mL/min, because of increased risk of peripheral neuritis and myopathy.144 Not recommended in patients who require hemodialysis.144

Dosage Forms. Tab 500, 600 µg; Inj 500 µg/mL.

Patient Instructions. You should always have a supply of this drug at hand, and you should take it promptly at the earliest symptoms of a gouty attack. Relief of gout pain or occurrence of nausea, vomiting, stomach pain, or diarrhea indicate that the full therapeutic dosage has been attained and no more drug should be taken. After treatment of an attack, do not take any more colchicine for at least 3 days. Immediately report black tarry stools or bright red blood in the stools, which can indicate gastrointestinal bleeding. Report any tiredness, weakness, numbness, or tingling. Also immediately report sore throat, fever, oral lesions, or unusual bleeding that can be an early sign of a severe, but rare, blood disorder.

Missed Doses. If you are taking this drug at regular intervals, such as daily, and you miss a dose, take it as soon as you remember. If it is about time for the next dose, take that dose only. Do not double the dose or take extra.

Pharmacokinetics. Fate. Rapidly but variably absorbed after oral administration (healthy young adults, 44 ± 17%; elderly, 45 ± 19%), with partial hepatic deacetylation. Plasma protein binding is approximately 50%; extensive leukocyte uptake occurs with levels found for up to 9 days. Distribution after IV administration is triphasic; Vd of the terminal phase is 6.7 ± 1.4 L/kg for healthy young adults and 6.3 ± 2.3 L/kg for the elderly. CI is 0.15 ± 0.02 L/hr/kg for healthy young adults and 0.12 ± 0.01 L/hr/kg for the elderly. Urinary (about 10% unchanged), biliary, and fecal elimination occur.143,145,146

t½. (Healthy young adults) second phase 1.2 ± 0.2 hr; terminal phase 30 ± 6 hr; (elderly) second phase 1.2 ± 0.1 hr; terminal phase 34 ± 8 hr.145

Adverse Reactions. Nausea, vomiting, stomach pain, and diarrhea are frequent and can occur several hours after oral or IV drug administration; discontinue drug at first signs. Prolonged administration occasionally can cause bone marrow depression with agranulocytosis or thrombocytopenia, aplastic anemia, and purpura. Peripheral neuritis and myopathy with characteristically elevated creatine kinase occur occasionally. This reaction is associated with standard (unadjusted) dosage in renal insufficiency and usually resolves in 3–4 weeks after drug withdrawal.130,142-144,147 Alopecia, reversible malabsorption of vitamin B₁₂, and reversible azoospermia occur. Tissue and nerve damage can occur with IV extravasation. Overdosage can cause hemorrhagic gastroenteritis, vascular damage leading to shock, nephrotoxicity, and paralysis. As little as 7 mg has proved fatal, but much larger dosages have been survived.143,148-150
Contraindications. Serious GI, renal, hepatic, or cardiac disorders; combined hepatic, and renal dysfunction; blood dyscrasias.

Precautions. Use with great caution in elderly or debilitated patients, especially those with early manifestations of hepatic, renal, GI, or heart disease. Reduce dosage if weakness, anorexia, nausea, vomiting, stomach pain, or diarrhea occurs.

Drug Interactions. None known.

Notes. Colchicine is most effective when used early in the attack before most WBC chemotaxis takes place. For acute gout, an NSAID or a corticosteroid (systemically or intra-articularly) may be preferred, but daily colchicine often is given for prophylaxis against recurrent gouty attacks before and during the first one to several months of allopurinol or uricosuric treatment. Continuous prophylactic colchicine therapy can be effective in suppressing the acute attacks and renal dysfunction of familial Mediterranean fever. Colchicine therapy also might be effective for primary biliary cirrhosis and certain inflammatory dermatoses.

Pharmacology. Probenecid, a sulfonamide, is an organic acid that inhibits renal tubular reabsorption of urate, thereby increasing the urinary excretion of uric acid and lowering serum urate. Probenecid also interferes with renal tubular secretion of many drugs, causing an increase or prolongation in their serum levels. (See Notes.)

Administration and Adult Dosage. PO for chronic gout 250 mg bid for 1 week, then 500 mg bid (not to be started during an acute attack). Colchicine 0.5–1.2 mg/day or an NSAID started before and continued for 1 to several months after initiation of uricosuric treatment diminishes exacerbation of uricosuric-induced gouty attacks. To prevent hematuria, renal colic, costovertebral pain, and urate stone formation, liberal fluid intake and alkalinization of the urine with 3–7.5 g/day sodium bicarbonate or 7.5 g/day potassium citrate are recommended, at least until serum uric acid levels normalize and tophaceous deposits disappear. If an acute gouty attack is precipitated during therapy, increase the dosage of colchicine or add a corticosteroid or an NSAID to control the attack. (See Precautions.) Decrease daily dosage by 500 mg q 6 months if no acute attacks occur, adjusted to maintain normal serum uric acid levels. PO to prolong penicillin or cephalosporin action 2 g/day in 4 divided doses, except with known renal impairment. PO with procaine penicillin G for uncomplicated gonorrhea 1 g as a single dose. PO with procaine penicillin G for neurosyphilis 2 g/day in 4 divided doses for 10–14 days. PO with cefoxitin for outpatient treatment of pelvic inflammatory disease 1 g as a single dose.

Special Populations. Pediatric Dosage. (<2 yr) contraindicated. PO to prolong penicillin or cephalosporin action (<50 kg) 25 mg/kg initially, then maintain at 40 mg/kg/day or 1.2 g/m²/day in 4 divided doses; (>50 kg) same as adult dosage.

Geriatric Dosage. Same as adult dosage unless renal impairment is present.
Other Conditions. For chronic gout in renal impairment (although probably ineffective when Clcr ≤ 30 mL/min), increase initial dosage of 500 mg bid in 500 mg/day increments q 4 weeks to the dosage that maintains normal serum uric acid levels, to a maximum of 2 g/day in divided doses. Reduce dosage for prolonging penicillin or cephalosporin action in patients with renal impairment.

Dosage Forms. Tab 500 mg.

Patient Instructions. This drug may be taken with food, milk, or an antacid to minimize stomach upset. Drink a large amount (10 to 12 full glasses) of fluids each day and avoid the use of aspirin- or salicylate-containing products unless directed otherwise.

Missed Doses. If you are taking this drug at regular intervals, such as daily, and you miss a dose, take it as soon as you remember. If it is about time for the next dose, take that dose only. Do not double the dose or take extra.

Pharmacokinetics. Fate. Rapidly and completely absorbed from the GI tract; 74–99% plasma protein bound (decreasing with increasing dose), mostly to albumin. Vd is 0.17 ± 0.03 L/kg. Probenecid is extensively metabolized or conjugated, exhibiting Michaelis–Menten elimination; about 40% is excreted in urine as the monoacylglycuronide, <5% as unchanged drug, and the remainder as hydroxylated metabolites, which can have uricosuric activity.

t1/2. Dose dependent (increases with increasing dose): 4.5 ± 0.6 hr with 0.5 g; 12 hr with 2 g.

Adverse Reactions. Headache, nausea, vomiting, urinary frequency, rash, and dizziness occur frequently. Exacerbation of gout, hematuria, renal colic, costovertebral pain, and uric acid stones can occur. Nephrotic syndrome, hepatic necrosis, aplastic anemia, hemolytic anemia (possibly related to G-6-PD deficiency), and severe allergic reactions occur rarely.

Contraindications. Children <2 yr; known blood dyscrasias or uric acid kidney stones; initiation during an acute gouty attack.

Precautions. Hypersensitivity reactions require drug discontinuation. Use with caution in patients with histories of sulfonamide allergy, peptic ulcer, or G-6-PD deficiency. (See Notes.)

Drug Interactions. Salicylates and pyrazinamide antagonize the uricosuric action of probenecid. Probenecid can increase the serum concentration of many drugs, including acyclovir, benzodiazepines, some β-lactams, clofibrate, dapsone, methotrexate, NSAIDs, penicillamine, sulfonamides, sulfonyleureas, thiopeptil, and zidovudine. NSAID clearance might be decreased by competitively inhibiting formation or renal excretion of acylglucuronide metabolites.

Parameters to Monitor. Serum uric acid weekly until stable when treating hyperuricemia; pretreatment 24-hr urinary uric acid excretion. If alkali is administered, periodically determine acid–base balance.

Notes. Current data do not support the treatment of patients with asymptomatic hyperuricemia caused by undersecretion of uric acid. Most useful in symptomatic patients with reduced urinary excretion of urate: <800 mg/day on an unrestricted diet or <600 mg/day on a purine-restricted diet. Ineffective in pro-
longing the half-life of β-lactams that do not undergo renal tubular secretion (eg, cefazidime, ceftriaxone).158

**SULFINPYRAZONE**

**Pharmacology.** Sulfinpyrazone is an analogue of phenylbutazone that lacks anti-inflammatory and analgesic properties. It is a uricosuric agent with a mechanism and site of action resembling those of probenecid. Sulfinpyrazone, like probenecid, interferes with the renal tubular secretion of many drugs. It also has antiplatelet and antithrombotic activities but currently is not used clinically for these indications.128–130,159

**Adult Dosage.** PO as a uricosuric 200–400 mg/day orally in 2 divided doses with meals or milk, increasing over 1 week to a maximum of 800 mg/day with adequate fluid intake and alkalization of the urine. Reduce to the lowest dosage needed to control serum uric acid (as low as 200 mg/day). In elderly, azotemic cardiovascular patients, initiate therapy with 200 mg/day and increase in 200 mg/day increments q 4 days or keep constant for another 4 days depending on Cr, and serum uric acid, to a maximum maintenance dosage of 800 mg/day.160

**Dosage Forms.** Tab 100 mg; Cap 200 mg.

**Pharmacokinetics.** Oral absorption is rapid and complete, with peak serum levels occurring in 1–2 hr. Vd is 0.73 ± 0.23 L/kg; Cl of the parent compound is 0.14 ± 0.044 L/hr/kg. Hepatic metabolism yields four metabolites. The parent compound is mainly responsible for uricosuric activity; the sulfide metabolite produces the antiplatelet effect. Half-lives are 10 ± 1.3 hr (sulfinpyrazone) and 14.3 ± 4.5 hr (sulfide metabolite).161,162

**Adverse Reactions.** Adverse effects are similar to those of probenecid, with occasional acute renal insufficiency, possibly caused by precipitation of uric acid in renal tubules or decrease in prostaglandin synthesis.163,164 **Colchicine** 0.5–1.2 mg/day started before and continued for 1 to several months after initiation of uricosuric treatment diminishes exacerbation of uricosuric-induced gouty attacks. Treat acute exacerbations of gout by increasing the colchicine dosage or adding an NSAID or a corticosteroid. Sulfinpyrazone is contraindicated in patients with peptic ulcers, symptoms of GI inflammation or ulceration, and blood dyscrasias. Avoid sulfinpyrazone in renal insufficiency because it might not be effective. Salicylates can antagonize the action of sulfinpyrazone.

**REFERENCES**


Antiasthatics

**Class Instructions.** **Antiasthmatic Inhalers.** (Aerosols) Remove inhaler cap and hold inhaler upright. Shake inhaler. Tilt your head back and breathe out slowly. To position inhaler, open your mouth with the inhaler 1 to 2 inches away or in your mouth. (For young children and corticosteroid inhalers, use a spacer or holding chamber.) Press down on inhaler to release medication as you start to breathe slowly. Breathe slowly for 3 to 5 seconds. Hold your breath for 10 seconds to allow the medication to reach deep into the lungs. Repeat as directed. (Dry Powder) Close your mouth tightly around the mouthpiece and inhale rapidly. Hold the device horizontally (parallel to the ground) after it has been activated. Do not exhale into the device.

Do not exceed the prescribed dosage. Report if symptoms do not completely clear or the inhaler is required more than prescribed. Clean the mouthpiece weekly with hot water and soap. Store away from heat and direct sunlight. Bronchodilators can cause nervousness, tremors (especially with terbutaline or albuterol), or rapid heart rate. Report if these effects continue after dosage reduction; if chest pain, dizziness, or headache occur; or if asthmatic symptoms are not relieved.

**Missed Doses.** Take missed doses as soon as possible. However, if it is almost time for your next dose, skip the missed dose and go back to your regular schedule. Do not double doses.

**ALBUTEROL SULFATE** Proventil, Ventolin, Volmax, Various

**Pharmacology.** Albuterol is a selective β₂-adrenergic agonist that produces bronchodilation, vasodilation, uterine relaxation, skeletal muscle stimulation, peripheral vasodilation, and tachycardia.¹

**Administration and Adult Dosage.** **Inhal for asthma** (metered-dose inhaler) 90–180 µg (1–2 puffs) q 4–6 hr prn and just before exercise; (inhalation solution) 2.5 mg by nebulization tid–qid; (inhalation capsule) 1–2 inhalation capsules q 4–6 hr or 1 capsule 15 min before exercise. **Inhal for severe bronchospasm** nebulized by compressed air or oxygen 2.5–5 mg (0.5–1 mL of 0.5% in 2–3 mL NS) q 4–6 hr prn (q 1–2 hr under medical supervision). **PO for asthma** 2–4 mg q 6–8 hr, increase as tolerated to a maximum of 32 mg/day; **SR Tab** 4–8 mg q 12 hr, to a maximum of 32 mg/day.

**Special Populations.** **Pediatric Dosage.** **Inhal for asthma** (metered-dose inhaler) (<12 yr) 90–180 µg (1–2 puffs) q 4–6 hr using spacer; (≥12 yr) same as adult dosage; (inhalation solution) (<12 yr) 0.05–0.15 mg/kg q 4–6 hr prn, or (<20 kg) 0.25 mL of 0.5% solution; (>20 kg) 0.5 mL of 0.5% solution to a maximum of
1 mL diluted in 2–3 mL NS q 4–6 hr prn (q 1–2 hr for severe bronchospasm under medical supervision); (≥12 yr) same as adult dosage. **PO for asthma** (2–6 yr) 100–200 μg/kg/dose q 8 hr, to a maximum of 4 mg q 8 hr; (6–12 yr) 2 mg q 6–8 hr, to a maximum of 24 mg/day; (>12 yr) same as adult dosage. **SR Tab** (<12 yr) dosage not established; (>12 yr) same as adult dosage.

**GeriatricDosage.** **Inhal for asthma** same as adult dosage. **PO** 2 mg tid–qid initially, increasing prn to a maximum of 8 mg tid–qid.

**Dosage Forms.*** **Inhal** (metered-dose) 90 μg/puff (200 puffs/inhaler); **Inhal** (metered-dose, HFA, does not contain chlorofluorocarbons as a propellant) (Proventil HFA, Ventolin HFA) 90 μg/puff (200 puffs/inhaler); **Inhal Soln** 0.5% (5 mg/mL), 0.083% (unit dose solution, 3 mL); **Inhal Cap** (Rotacap) 200 μg for use with powder inhaler; **Tab** 2, 4 mg; **Syrup** 0.4 mg/mL; **SR Tab** 4, 8 mg. **Inhal** 90 μg plus ipratropium bromide 18 μg/puff (Combivent); **Inhal Soln** 3 mg plus ipratropium bromide 0.5 mg/3 mL (DuoNeb).

**Patient Instructions.** *(See Class Instructions: Antiasthmatic Inhalers.)*

**Pharmacokinetics.** **Onset and Duration.** (Inhal) onset within 15 min, peak 60–90 min or less; (PO) onset 30–60 min, peak 2–3 hr. Duration 4–6 hr, depending on the dose, dosage form, and clinical condition. *(See Sympathomimetic Bronchodilators Comparison Chart.)*

**Fate.** Peak serum level after 0.15 mg/kg by inhalation is 5.6 μg/L (23 nmol/L). Oral bioavailability is 50% because of hepatic first-pass metabolism; peak after 4 mg tablet is 10 μg/L (42 nmol/L); 50% is excreted in urine as an inactive sulfate conjugate. The drug does not appear to be metabolized in the lung.2

*tk1/2.* (IV) 2–3 hr; apparent half-life is 5–6 hr after oral and up to 7 hr after inhalation because of prolonged absorption.2

**Adverse Reactions.** Dose-related reflex tachycardia from peripheral vasodilation and direct stimulation of cardiac β2-receptors. Tremor, palpitations, and nausea are other dose-related effects that are markedly reduced with aerosol administration. All β2-agonists lower serum potassium concentrations.

**Precautions.** Pregnancy; cardiac disorders including coronary insufficiency and hypertension; diabetes. Excessive or prolonged use may lead to tolerance.

**Drug Interactions.** Concurrent β-blockers may antagonize effects.

**Parameters to Monitor.** Inhalation technique, asthma symptoms, frequency of use, pulmonary function, and heart rate.

**Notes.** A relationship between regular (ie, not prn) use of inhaled β2-agonists and death from asthma has been a concern.3,4 Regardless of whether β2-agonists are directly responsible or simply a marker for more severe asthma, heavy use (>1 canister/month or 12 puffs/day) of these agents should alert clinicians that it is necessary to re-evaluate the patient’s condition. Proventil HFA inhalers use a nonchlorofluorocarbon propellant; drug delivery is similar, but not identical, to Ventolin and Proventil. **Levalbuterol** (Xopenex) is the active L-isomer of albuterol. It is available as solution for inhalation 0.63 mg and 1.25 mg/3 mL.
Pharmacology. Cromolyn stabilizes the membranes of mast cells and other inflammatory cells (e.g., eosinophils), thereby inhibiting release and production of soluble mediators (e.g., histamine, leukotrienes) that produce inflammation and bronchospasm. The mechanism appears to be the inhibition of calcium ion influx through the cell membrane. Cromolyn inhibits the early and late responses to specific allergen and exercise challenges. It also prevents the increase in nonspecific bronchial hyperreactivity that occurs during a specific allergen season in atopic asthmatics.5

Administration and Adult Dosage. **Inhal** for asthma 20 mg qid at regular intervals in nebulizer (1 ampule inhalant solution) or 0.8–1.6 mg qid via a pressurized metered-dose inhaler. Initiate therapy in conjunction with an aerosolized β2-agonist. (See Notes.) **Inhal** for prevention of exercise-induced bronchospasm single dose (as above) just before exercise. **Intranasal** for prophylaxis of allergic rhinitis 5.2 mg/nostril 3–6 times/day at regular intervals. **Ophth** for allergic ocular disorders 1–2 drops (1.6–3.2 mg) in each eye 4–6 times/day at regular intervals. For chronic conditions, the drug must be used continuously to be effective. **PO** for mastocytosis 200 mg qid, 30 min before meals and hs.

Special Populations. **Pediatric Dosage.** **Inhal** (<2 yr) dosage not established; (≥2 yr) same as adult dosage. **Intranasal** or **Ophth** same as adult dosage. **PO** for mastocytosis (term infants–2 yr) 20 mg/kg/day in 4 divided doses, to a maximum of 30 mg/kg/day; (2–12 yr) 100 mg qid, 30 min before meals and hs, increasing, if necessary, to a maximum of 40 mg/kg/day.

**Geriatric Dosage.** Same as adult dosage.

Other Conditions. The therapeutic effect is dose dependent, and patients with more severe disease may require more frequent administration initially. After a patient becomes symptom free, the frequency of administration may be reduced to bid–tid.

Dosage Forms. **Inhal** Soln 10 mg/mL; **Inhal** 800 μg/puff (112, 200 doses/inhaler); **Nasal** Inh 5.2 mg/spray (100, 200 doses/inhaler); **Ophth** Drp 4% (40 mg/mL, 250 drops/container); **PO** Soln 20 mg/mL.

Patient Instructions. (See Class Instructions.) (Asthma) this medication must be used regularly and continuously to be effective. Do not stop therapy abruptly, except on medical advice. Carefully follow directions for inhaler use included with the device. You may mix the nebulizer solution with any bronchodilator inhalant solution that does not contain benzalkonium chloride. (Mastocytosis) dissolve oral capsules in one-half glass (4 fluid ounces) of hot water, add an equal amount of cold water, and drink the entire amount. Do not mix with fruit juice, milk, or foods.

Missed Doses. Take this drug at regular intervals. If you miss a dose, take it as soon as you remember. If it is about time for the next dose, take that dose only. Do not double the dose or take extra.

Pharmacokinetics. **Onset and Duration.** (Asthma) onset within 1 min for prevention of allergen-induced mast cell degranulation; duration dose dependent,
2–5 hr. It may require 4–6 weeks to achieve maximal response, although most asthmatics respond within 2 weeks.

**Fate.** Oral bioavailability is 0.5–1%. Amount absorbed after inhalation depends on the delivery system; about 10% of the dosage for a Spinhaler and <2% with the nebulizer solution. Peak serum levels occur 15–20 min after inhalation. Vd is 0.2 ± 0.04 L/kg; Cl is 0.35 ± 0.1 L/hr/kg. Rapidly excreted unchanged in equal portions in the bile and urine.

\[ t_{1/2} = 22.5 ± 1.6 \text{ min}. \]

**Adverse Reactions.** Mild burning or stinging can occur with ophthalmic solution. Occasionally, headache and diarrhea occur with oral capsules.

**Precautions.** Use with caution in patients with lactose sensitivity (capsules only). Watch for worsening of asthma in patients discontinuing the drug. The ophthalmic solution contains 0.01% benzalkonium chloride; therefore, do not wear soft contact lenses during therapy.

**Drug Interactions.** None known.

**Parameters to Monitor.** Monitor relief of asthmatic symptoms and the proper dosage and inhalation technique. Patient noncompliance or inappropriate inhalation technique often contributes to treatment failure. The measurement of peak expiratory flow rate with a peak flow meter is useful in severe chronic asthma. Periodic standard pulmonary function tests are indicated q 1–6 months.

**Notes.** Comparative studies have shown cromolyn and **theophylline** to be equally effective for the prophylaxis of chronic asthma, although cromolyn produces fewer side effects. The inhalant solution is stable with all β2-agonist and anticholinergic solutions for nebulization, although benzalkonium chloride–free solutions are preferred. The nasal spray is most effective if started 1 week before the allergen season; however, patients receive benefit even if treatment is begun after symptoms occur. Oral cromolyn has been used in the management of GI conditions such as food allergy and irritable bowel syndrome. Cromolyn solution is incompatible with benzalkonium chloride.

**IPRATROPIUM BROMIDE**

**Pharmacology.** Ipratropium is a competitive antagonist of acetylcholine at peripheral, but not central, muscarinic receptors because of its quaternary structure. It is used primarily as a bronchodilator in COPD, emphysema, and bronchitis.

**Administration and Adult Dosage.** Inhal for bronchospasm of COPD (including chronic bronchitis) 36–72 μg (2–4 puffs) qid by metered-dose inhaler, to a maximum of 288 μg (16 puffs)/day. Inhal for acute, severe asthma 500 μg tid–qid by nebulizer. Combivent or extemporaneous ipratropium/albuterol mixtures have the same dosage as above. Nasal spray for rhinorhea of perennial rhinitis 2 sprays (84 μg)/nostril of 0.03% solution bid–tid; Nasal spray for rhinorhea of the common cold 2 sprays (84 μg)/nostril tid–qid for up to 4 days.

**Special Populations.** Pediatric Dosage. (<12 yr) safety and efficacy not established. Inhal (<2 yr) 125 μg/dose by nebulizer, (>2 yr) 18–36 μg (1–2 puffs) q 6–8 hr by metered-dose inhaler, or 250 μg q 6–8 hr by nebulizer has been
used. Nasal spray for rhinorrhea of perennial rhinitis (<6 yr) safety and efficacy not established; (6–11 yr) 1 spray (42 μg)/nostril of 0.03% solution bid–tid; (≥12 yr) same as adult dosage.

**Geriatric Dosage.** Same as adult dosage.

**Dosage Forms.** Inhal 18 μg/puff (200 doses/inhaler); Inhal Soln 200 μg/mL (500 μg/vial); Nasal Spray 0.03, 0.06%. Inhal 18 μg plus 90 μg albuterol/puff (Combivent); 500 μg plus albuterol 3 mg/3 mL (DuoNeb).

**Patient Instructions.** (See Class Instructions: Antiasthmatic Inhalers.) Temporary blurring of vision can occur if the drug is sprayed into eyes.

**Pharmacokinetics.** **Onset and Duration.** Onset 3 min; peak 1–2 hr; duration 4–6 hr, depending on intensity of response.13

**Fate.** Only ≤32% is orally absorbed and <1% of inhaled dose is absorbed.10 Metabolized to eight metabolites, which are excreted in urine and bile.

**t\(^1/2\).** 1.5–4 hr.10

**Adverse Reactions.** Dryness of the mouth. Because of the quaternary nature of the molecule, typical systemic anticholinergic side effects are absent.10,13 With the nasal spray, epistaxis, nasal dryness, dry mouth, or throat and nasal congestion occur in 1–10% of patients. During long-term use, headache, nausea, and upper respiratory tract infections also occur frequently.

**Contraindications.** (Aerosol inhaler) hypersensitivity to soy lecithin, soybeans, or related products.

**Precautions.** Use with caution in narrow-angle glaucoma, prostatic hypertrophy, or bladder neck obstruction.

**Drug Interactions.** None known.

**Parameters to Monitor.** Inhalation technique, asthma symptoms, frequency of use, pulmonary function, and anticholinergic symptoms.

**Notes.** Anticholinergics appear to be as potent bronchodilators as β2-adrenergic drugs in bronchitis and emphysema but less potent in asthma.8,10 Anticholinergics produce an additive bronchodilation with β2-adrenergic agents in severe asthma.10,12 Ipratropium and albuterol nebulizer solutions can be mixed if the mixture is used within 1 hr. Tiotropium bromide (Spiriva—Boehringer-Ingelheim) is similar to ipratropium and is being studied in COPD.

**MONTELUKAST SODIUM**

**Pharmacology.** Montelukast sodium is a selective and orally active leukotriene-receptor antagonist that inhibits the cysteinyll leukotriene CysLT\(_1\) receptor.14,15

**Administration and Adult Dosage.** PO for mild persistent asthma 10 mg/day in the evening.

**Special Populations.** **Pediatric Dosage.** PO for mild persistent asthma (<2 yr) safety and efficacy not established; (2–5 yr) 4 mg chewable tablet every evening; (6–14 yr) 5 mg chewable tablet every evening; (≥15 yr) same as adult dosage.

**Geriatric Dosage.** Same as adult dosage.
Dosage Forms. Chew Tab 4, 5 mg; Tab 10 mg.

Patient Instructions. This drug is used for long-term control and prevention of mild persistent asthma symptoms. Take this medication daily, even when you are having no symptoms, and during periods of worsening asthma. This medication is not for the treatment of acute asthma attack management of exercise-induced bronchospasm. You should have appropriate short-acting $\beta_2$-agonist medication available to treat acute symptoms. Seek medical attention if short-acting inhaled bronchodilators are needed more often than usual, or if the maximum number of inhalations of short-acting bronchodilator treatment prescribed for a 24-hour period is needed.

Missed Doses. Take a missed dose as soon as possible. If it is almost time to take the next dose, skip the missed dose and go back to your regular dosage schedule. Do not double doses.

Pharmacokinetics. Onset and Duration. Duration is 24 hr.$^{14,15}$

Fate. Montelukast is rapidly absorbed after oral administration. Mean oral bioavailabilities are 64% for the film-coated tablet and 73% for the chewable tablet in the fasted state, and 63% with a standard morning meal. Peak concentrations occur 3–4 hr after administration of a 10 mg film-coated tablet and 2–2.5 hr after the 5 mg chewable tablet in fasted adults. Montelukast is $>99\%$ protein bound; $V_{dss}$ is 8–11 L; Cl is 2.7 L/hr. CYP3A4 and 2C9 are involved in the metabolism of montelukast. Montelukast and its metabolites are excreted almost exclusively via the bile.

$t_{1/2}$, 2.7–5.5 hr in healthy young adults.

Adverse Reactions. Generally well tolerated. Adverse events that occur with a frequency of $\geq 2\%$ and more frequently in patients on montelukast than on placebo are diarrhea, laryngitis, pharyngitis, nausea, otitis, sinusitis, and viral infection.

Contraindications. Patients with known aspirin sensitivity should continue to avoid aspirin or other NSAIDs while taking montelukast. Inform phenylketonurics that the chewable tablets contain phenylalanine (a component of aspartame) 0.824 mg/tablet.

Precautions. (See Patient Instructions.) Reduction in systemic corticosteroid dosage in patients on a leukotriene modifier has been followed rarely by eosinophilia, vasculitic rash, worsening pulmonary symptoms, cardiac complications, and/or neuropathy, sometimes presenting as Churg-Strauss syndrome. A causal relationship with leukotriene-receptor antagonists is not established.

Parameters to Monitor. Clinical symptoms of asthma. Appropriate monitoring recommended when systemic corticosteroid reduction is considered.

Notes. Zafirlukast (Accolate) is similar to montelukast but has the disadvantages of twice-daily administration, the need to take on an empty stomach, cases of severe liver damage and several drug–drug interactions; the anticoagulant effect of warfarin is increased by zafirlukast; erythromycin and theophylline decrease zafirlukast serum concentrations, whereas aspirin increases zafirlukast serum concentrations. Interactions with other drugs are not well studied. (See Precautions.) The dosages of zafirlukast are 10 mg bid on a empty stomach in patients 7–11 yr and
20 mg bid on an empty stomach in those ≥12 yr; it is available as 10 and 20 mg tablets.

**NEDOCROMIL SODIUM**

**Pharmacology.** Nedocromil sodium is the disodium salt of a pyranoquinolone dicarboxylic acid that is chemically dissimilar, but pharmacologically similar, to cromolyn sodium. Like cromolyn, nedocromil inhibits the activation of and mediator release from inflammatory cells important in asthma and allergy. Nedocromil appears to have more potent in vitro activity against allergic response than cromolyn.

**Adult Dosage.** *Inhal for asthma* 2 metered-dose actuations qid. In patients under good control with qid administration (ie, patients requiring inhaled or oral β-agonists not more than twice a week), a lower dosage can be tried. First reduce to a tid regimen, then, after several weeks of continued good control, attempt to reduce to a bid regimen. *Ophth for allergic conjunctivitis* 1–2 drops into each eye bid.

**Pediatric Dosage.** (>12 yr) same as adult dosage.

**Dosage Forms.** *Inhal* 16.2 g, containing at least 104 actuations of 2 mg doses (1.75 mg reaches the patient); *Ophth Soln* 2%.

**Pharmacokinetics.** Oral bioavailability is only 2–3%. After inhalation, bioavailability is 5%, with peak serum concentrations occurring in 20–40 min; concentrations fall monoexponentially, with a half-life of 1.5–2.3 hr, reflecting absorption from lungs.

**Adverse Reactions.** Bronchospasm, headache, distinctive taste, nausea, and vomiting occur frequently. In a limited number of trials, nedocromil was effective for long-term prophylaxis of asthma. Like cromolyn, it can decrease bronchial hyperreactivity but is only partly effective in steroid-dependent asthmatics. Nedocromil sodium is intended for regular maintenance treatment and should not be used in acute asthma attacks.

**SALMETEROL XINAFOATE**

**Pharmacology.** Salmeterol is a β2-agonist structurally and pharmacologically similar to albuterol. Salmeterol is intended for regular treatment of reversible airway obstruction and not for immediate symptomatic relief. The place of salmeterol in asthma therapy is being debated, in part because patients in need of regular β2-agonist therapy should be regarded as candidates for an inhaled corticosteroid to treat underlying inflammation. (See Sympathomimetic Bronchodilators Comparison Chart.)

**Administration and Adult Dosage.** *Inhal for asthma prophylaxis or COPD* 42 μg (2 puffs) q 12 hr by metered-dose inhaler or 1 dry powder blister inhaled q 12 hr. *Inhal to prevent exercise-induced bronchospasm* 2 puffs 30–60 min before exercise. (See also Inhaled Corticosteroid Comparison Chart for combination product dosages.)

**Pediatric Dosage.** *Inhal* (Aerosol) (<12 yr) safety and efficacy not established; (≥12 yr) same as adult dosage. (Dry powder) (≥4 yr) same as adult dosage.
**Dosage Forms.** *Inhal* 21 μg/metered-dose puff, in 6.5 g (60 actuations) and 13 g (120 actuations) canisters; *Dry Pwdr Inhal* 50 μg/blister. *Dry Pwdr Inhal* 50 μg plus fluticasone 100, 250, or 500 μg/blister.

**Patient Instructions.** Shake metered-dose canister well before using. For asthma, use this medication regularly every 12 hours. If asthma symptoms occur between doses, use a short-acting inhaler to treat symptoms. If you regularly need more than 4 inhalations of the short-acting inhaler, see your health care provider.

**Pharmacokinetics.** Onset of effective bronchodilation is achieved in 20–30 min; peak effect occurs within 3–4 hr. Bronchodilation lasts for at least 12 hr after inhalation of a single dose of 50 μg. After inhalation, salmeterol is extensively metabolized by hydroxylation, with the majority of a dose being eliminated within 72 hr. About 23% of administered radioactivity was recovered in the urine and 57% in the feces over 168 hr.

**Adverse Reactions.** *(See Albuterol.)*

**Notes.** Formoterol (Foradil—Novartis) is a long-acting β2-adrenergic agonist that is similar to salmeterol in duration but with a more rapid onset. Dosage (≥12 yr) is 12 μg bid for maintenance or 15 min before exercise. It is available as a dry powder for inhalation. A fixed-dose combination with budesonide (Symbicort—Astra Zeneca) is being investigated.
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<th>DOSAGE</th>
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<td>AccuNeb</td>
<td>(unit dose) 0.021%,</td>
<td>Inhal (soln) 0.5%; Inhal (soln) 2.5–5 mg in 2–3 mL NS</td>
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<td>Proventil</td>
<td>0.042%, 0.083%,</td>
<td>q 4–6 hr prn by nebulizer; may use q 1–2 hr prn status asthmatics under medical supervision; (metered-dose) 1–2 puffs q 4–6 hr prn and before exercise; 1 Rota-cap is equivalent to 2 metered-dose puffs PO 2–4 mg q 6–8 hr, to a maximum of 32 mg/day SR Tab 4–8 mg q 12 hr, to a maximum of 32 mg/day.</td>
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<td>Ventolin</td>
<td>(metered-dose)</td>
<td>Inhal (soln) 0.05–0.15 mg/kg in 2–3 mL NS</td>
<td>q 4–6 hr prn by nebulizer, may use q 1–2 hr prn or 0.5 mg/kg/hr continuously nebulized for status asthmatics under medical supervision; (metered-dose) 1–2 puffs q 4–6 hr prn and before exercise PO 0.1–0.2 mg/kg q 6–8 hr, to a maximum of 24 mg/day.</td>
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<td>Volmax</td>
<td>90 µg/puff; (Rotacaps) 200 µg/cap</td>
<td>Inhal (soln) 0.05–0.15 mg/kg in 2–3 mL NS</td>
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<td>Various</td>
<td>SR Tab 4, 8 mg</td>
<td>Inhal (soln) 0.05–0.15 mg/kg in 2–3 mL NS</td>
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<td>Syrup 0.4 mg/mL</td>
<td>q 4–6 hr prn by nebulizer, may use q 1–2 hr prn or 0.5 mg/kg/hr continuously nebulized for status asthmatics under medical supervision; (metered-dose) 1–2 puffs q 4–6 hr prn and before exercise PO 0.1–0.2 mg/kg q 6–8 hr, to a maximum of 24 mg/day.</td>
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<td>Tab 2, 4 mg.</td>
<td>Inhal (soln) 0.05–0.15 mg/kg in 2–3 mL NS</td>
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<td>q 4–6 hr prn by nebulizer, may use q 1–2 hr prn or 0.5 mg/kg/hr continuously nebulized for status asthmatics under medical supervision; (metered-dose) 1–2 puffs q 4–6 hr prn and before exercise PO 0.1–0.2 mg/kg q 6–8 hr, to a maximum of 24 mg/day.</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

(continued)
<table>
<thead>
<tr>
<th>DRUG</th>
<th>DOSAGE FORMS</th>
<th>DOSAGE</th>
<th>RECEPTOR SELECTIVITY</th>
<th>RELATIVE POTENCY</th>
<th>DURATION OF ACTION BY INHALATION (HR)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Bitolterol</td>
<td>Inhal (metered-dose) 0.37 mg/puff; (soln) 0.2%.</td>
<td>1–3 puffs 4–6 hr pm.</td>
<td>+ 2.5 4–8</td>
<td>++</td>
<td></td>
</tr>
<tr>
<td>Tornalate</td>
<td>Inhal (metered-dose) 0.2%.</td>
<td>1–2 puffs 4–6 hr pm.</td>
<td>+++</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Epinephrine</td>
<td>Inhal (soln) 2.25% (racemic); Inj 0.1, 1 mg/mL.</td>
<td>SC 0.2–0.5 mg q 20 min–4 hr pm.</td>
<td>+++</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Adrenalin</td>
<td>SC 0.01 mL/kg of 1:1000 q 15–20 min for 2 doses, then q 4 hr pm.</td>
<td>Inhal not recommended.</td>
<td>+++</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Various</td>
<td>Inhal (soln) 0.1, 1 mg/mL.</td>
<td>Foradil 12 µg bid, (≥5 yr) same as adult dosage.</td>
<td>+++</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Formoterol</td>
<td>Inhal (dry pwdr) 12 µg.</td>
<td>Inhal 12 µg bid, or 15 min before exercise pm. (≥5 yr) same as adult dosage.</td>
<td>++</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Foradil</td>
<td>Inh (soln) 2.5–10 mg diluted 1.3 in NS q 2–4 hr pm.</td>
<td>Inhal (soln) 0.1–0.2 mg/kg q 2–4 hr pm.</td>
<td>++</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Isoetharine</td>
<td>Inh (soln) 1%.</td>
<td>Inhal (soln) 0.1–0.2 mg/kg q 2–4 hr pm.</td>
<td>+++</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Various</td>
<td>Inh (soln) 0.5 1%; (metered-dose) 80, 103 µg/puff Inj 0.02, 0.2 mg/mL.</td>
<td>Not recommended because of short duration and lack of selectivity.</td>
<td>++++</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Isoproterenol</td>
<td>Inh (soln) 0.5 1%; (metered-dose) 80, 103 µg/puff Inj 0.02,</td>
<td>Not recommended because of short duration and lack of selectivity.</td>
<td>++++</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

(continued)
<table>
<thead>
<tr>
<th>DRUG</th>
<th>DOSAGE FORMS</th>
<th>DOSAGE</th>
<th>RECEPTOR SELECTIVITY(^b)</th>
<th>RELATIVE POTENCY(^c)</th>
<th>DURATION OF ACTION BY INHALATION (HR)(^d)</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Metaproterenol</strong></td>
<td><strong>Adult</strong> Inh (soln) 0.4% (unit dose 10 mg), q 2–4 hr prn; (q 1–2 hr under medical supervision); (metered-dose) 1–3 puffs q 4–6 hr prn and before exercise; PO 20 mg 3–4 times/day.</td>
<td><strong>Pediatric</strong> Inh (soln) 5–15 mg q 2–4 hr prn up to 15 mg q 2–4 hr pm; (metered-dose) 1–2 puffs pm and before exercise; PO 0.5 mg/kg q 4–6 hr, increase by 0.25 mg/kg as tolerated.</td>
<td>++</td>
<td>++</td>
<td>1</td>
</tr>
<tr>
<td><strong>Alupent</strong></td>
<td>0.6% (unit dose 15 mg), 5%; (metered-dose) 0.65 mg/puff Syrup 2 mg/mL Tab 10, 20 mg.</td>
<td>Inh (soln) q 2–4 hr prn up to 15 mg q 2–4 hr pm; (metered-dose) 1–2 puffs pm and before exercise; PO 0.5 mg/kg q 4–6 hr, increase by 0.25 mg/kg as tolerated.</td>
<td>0.25–0.5 mg/kg</td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Pirbuterol</strong></td>
<td><strong>Maxair</strong> Inh (metered-dose) 0.2 mg/puff. q 4–6 hr and before exercise.</td>
<td><strong>Serevent</strong> Inh (metered-dose) 2 puffs q 12 hr¹ Inh (dry pwdr) 1 blister q 12 hr.</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td><strong>Serevent</strong> Inh (aerosol) (&lt;12 yr) not established; (≥12 yr) same as adult dosage; (dry powder) (≥4 yr) same as adult dosage.</td>
<td>+</td>
<td>++++</td>
</tr>
</tbody>
</table>

\(^{a}\) Relative Potency

\(^{b}\) Receptor Selectivity

\(^{c}\) Duration of Action by Inhalation (HR)

\(^{d}\) Adult Pediatric
### SYMPATHOMIMETIC BRONCHODILATORS COMPARISON CHART (continued)

<table>
<thead>
<tr>
<th>DRUG</th>
<th>DOSAGE FORMS</th>
<th>DOSAGE</th>
<th>RECEPTOR SELECTIVITY</th>
<th>DURATION OF RELATIVE POTENCY</th>
<th>ACTION BY INHALATION (HR)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Terbutaline</td>
<td>Inhal (metered-dose)</td>
<td>Inhal 1–3 puffs q 4–6 hr pm; 5–7 mg undiluted by nebulizer q 4–6 hr pm; SC 0.25–0.5 mg q 2–6 hr pm; PO 5 mg q 6–8 hr.</td>
<td>+++</td>
<td>2.5</td>
<td>4–8</td>
</tr>
<tr>
<td>Brethaire</td>
<td>0.2 mg/puff</td>
<td>Inhal 1–2 puffs q 4–6 hr pm; 0.1–0.3 mg/kg</td>
<td>+</td>
<td>2.5</td>
<td>4–8</td>
</tr>
<tr>
<td>Brethine</td>
<td>Inj 1 mg/mL</td>
<td>q 4–6 hr pm</td>
<td>+</td>
<td>2.5</td>
<td>4–8</td>
</tr>
<tr>
<td>Bricanyl</td>
<td>Tab 2.5, 5 mg.</td>
<td>+</td>
<td>2.5</td>
<td>4–8</td>
<td></td>
</tr>
</tbody>
</table>

#### COMBINATION PRODUCTS

<table>
<thead>
<tr>
<th>Albuterol and Ipratropium</th>
<th>Inhal (metered-dose) abuterol 90 µg plus ipratropium 18 µg/puff. Inhal Soln abuterol 3 mg plus ipratropium 0.5 mg/3 mL.</th>
<th>Inhal 2 puffs qid to a maximum of 12 puffs/day.</th>
<th>—</th>
<th>—</th>
<th>—</th>
</tr>
</thead>
<tbody>
<tr>
<td>Combivent</td>
<td>Duoneb</td>
<td></td>
<td>---</td>
<td>---</td>
<td>---</td>
</tr>
</tbody>
</table>

(continued)
### SYMPATHOMIMETIC BRONCHODILATORS COMPARISON CHART (continued)

<table>
<thead>
<tr>
<th>DRUG</th>
<th>DOSAGE FORMS</th>
<th>DOSAGE</th>
<th>RECEPTOR SELECTIVITY&lt;sup&gt;a&lt;/sup&gt;</th>
<th>DURATION OF RELATIVE POTENCY&lt;sup&gt;b&lt;/sup&gt;</th>
<th>ACTION BY INHALATION (HR)&lt;sup&gt;d&lt;/sup&gt;</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>Adult</td>
<td>Pediatric&lt;sup&gt;a&lt;/sup&gt;</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Salmeterol and</td>
<td>Inhal (dry pwdr)</td>
<td>Inhal 1 inhal bid.</td>
<td>(≥12 yr) Same as adult dosage.</td>
<td>—</td>
<td></td>
</tr>
<tr>
<td>Fluticasone 50 mg</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Advair Diskus 100, 250 or 500 µg/inhal</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

<sup>a</sup>Minimal effect; ++++ = Pronounced effect.

<sup>b</sup>β<sub>2</sub>-selectivity does not equate to bronchoselectivity; β<sub>2</sub>-stimulation produces reflex tachycardia from vasodilation as well as stimulation of cardiac β<sub>2</sub>-receptors.

<sup>c</sup>Molar potency relative to metaproterenol; large numbers indicate more potent compounds.

<sup>d</sup>Onset and duration data apply to aerosol therapy only. Duration of bronchodilation only applies to otherwise stable asthmatics and is not applicable to acute severe asthma or protection from severe provocation (eg, allergen, exercise, ozone). Duration may be shorter during acute exacerbation or with long-term therapy because of downregulation of β-receptors (tolerance). Oral tablets (especially SR tablets) and syrups are slower in onset but may be slightly longer acting than aerosols.

<sup>e</sup>Biltolterol is a prodrug converted in the body to colterol, the active drug, which is more potent than isoproterenol; the relative potency value because of incomplete conversion.

<sup>f</sup>For prophylaxis only; acute attacks must be treated with a short-acting agent.

<sup>g</sup>Use injectable solution; not a labeled indication.
Pharmacology. Theophylline directly relaxes smooth muscles of bronchial airways and pulmonary blood vessels to act as a bronchodilator and pulmonary vasodilator. It is also a diuretic, coronary vasodilator, cardiac stimulant, and cerebral stimulant; it improves diaphragmatic contractility; and it lessens diaphragmatic fatigue. The exact cellular mechanism of smooth muscle relaxation is unknown, but intracellular calcium sequestration, inhibition of specific phosphodiesterase isozymes, adenosine-receptor antagonism, and stimulation of endogenous catecholamine release have been postulated to play a role. Aminophylline is the ethylenediamine salt of theophylline.

Administration and Adult Dosage. PO (theophylline) or IV (aminophylline) for acute asthma symptoms in the emergency department or in the hospital no longer recommended because it appears to provide no additional benefit over optimal inhaled β2-agonist therapy and might increase adverse effects; addition of IV theophylline to other therapies in hospitalized adults remains controversial.

The following dosages have been used: 5 mg/kg (6 mg/kg aminophylline), if patient has taken no theophylline in previous 24 hr. In emergencies, 2.5 mg/kg (3 mg/kg aminophylline) may be given if an immediate serum level cannot be obtained. Each 1 mg/kg (1.25 mg/kg aminophylline) results in about a 2 mg/L increase in serum theophylline. Infuse IV aminophylline no faster than 25 mg/min. Maintenance dosage (see Theophylline Dosage Adjustment Chart.) PO for chronic asthma (theophylline) (see Theophylline Dosage Adjustment Chart.) Adjust dosage to achieve serum concentration of 5–15 mg/L.

IM, PR Supp not recommended.

Special Populations. All dosage recommendations are based on the average theophylline clearance for a given population group. There is a wide interpatient variability (often >2-fold) within all patient groups. Therefore, it is essential that serum concentrations be monitored in all patients. If no doses have been missed or extra doses have been taken during the previous 48 hr, and if peak serum concentrations have been obtained (1–2 hr after liquid or plain uncoated tablet and 4–6 hr after most SR products), adjust dosage using the Theophylline Dosage Adjustment Chart.

Pediatric Dosage. PO or IV for acute symptoms not recommended for children hospitalized for severe asthma. PO for chronic asthma (theophylline) (<1 yr) 0.2 × (age in weeks) + 5 = dosage in mg/kg/day. IM, PR Supp not recommended.

Geriatric Dosage. Not established, but the elderly as a group have slower hepatic clearance. Therefore, use lower initial doses and monitor closely for response and adverse reactions.

Other Conditions. Many factors can alter theophylline dosage requirements. (See Precautions and Factors Affecting Serum Theophylline Concentrations Chart.) Use IBW for dosage calculations in obese patients.
Dosage Forms.  (See Theophylline Products Comparison Chart.)

Patient Instructions. Do not chew or crush sustained-release tablets or capsules. Take at equally spaced intervals around the clock. Report any nausea, vomiting, gastrointestinal pain, headache, or restlessness. Contents of sustained-release bead-filled capsules may be mixed with a vehicle (applesauce or jam) and swallowed without chewing for patients who have difficulty swallowing capsules. Take Theo-24 and Uniphyl products at least 1 hour before meals to avoid too rapid absorption of the drug.

Missed Doses. Take the missed dose as soon as possible. However, if it is almost time to take the next dose, skip the missed dose and go back to the regular dosage schedule. Do not double doses. Do not have your theophylline levels measured until you have missed no doses for 3 days.

Pharmacokinetics. Onset and Duration. IV onset within 15 min with loading dose. Serum Levels. Well correlated with clinical effects: therapeutic is 10–15 mg/L (56–83 μmol/L); however, improvement in respiratory function can be observed...
with serum concentrations of 5 mg/L (28 μmol/L). A serum concentration of 5–10 mg/L is often adequate for treatment of neonatal apnea. Toxicity increases at levels >20 mg/L. (See Adverse Reactions.)

**Fate.** Plain uncoated tablets and solution are well absorbed orally; enteric-coated tablets and some SR dosage forms might be unreliably absorbed. Food can affect the rate and extent of absorption of some SR formulations but has minimal effects on rapid-release forms. Food can increase the rate of absorption (Theo-24, Uniphyl), producing dose dumping, or impair absorption (Theo-Dur Sprinkle). Rectal suppository absorption is slow and erratic, and suppositories (including aminophylline) are not recommended under any circumstances. Rectal solutions might result in serum concentrations comparable to oral solution. About 60% is plasma protein bound (less in neonates); \( V_d \) is 0.5 ± 0.1 L/kg (greater in neonates). There can be marked intrapatient variability in clearance over time. Cl also is affected by many factors. (See Precautions.) Smoking increases theophylline metabolism; this effect can last for 3 months–2 yr after cessation of smoking. Clearance progressively increases in infants during the first year of life. Dose-dependent pharmacokinetics in the therapeutic range occur often in children and rarely in adults. In the elderly, clearance declines with age to about 35 mL/hr/kg. Extensively metabolized in the liver to several inactive metabolites; 10% excreted unchanged in the urine.

\( t_1/2 \). 8 ± 2 hr in adult nonsmokers, 4.4 ± 1 hr in adult smokers (1–2 packs per day); 3.7 ± 1.1 hr in children 1–9 yr. In newborn infants, older patients with COPD or cor pulmonale, and patients with CHF or liver disease, the drug can have a half-life >24 hr.

**Adverse Reactions.** Local GI irritation can occur. Reactions occur more frequently at serum concentrations >20 mg/L and include anorexia, nausea, vomiting, epigastric pain, diarrhea, restlessness, irritability, insomnia, and headache. Serious arrhythmias and convulsions (frequently leading to death or permanent brain damage) usually occur at levels >35 mg/L but have occurred at lower concentrations and might not be preceded by less serious toxicity; cardiovascular reactions include sinus tachycardia and life-threatening ventricular arrhythmias with PVCs. Rapid IV administration can cause hypotension, syncope, cardiac arrest (particularly if administered directly into central line), and death. IM administration is painful and offers no advantage.

**Contraindications.** Active peptic ulcer disease; untreated seizure disorder. (Aminophylline) hypersensitivity to ethylenediamine.

**Precautions.** Use with caution in severe cardiac disease, hypoxemia, hepatic disease, acute myocardial injury, cor pulmonale, CHF, fever, viral illness, underlying seizure disorder, migraine, hepatic cirrhosis, and neonates. Do not give with other xanthine preparations. The alcohol in some oral liquid preparations might cause side effects in infants.

**Drug Interactions.** Numerous drugs and conditions can alter theophylline clearance and serum levels. Factors that can decrease serum levels are carbamazepine, charcoal-broiled beef, high-protein/low-carbohydrate diet, isoproterenol (IV),
phenytoin, rifampin, and smoking. Factors that can increase serum levels are allopurinol (>600 mg/day), cimetidine, ciprofloxacin, cor pulmonale, macrolides (eg, erythromycin, troleandomycin), oral contraceptives, and propranolol. (See Factors Affecting Serum Theophylline Concentrations Chart.)

**Parameters to Monitor.** (Inpatients) obtain serum theophylline concentrations before starting therapy (if patient previously took theophylline) and 1, 6, and 24 hr after start of infusion; monitor daily during continuous infusion. (Outpatients) monitor serum concentrations q 6 months, 3–5 days after any dosage change, and whenever there are symptoms of toxicity.²¹,²²

**Notes.** The oral theophylline preparations of choice for long-term use, to achieve sustained therapeutic concentrations and improved compliance, are completely and slowly absorbed SR formulations that are minimally affected by food and pH.²³ (See Theophylline Products Comparison Chart.) Combination products containing ephedrine increase CNS toxicity and have no therapeutic advantage over adequate serum concentrations of theophylline alone. **Diphylline** is chemically related to, but not a salt of, theophylline; the amount of diphylline equivalent to theophylline is unknown. Because its potency is less than that of theophylline and it has a short half-life (2 hr), its dosage is greater than that of theophylline and it must be given more frequently.
<table>
<thead>
<tr>
<th>FACTOR</th>
<th>DECREASES IN THEOPHYLLINE CONCENTRATIONS</th>
<th>INCREASES IN THEOPHYLLINE CONCENTRATIONS</th>
<th>ACTION</th>
</tr>
</thead>
<tbody>
<tr>
<td>Age</td>
<td>↑ metabolism (1–9 yr)</td>
<td>↓ metabolism (&lt;6 months, elderly)</td>
<td>Adjust dosage according to serum concentration.</td>
</tr>
<tr>
<td>Diet</td>
<td>↑ metabolism (high protein)</td>
<td>↓ metabolism (high carbohydrate)</td>
<td>Inform patient that major changes in diet are not recommended while taking theophylline.</td>
</tr>
<tr>
<td>Food</td>
<td>↓ or delays absorption (fatty food)</td>
<td>↑ rate of absorption (SR preparations)</td>
<td>Select theophylline product that is not affected by food.</td>
</tr>
<tr>
<td>Hypoxia, cor pulmonale, decompensated CHF, cirrhosis</td>
<td>—</td>
<td>↓ metabolism</td>
<td>Decrease dosage according to serum concentration.</td>
</tr>
<tr>
<td>Cimetidine</td>
<td>—</td>
<td>↓ metabolism</td>
<td>Use alternative H₂ blocker (eg, famotidine, nizatadine, ranitidine).</td>
</tr>
<tr>
<td>Macrolides: troleandomycin, erythromycin, clarithromycin</td>
<td>—</td>
<td>↓ metabolism</td>
<td>Use alternative antibiotic or decrease theophylline dosage.</td>
</tr>
<tr>
<td>Phenobarbital, phenytoin, carbamazepine</td>
<td>↑ metabolism</td>
<td>—</td>
<td>Increase dosage according to serum concentration.</td>
</tr>
</tbody>
</table>

(continued)
<table>
<thead>
<tr>
<th>FACTOR</th>
<th>DECREASES IN THEOPHYLLINE CONCENTRATIONS</th>
<th>INCREASES IN THEOPHYLLINE CONCENTRATIONS</th>
<th>ACTION</th>
</tr>
</thead>
<tbody>
<tr>
<td>Quinolones: ciprofloxacin, enoxacin</td>
<td>—</td>
<td>↓ metabolism</td>
<td>Use alternative antibiotic or adjust theophylline dosage. Circumvent with levofloxacin if quinolone therapy is required.</td>
</tr>
<tr>
<td>Rifampin</td>
<td>↑ metabolism</td>
<td>—</td>
<td>Increase dosage according to serum concentration.</td>
</tr>
<tr>
<td>Smoking</td>
<td>↑ metabolism</td>
<td>—</td>
<td>Advise patient to stop smoking; increase dosage according to serum concentration.</td>
</tr>
<tr>
<td>Ticlopidine</td>
<td>—</td>
<td>↓ metabolism</td>
<td>Decrease dosage according to serum concentration.</td>
</tr>
<tr>
<td>Viral illness, systemic febrile (eg, influenza)</td>
<td>—</td>
<td>↓ metabolism</td>
<td>Decrease theophylline dosage according to serum concentration. Decrease dosage by 50% if serum concentration is not available.</td>
</tr>
</tbody>
</table>

*This chart is not all inclusive; for other factors, see Cytochrome P450 Interactions and product information.
From reference 20.*
<table>
<thead>
<tr>
<th>PRODUCT</th>
<th>ANHYDROUS THEOPHYLLINE CONTENT</th>
<th>MEASURABLE DOSE INCREMENT (MG)</th>
<th>COMMENTS</th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>RAPIDLY ABSORBED</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Plain Uncoated Tablets</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Various</td>
<td>Tab 100 mg scored.</td>
<td>50</td>
<td>Serum level fluctuations are 459%/117%.</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Tab 125 mg scored.</td>
<td>62.5</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Tab 200 mg scored.</td>
<td>100</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Tab 250 mg scored.</td>
<td>125</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Tab 300 mg scored.</td>
<td>150</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Oral Liquids (Alcohol-Free)</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Aerolate</td>
<td>10 mg/mL.</td>
<td>5</td>
<td>Sugar free.</td>
<td></td>
</tr>
<tr>
<td>Slo-Phyllin 80 Syrup</td>
<td>5.3 mg/mL.</td>
<td>5</td>
<td>Sugar free.</td>
<td></td>
</tr>
<tr>
<td>Intravenous Solution</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Aminophylline&lt;sup&gt;2&lt;/sup&gt;</td>
<td>20 mg/mL.</td>
<td>5</td>
<td>Use rubber-stoppered vials to avoid glass particles from the breaking of ampules.</td>
<td></td>
</tr>
<tr>
<td>Theophylline</td>
<td>0.4, 0.8, 1.6, 2, 3.2, 4 mg/mL.</td>
<td>—</td>
<td>Available in large volume solutions only.</td>
<td></td>
</tr>
</tbody>
</table>

(continued)
### THEOPHYLLINE PRODUCTS COMPARISON CHART

#### SLOW-RELEASE PRODUCTS

<table>
<thead>
<tr>
<th>PRODUCT</th>
<th>ANHYDROUS THEOPHYLLINE CONTENT</th>
<th>MEASURABLE DOSE INCREMENT (MG)</th>
<th>COMMENTS</th>
</tr>
</thead>
<tbody>
<tr>
<td>Slo-bid Gyrocaps</td>
<td>Cap 50 mg.</td>
<td>25</td>
<td>Excellent bioavailability in young infants; beads can be sprinkled on small amount of food; serum level fluctuations are 43%/18%.</td>
</tr>
<tr>
<td></td>
<td>Cap 75 mg.</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Cap 100 mg.</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Cap 125 mg.</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Cap 200 mg.</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Cap 300 mg.</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Theo-Dur</td>
<td>Tab 100 mg scored.</td>
<td>25</td>
<td>Serum level fluctuations are 38%/16% for 200, 300, and 450 mg, and 87%/34% for 100 mg tablets; some rapid metabolizers may require 8-hr dosage intervals to avoid breakthrough of symptoms.</td>
</tr>
<tr>
<td></td>
<td>Tab 200 mg scored.</td>
<td>100</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Tab 300 mg scored.</td>
<td>150</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Tab 450 mg scored.</td>
<td>225</td>
<td></td>
</tr>
<tr>
<td>Uni-Dur</td>
<td>Tab 400 mg scored.</td>
<td>200</td>
<td>The extent of absorption of Uni-Dur does not appear to be affected by food; however, large serum level fluctuations (78% in adults) may render this agent unreliable for once-daily administration.</td>
</tr>
<tr>
<td></td>
<td>Tab 600 mg scored.</td>
<td>300</td>
<td></td>
</tr>
</tbody>
</table>

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*Only products with documented bioavailability that are minimally affected by food and with dosage forms that permit incremental changes in dose are listed.*

*Accuracy of measurement decreases below 0.5 mL with suspensions and syrups because of viscosity; smaller amounts cannot be accurately measured; measure all liquid dosage forms with a syringe.*

*Predicted child/adult fluctuation between peak and trough (%) for 12-hr dosage interval: average child \( t_{1/2} = 3.7 \) hr, average adult \( t_{1/2} = 8.2 \) hr.*

*The ethylenediamine portion of aminophylline may cause urticaria or exfoliative dermatitis rarely.*

*Only Slo-bid Gyrocaps and Theo-Dur tablets have sufficiently slow and complete absorption to allow 12-hr dosage intervals with minimal serum concentration fluctuations in most patients. Many products advertised for bid dosage do not maintain serum concentrations within the therapeutic range in many patients, especially children. Some once-daily dosage products (eg, Uniphyl) are affected by food and may be unreliable.*
Pharmacology. Zileuton is an inhibitor of leukotriene synthesis. It has anti-inflammatory activity and inhibits the antigen-induced contraction of the trachea and bronchospasm that occurs in asthma.

**Adult Dosage.** PO for asthma prophylaxis 600 mg qid; dosage reduction may be necessary in hepatic dysfunction.

**Pediatric Dosage.** PO (<12 yr) safety and efficacy not established; (≥12 yr) same as adult dosage.

**Dosage Forms.** Tab 600 mg.

**Pharmacokinetics.** Zileuton is orally absorbed; food has no important effect on absorption. It is 93% plasma protein bound; V_d is about 1.2 L/kg. It is metabolized by CYP1A2, 2C9, and 3A4 and has a half-life of 2.5 hr.

**Adverse Reactions.** It is generally well tolerated, with headache reported in about 10% of patients in clinical trials. GI effects such as nausea and dyspepsia occur occasionally. Hepatic enzyme abnormalities have been reported. It is contraindicated in patients with active hepatic disease. Low WBC counts occur at rates greater than those in placebo-treated patients.

**Drug Interactions.** Zileuton markedly increases the effects of propranolol, theophylline, and warfarin.

**Parameters to Monitor.** Obtain ALT at baseline and monthly for 3 months, then q 2–3 months for the remainder of the first year, then periodically.

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**Antihistamines**

**Class Instructions.** Antihistamines. This drug (with the exceptions of fexofenadine and loratadine) can cause drowsiness, dry mouth, or occasional dizziness. Until the extent of drowsiness is known, use caution when driving, operating machinery, or performing other tasks requiring mental alertness or motor coordination. Avoid excessive concurrent use of alcohol and other central nervous system depressants that cause drowsiness. This drug effectively suppresses seasonal allergic rhinitis only when taken continuously.

**Missed Doses.** Missed doses should be taken as soon as possible. However, if it is almost time for the next dose, skip the missed dose and go back to the regular dosage schedule. Do not double doses.

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**CETIRIZINE**

**Pharmacology.** Cetirizine is a low-sedating, long-acting H_1-receptor antagonist that is a metabolite of hydroxyzine. Cetirizine competitively inhibits the interaction of histamine with H_1 receptors, thereby preventing the allergic response.

**Administration and Adult Dosage.** PO for allergic rhinitis or urticaria 5–10 mg/day depending on symptom severity.

**Special Populations.** Pediatric Dosage. PO for allergic rhinitis or urticaria (2–5 yr) 2.5–5 mg/day; (≥6 yr) same as adult dosage.
Geriatric Dosage. PO Same as adult dosage. Reducing dosage in geriatric patients might be necessary because of a 50% increase in cetirizine’s half-life and a 40% decrease in clearance.

Other Conditions. In patients with Clcr of 11–31 mL/min, those on hemodialysis, and in hepatically impaired patients, give 5 mg/day.

Dosage Forms. Tab 5, 10 mg; Syrup 1 mg/mL.

Patient Instructions. (See Class Instructions: Antihistamines.)

Pharmacokinetics. Onset and Duration. Onset is within 1 hr; duration is 24 hr.

Fate. Cetirizine is rapidly absorbed after oral administration. Peak serum levels are reached within 1 hr. Food does not affect the amount absorbed but might decrease the absorption rate. Protein binding averages 93%. CI in normal adults is 0.04–0.05 L/kg/hr. Cetirizine is oxidized to a small extent to inactive metabolites. After a 10 mg dose, 70% of the drug is excreted unchanged in the urine within 72 hr and 10% is excreted in feces. Cetirizine is not appreciably dialyzable.\( t_{1/2} \) (Adults) 7–10 hr; (children) 6–7 hr; (elderly/renal insufficiency) 18–21 hr.30–34

Adverse Reactions. The most frequent side effects are sedation, headache, dry mouth, fatigue, and nausea. Cetirizine 10 mg/day produces more sedation than loratadine 10 mg/day or placebo. Cetirizine has not been implicated in cardiac adverse events. Higher-than-recommended doses of cetirizine (up to 60 mg daily) did not prolong the QT interval in 25 healthy volunteers.

Contraindications. Hypersensitivity to hydroxyzine or cetirizine.

Precautions. Sedative effects may be dose dependent.

Drug Interactions. Exercise caution when cetirizine is combined with anticholinergic agents, alcohol, or other CNS depressants. Because most of cetirizine is eliminated renally, cytochrome P450 interactions are not likely. Clinically important drug interactions have not been found with cetirizine and azithromycin, pseudoephedrine, ketoconazole, or erythromycin. Clearance of cetirizine was reduced slightly by a 400 mg dose of theophylline, but this reduction was not clinically important; however, larger doses might have a greater effect. Therefore, it seems appropriate to monitor patients for increased sedation or other CNS-related side effects when administering theophylline concomitantly with cetirizine.

Parameters to Monitor. (Allergic rhinitis) observe for sneezing, rhinorrhea, itchy nose, and conjunctivitis. Monitor for side effects such as sedation.

Notes. Cetirizine and loratadine have the advantages of noncardiotoxicity, once-daily administration, and availability of liquid dosage forms. (See Antihistamines Comparison Chart.)

CHLORPHENIRAMINE MALEATE

Pharmacology. Chlorpheniramine is a competitive antagonist of histamine at the \( \text{H}_1 \)–histamine receptor. It also has anticholinergic and transient sedative effects when used intermittently.

Administration and Adult Dosage. PO for seasonal allergic rhinitis (effectiveness is maximized if given continuously, starting just before the pollen season)
4 mg hs initially, increasing gradually over 10 days as tolerated to 24 mg/day in 1–2 divided doses until the end of the season. **PO for acute allergic reactions** 12 mg in 1–2 divided doses. **SR** (see Notes.)

**Special Populations.** **Pediatric Dosage.** PO for **seasonal allergic rhinitis** (2–6 yr) 1 mg q 4–6 h up to 4 mg/day; (6–12 yr) 2 mg hs initially, increasing gradually over 10 days as tolerated to 12 mg/day in 1–2 divided doses until the end of the season. **SR** (see Notes.)

**Geriatric Dosage.** PO (≥60 yr) 4 mg daily–bid.

**Dosage Forms.** Chew **Tab** 2 mg; **Tab** 4 mg; **Syrup** 0.4 mg/mL; **SR Cap** 8, 12 mg (see Notes); **SR Tab** 8, 12 mg (see Notes); **Cap** 4, 10 mg with pseudoephedrine HCl 60 and 65 mg, respectively (various), and 8, 12 mg with pseudoephedrine HCl 120 mg (various).

**Patient Instructions.** (See Class Instructions: Antihistamines.)

**Pharmacokinetics. Onset and Duration.** Onset is 0.5–1 hr; duration of suppression of wheal and flare response (IgE mediated) to skin tests with allergenic extract is 2 days. Fast metabolizers have an earlier, greater, and more prolonged antihistaminic response than slow metabolizers because of rapid conversion to active metabolite. In the elderly, duration of action can be ≥36 hr, even when serum concentrations are low.

**Serum Levels.** Serum chlorpheniramine levels do not correlate with histamine antagonistic activity because of an unidentified active metabolite. (Children) 2.3–12 μg/L (6–31 nmol/L) suppress allergic rhinitis symptoms; (children) 4–10 μg/L (11–26 nmol/L) suppress histamine-induced wheal and flare.

**Fate.** Oral bioavailability is about 34%; 72% is plasma protein bound. Vd is (adults) 3.2 ± 0.3 L/kg; (children) 7 ± 2.8 L/kg; Cl is (adults) 0.1 ± 0.006 L/hr/kg; (children) 0.43 ± 0.19 L/hr/kg. Rapidly and extensively metabolized by CYP2D6 to mono- and didesmethylchlorpheniramine and unidentified metabolites, one or more of which are active. Metabolites and a small amount of parent drug are excreted in urine.

\[ t_{1/2} \]
- (Adults) 20 ± 5 hr; (children) 13 ± 6 hr; (chronic renal failure) 280–330 hr.

**Adverse Reactions.** Frequent drowsiness, dry mouth, dizziness, and irritability occur with intermittent therapy; however, most patients develop tolerance to these side effects during continuous therapy, particularly if the dosage is increased slowly.

**Contraindications.** Lactation; premature and newborn infants.

**Precautions.** Use chlorpheniramine with caution in patients ≥60 yr. It might cause paradoxical CNS stimulation in children. OTC labeling states to avoid in patients with narrow-angle glaucoma, symptomatic prostatic hypertrophy, asthma, emphysema, chronic pulmonary disease, shortness of breath, or breathing difficulties except under physician supervision; however, many studies have shown some bronchodilator effect of H1-receptor antagonists.
Drug Interactions. MAOIs prolong and intensify the anticholinergic effects of antihistamines.\textsuperscript{41} Alcohol or sedative-hypnotics can increase CNS depressant effects.

Parameters to Monitor. In seasonal allergic rhinitis, observe for sneezing, rhinorrhea, itchy nose, and conjunctivitis.

Notes. Not effective for nasal stuffiness. SR formulations offer no advantage over syrup or plain, uncoated tablets because the drug has an inherently long duration of action. (See Antihistamines Comparison Chart.)

DIPHENHYDRAMINE HYDROCHLORIDE Benadryl, Various

Pharmacology. (See Chlorpheniramine.) Diphenhydramine has strong sedating and anticholinergic properties.

Administration and Adult Dosage. PO as an antihistamine or for parkinsonism 25–50 mg tid–qid. PO for motion sickness 50 mg 30 min before exposure, then ac and hs. PO as a nighttime sleep aid 25–50 mg hs. PO as an antitussive 25 mg q 4–6 hr. Deep IM or IV as an antihistamine, or for allergic reactions to blood or plasma, motion sickness, adjunctive treatment of anaphylaxis, or parkinsonism 10–50 mg/dose, 100 mg if required, to a maximum of 400 mg/day.

Special Populations. Pediatric Dosage. PO as an antihistamine 5 mg/kg/day, or (≤9 kg) 6.25–12.5 mg tid–qid; (>9 kg) 12.5–25 mg tid–qid, to a maximum of 300 mg/day. PO as an antitussive (2–6 yr) 6.25 mg q 4 hr, to a maximum of 25 mg/day; (6–12 yr) 12.5 mg q 4 hr, to a maximum of 75 mg/day. Deep IM or IV 5 mg/kg/day in 4 divided doses, to a maximum of 300 mg/day.

Geriatric Dosage. PO as an antihistamine 25 mg bid–tid initially, then increase as needed.\textsuperscript{42} (See Notes.)

Other Conditions. In renal impairment, increase dosage interval as follows: (Cl\textsubscript{cr} 10–50 mL/min), increase to 6–12 hr; (Cl\textsubscript{cr} <10 mL/min), increase to 12–18 hr.\textsuperscript{42}

Dosage Forms. Cap 25, 50 mg; Chew Tab 12.5 mg; Elxr 2.5 mg/mL; Syrup 1.25, 2.5 mg/mL; Tab 25, 50 mg; Inj 50 mg/mL.

Patient Instructions. (See Class Instructions: Antihistamines.)

Pharmacokinetics. Onset and Duration. Onset is 15 min after single oral dose; duration of suppression of wheal and flare is up to 2 days.\textsuperscript{37,43} Duration of effect does not appear to be related to serum levels.

Serum Levels. (Antihistaminic effect) >25 \(\mu\)g/L (0.09 \(\mu\)mol/L); (sedation) 30–50 \(\mu\)g/L (0.1–0.17 \(\mu\)mol/L); (mental impairment) >60 \(\mu\)g/L (0.2 \(\mu\)mol/L).\textsuperscript{39,43}

Fate. As a result of first-pass metabolism, oral bioavailability is variable, 61 ± 25%.\textsuperscript{39,44} A single 50 mg oral dose in adults usually produces serum concentrations of 25–50 \(\mu\)g/L.\textsuperscript{43} About 85% is plasma protein bound and lower in Asians and those with cirrhosis.\textsuperscript{35,46} \(V_d\) is 17.4 ± 4.8 L/kg in adults and larger in Asians and those with cirrhosis. Cl is 1.4 ± 0.6 L/hr/kg in adults and higher in Asians and 0.7 ± 0.2 L/hr/kg in the elderly.\textsuperscript{39,42,44,45} Metabolized to N-dealkylated and acidic metabolites.\textsuperscript{44,47} Less than 4% is excreted unchanged in urine.\textsuperscript{48}
t_{1/2}. (Adults) 9.2 ± 2.5 hr; (elderly >65 yr) 13.5 ± 4.2 hr; (children 8–12 yr) 5.4 ± 1.8 hr, \textsuperscript{39,44} (cirrhosis) 15 hr.\textsuperscript{46}

Adverse Reactions. (See Chlorpheniramine.)

Contraindications. (See Chlorpheniramine.)

Precautions. (See Chlorpheniramine.)

Drug Interactions. (See Chlorpheniramine.)

Parameters to Monitor. (See Chlorpheniramine.)

Notes. Because of its low degree of efficacy for pruritus, weak suppression of IgE-mediated skin tests, and high sedative potential, diphenhydramine is not the antihistamine of choice for most conditions. In the elderly, diphenhydramine is discouraged as a nighttime sleep aid because of its high anticholinergic potential.

Dimenhydrinate (Dramamine), used for motion sickness, is the 8-chlorotheophyllinate salt of diphenhydramine; 100 mg dimenhydrinate is about equal to 50 mg diphenhydramine.

FEXOFENADINE

Pharmacology. Fexofenadine is a histamine H\textsubscript{1}-receptor antagonist that is a metabolite of terfenadine. It causes little sedation and has little anticholinergic activity. (See Antihistamines Comparison Chart.)

Adult Dosage. PO for allergic rhinitis 180 mg once daily or 60 mg bid; PO for chronic idiopathic urticaria 60 mg bid. In renal impairment, reduce initial dosage to 60 mg/day.

Pediatric Dosage. PO (<6 yr) safety and efficacy not established; (6–11 yr) 30 mg bid; (≥12 yr) same as adult dosage.

Dosage Forms. Cap 60 mg; Tab 30, 60, 180 mg; Tab 60 mg with pseudoephedrine 120 mg (Allegra-D).

Pharmacokinetics. Onset and Duration. Onset is rapid; peak serum levels occur at 2.6 hr. Food decreases oral absorption. It is 60–70\% plasma protein bound and excreted unchanged in urine and feces. Its half-life is about 14 hr in normal renal function and increases to about 19 hr in severe renal impairment.

Adverse Reactions. Drowsiness or fatigue occur in <2\% of patients; GI effects occur in about 1.5\% and headache in >1\%. Pharyngitis and menstrual disturbances have been reported.

HYDROXYZINE HYDROCHLORIDE

Pharmacology. Hydroxyzine is a competitive antagonist of histamine at the H\textsubscript{1}-histamine receptor. It also has antiemetic and sedative effects, thought to be a result of CNS subcortical suppression. Claims of long-term antianxiety properties have not been substantiated by well-designed studies.

Administration and Adult Dosage. PO for pruritus 25 mg tid–qid. PO for seasonal allergic rhinitis (effectiveness is maximized if given continuously just before the pollen season) 25 mg initially q hs until no sedation in the morning, then
increase dosage q 2–3 days, to a maximum of 150 mg/day in 1–2 divided doses and maintain until the end of the season. Reduce dosage by one-third or more if sedation persists. Dosage may be increased, if tolerated, for symptoms during the peak of pollen season.\textsuperscript{49} IM for sedation before and after general anesthesia 50–100 mg. IM for nausea and vomiting and pre- and postoperative adjunctive medication 25–100 mg. Preferred IM injection site is upper outer quadrant of gluteus maximus or midlateral thigh. Not for SC or intra-arterial use.

\textbf{Special Populations. Pediatric Dosage.} PO for pruritus (<6 yr) 50 mg/day in 2–3 divided doses; (≥6 yr) 50–100 mg/day in divided doses. PO for seasonal allergic rhinitis 10 mg initially q hs until no sedation in the morning, then increase dosage q 2–3 days, to a maximum of 75 mg/day in 1–2 divided doses and maintain until the end of the season. Reduce dosage by one-third or more if sedation persists. Dosage may be increased, if tolerated, for symptoms during the peak of pollen season.\textsuperscript{49} IM for pre- and postoperative sedation 0.7 mg/kg. IM for nausea and vomiting and pre- and postoperative adjunctive medication 1.1 mg/kg. Preferred site in children is midlateral muscles of the thigh.

\textbf{Geriatric Dosage.} PO for pruritus 10 mg tid–qid, increasing to 25 mg tid–qid if necessary.\textsuperscript{51}

\textbf{Dosage Forms.} Cap (as pamoate equivalent of HCl salt) 25, 50, 100 mg; Susp (as pamoate equivalent of HCl salt) 5 mg/mL; Syrup (as HCl) 2 mg/mL; Tab (as HCl) 10, 25, 50, 100 mg; Inj (as HCl) 25, 50 mg/mL (IM only).

\textbf{Patient Instructions.} (See Class Instructions: Antihistamines.)

\textbf{Pharmacokinetics. Onset and Duration.} Onset 15–30 min after oral administration. Duration of suppression of wheal and flare response to allergenic extract skin test is 4 days.\textsuperscript{37,50}

\textbf{Serum Levels.} (Pruritus) 6–42 μg/L (14–102 nmol/L) suppress pruritus in children.\textsuperscript{52}

\textbf{Fate.} Peak serum level of 73 ± 11 μg/L occurs 2 ± 0.4 hr after a 0.7 mg/kg dose in healthy adults, 117 ± 61 μg/L at 2.3 ± 0.7 hr in primary biliary cirrhosis (mean dose 44 mg).\textsuperscript{53,54} \(V_d\) is (healthy adults) 16 ± 3 L/kg,\textsuperscript{53} (elderly) 23 ± 6 L/kg, (children) 19 ± 9 L/kg,\textsuperscript{52} and (primary biliary cirrhosis) 23 ± 13 L/kg.\textsuperscript{54} CI is (healthy young and elderly adults) 0.6 ± 0.2 L/hr/kg,\textsuperscript{51,53} (children) 1.9 L/hr/kg, (primary biliary cirrhosis) 0.5 ± 0.4 L/hr/kg.\textsuperscript{54}

\(t_{1/2}\). (Healthy adults) 20 ± 4 hr;\textsuperscript{53} (elderly) 29 ± 10 hr;\textsuperscript{51} (children) 7 hr, increasing with age;\textsuperscript{53} (primary biliary cirrhosis) 37 ± 13 hr.\textsuperscript{54}

\textbf{Adverse Reactions.} Transient drowsiness and dry mouth occur frequently when the drug is taken intermittently. Most patients develop tolerance to these effects when the drug is taken continuously, particularly if the dosage is slowly increased over 7–10 days. IM injection can be painful and has caused sterile abscess. Hemolysis has been associated with IV administration and tissue necrosis with SC or intra-arterial administration.

\textbf{Contraindications.} Early pregnancy; SC or intra-arterial use of injectable solution.

\textbf{Precautions.} Use with caution in the elderly.
**Drug Interactions.** MAOIs prolong and intensify the anticholinergic effects of antihistamines. Alcohol or sedative-hypnotics can increase CNS depressant effects.

**Parameters to Monitor.** In seasonal allergic rhinitis, observe for sneezing, rhinorrhea, itchy nose, and conjunctivitis.

**Notes.** Hydroxyzine suppresses wheal and flare response to the greatest degree and for the longest duration of all antihistamines,\(^\text{37}\) including the newer nonsedating antihistamines.\(^\text{50}\)

**LORATADINE**

**Pharmacology.** Loratadine is a long-acting piperadine antihistamine that is structurally similar to azatadine, with little or no action at \(\alpha\)-adrenergic or cholinergic receptors.\(^\text{55}\) (See Antihistamines Comparison Chart.)

**Adult Dosage.** PO for allergic rhinitis or urticaria 10 mg/day on an empty stomach. In patients with hepatic impairment, begin with 10 mg every other day. The dosage of Claritin-D is 1 tablet bid on an empty stomach; Claritin-D 24-hr is given once daily.

**Pediatric Dosage.** PO (<2 yr) safety and efficacy not established; (2–6 yr) 5 mg/day; (>6 yr) same as adult dosage.

**Dosage Forms.** Tab 10 mg (conventional and rapidly dissolving); Syrup 1 mg/mL; Tab 5 mg with pseudoephedrine 120 mg (Claritin-D); Tab 10 mg with pseudoephedrine 240 mg (Claritin-D 24-hr).

**Pharmacokinetics.** The drug is rapidly absorbed; bioavailability and peak serum levels are increased by about 50% in the elderly (66–78 yr) or when taken with food. It is 97% bound to plasma proteins and extensively metabolized to an active metabolite, descarboethoxyloratadine. Approximately 80% of a dose is excreted equally in urine and feces as metabolites after 10 days. The half-lives in healthy adults are 8.4 hr (range 3–20) for loratadine and 28 hr (range 8.8–92) for descarboethoxyloratadine.\(^\text{55,56}\)

**Adverse Reactions.** Headache and mild, dose-related drowsiness or fatigue occur occasionally.

**Notes.** Desloratadine (Clarinex) is the active metabolite of loratadine given in a dose of 5 mg once daily. It might have some advantages such as fewer adverse reactions and drug interactions.
<table>
<thead>
<tr>
<th>DRUG</th>
<th>DOSAGE FORMS</th>
<th>ADULT DOSAGE</th>
<th>PEDIATRIC DOSAGE</th>
<th>HALF-LIFE (HR)</th>
<th>SIDE EFFECTS</th>
<th>Sedation</th>
<th>Anticholinergic</th>
</tr>
</thead>
<tbody>
<tr>
<td>Acrivastine</td>
<td>Tab 8 mg with pseudoephedrine 60 mg (Semprex-D).</td>
<td>PO 1 tab q 4–6 hr, to a maximum of 4/day.</td>
<td>PO (&gt;12 yr) same as adult dosage.</td>
<td>1.5–3</td>
<td>+</td>
<td>±</td>
<td></td>
</tr>
<tr>
<td>Azatadine Maleate Optimine</td>
<td>Tab 1 mg.</td>
<td>PO 1–2 mg bid.</td>
<td>(&lt;12 yr) safety and efficacy not established.</td>
<td>12</td>
<td>++</td>
<td>++</td>
<td></td>
</tr>
<tr>
<td>Azelastine HCl Astelin</td>
<td>Nasal spray 125 µg/spray.</td>
<td>2 sprays/nostril bid.</td>
<td>(5–11 yr) 1 spray/nostril bid; (≥12 yr) same as adult dosage.</td>
<td>22 (metabolite: 54)</td>
<td>++</td>
<td>++</td>
<td></td>
</tr>
<tr>
<td>Brompheniramine Maleate Dimetapp Allergy Various</td>
<td>Cap 4 mg Inj 10 mg/mL.</td>
<td>PO 4 mg q 4–6 hr, to a maximum of 24 mg/day SC, IM, or slow IV 5–20 mg q 12 hr, to a maximum of 40 mg/day.</td>
<td>PO (&lt;12 yr) 0.5 mg/kg/day in 3–4 doses SC, IM, or slow IV 0.5 mg/kg/day in 3–4 divided doses.</td>
<td>25</td>
<td>+</td>
<td>++</td>
<td></td>
</tr>
</tbody>
</table>

(continued)
## ANTIHISTAMINES COMPARISON CHART (continued)

<table>
<thead>
<tr>
<th>DRUG</th>
<th>DOSAGE FORMS</th>
<th>ADULT DOSAGE</th>
<th>PEDIATRIC DOSAGE</th>
<th>HALF-LIFE (HR)</th>
<th>SIDE EFFECTS</th>
</tr>
</thead>
</table>
| Carboxinomine Maleate | Drp 2 mg with pseudoephedrine 25 mg/mL.  
Solv 0.4 mg/mL.  
Syrup 0.4, 0.8 mg with pseudoephedrine 6, 12 mg/mL, respectively  
Tab 4 mg with pseudoephedrine 60 mg Carbodec, Rondec.  
SR Tab 8 mg with pseudoephedrine 90 mg (Palgic-D). | PO 1 tab qid.          | PO 0.2–0.4 mg/kg/day;  
(1–3 yr) 2 mg tid–qid;  
(3–6 yr) 2–4 mg tid–qid;  
(>6 yr) 4–6 mg tid–qid  
(dosage refers to carboxinomine component). | 10–20           | ++                    | +++                   |
| Cetirizine HCl        | Tab 5, 10 mg  
Syrup 1 mg/mL. | PO 5–10 mg/day.         | PO (2–5 yr) 2.5–5 mg/day;  
(6–11 yr) 5–10 mg/day. | 7–10          | +                     | ±                     |
| Chlorpheniramine Maleate | Syrup 0.4 mg/mL.  
Chew Tab 2 mg  
Tabletab 4 mg  
SR Tab 8, 12 mg  
Inj 10, 100 mg/mL. | PO (acute allergic reactions) 12 mg/day  
in 1–2 divided doses;  
PO (seasonal allergic rhinitis) 24 mg/day  
in 1–2 divided doses;  
IV (acute allergic reactions) 5–40 mg. | PO (seasonal allergic rhinitis) (2–6 yr)  
1 mg tid up to 4 mg/day,  
SR not recommended;  
(6–12 yr) 2 mg tid up to 12 mg/day;  
SR not recommended. | 15–25         | +                     | ++                    |
## Antihistamines Comparison Chart (continued)

<table>
<thead>
<tr>
<th>Drug</th>
<th>Dosage Forms</th>
<th>Adult Dosage</th>
<th>Pediatric Dosage</th>
<th>Half-Life (HR)</th>
<th>Side Effects</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Clemastine</strong></td>
<td>Syrup 0.13 mg (equivalent to 0.1 mg clemastine)/mL Tab 1.34, 2.68 mg (equivalent to 1 and 2 mg clemastine, respectively).</td>
<td>PO 1.34 mg bid–2.68 mg tid, to a maximum of 8.04 mg/day.</td>
<td>PO (6–12 yr) 0.67–1.34 mg bid, to a maximum of 4.02 mg/day.</td>
<td>21</td>
<td>++</td>
</tr>
<tr>
<td><strong>Fumarate</strong></td>
<td><strong>Tavist</strong></td>
<td></td>
<td></td>
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<tr>
<td><strong>Various</strong></td>
<td></td>
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</tr>
<tr>
<td><strong>Cyproheptadine HCl</strong></td>
<td>Tab 4 mg Syrup 0.4 mg/mL.</td>
<td>PO 4–20 mg/day, usually 4 mg tid–qid, to a maximum of 0.5 mg/kg/day.</td>
<td>PO (2–6 yr) 2 mg bid–tid, to a maximum of 12 mg/day; (7–14 yr) 4 mg bid–tid, to a maximum of 16 mg/day.</td>
<td>—</td>
<td>+</td>
</tr>
<tr>
<td><strong>Periactin</strong></td>
<td></td>
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<td></td>
</tr>
<tr>
<td><strong>Various</strong></td>
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<tr>
<td><strong>Desloratadine</strong></td>
<td>Tab 5 mg.</td>
<td>PO 5 mg/day.</td>
<td></td>
<td></td>
<td>28</td>
</tr>
<tr>
<td><strong>Clarinex</strong></td>
<td></td>
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</tr>
<tr>
<td><strong>Dexchlorpheniramine Maleate</strong></td>
<td>Syrup 0.4 mg/mL. Tab 2 mg SR Tab 4, 6 mg.</td>
<td>PO 2 mg q 4–6 hr, to a maximum of 12 mg/day or SR 4–6 mg hs or q 6–10 hr during the day.</td>
<td>PO (2–5 yr) 0.5 mg q 4–6 hr, to a maximum of 3 mg/day, SR not recommended; (6–12 yr) 1 mg q 4–6 hr or SR 4 mg hs.</td>
<td>15–25</td>
<td>+</td>
</tr>
</tbody>
</table>
## Antihistamines Comparison Chart (continued)

<table>
<thead>
<tr>
<th>Drug</th>
<th>Dosage Forms</th>
<th>Adult Dosage</th>
<th>Pediatric Dosage</th>
<th>Half-Life (HR)</th>
<th>Side Effects</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Diphenhydramine HCl</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Benadryl</td>
<td>Cap 25, 50 mg</td>
<td>PO (antihistamine)</td>
<td>PO (&gt;9 kg) 5 mg/kg/day,</td>
<td>9</td>
<td>+++</td>
</tr>
<tr>
<td></td>
<td>Chew Tab 12.5 mg</td>
<td>25–50 mg tid–qid</td>
<td>usually 12.5–25 mg</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Elixir 2.5 mg/mL</td>
<td>PO (motion sickness)</td>
<td>tid–qid, to a maximum</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Syrup 1.25, 2.5 mg/mL</td>
<td>50 mg 30 min before</td>
<td>of 300 mg/day</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Tab 25, 50 mg</td>
<td>exposure, ac and hs</td>
<td>PO (antitussive) (6–12 yr)</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Inj 50 mg/mL</td>
<td>25 mg q 4–6 hr</td>
<td>12.5 mg q 4 hr, to a</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>PO (nighttime sleep aid)</td>
<td>maximum of 75 mg/day</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>25–50 mg hs</td>
<td>IM, IV 5 mg/kg, to a</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>IM, IV 10–50 mg, to a maximum of 400 mg/day.</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Ebastine</strong></td>
<td></td>
<td>PO 10–20 mg/day.</td>
<td>—</td>
<td>15 (metabolite)</td>
<td>±</td>
</tr>
<tr>
<td>Kestine (Investigational—RPR)</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>±</td>
</tr>
<tr>
<td><strong>Fexofenadine HCl</strong></td>
<td>Cap 60 mg</td>
<td>PO 60 mg bid or</td>
<td>PO (&gt;12 yr) same as</td>
<td>14</td>
<td>±</td>
</tr>
<tr>
<td>Allegra</td>
<td>Tab 30, 60, 180 mg</td>
<td>180 mg once daily.</td>
<td>adult dosage.</td>
<td></td>
<td>±</td>
</tr>
</tbody>
</table>

(continued)
<table>
<thead>
<tr>
<th>DRUG</th>
<th>DOSAGE FORMS</th>
<th>ADULT DOSAGE</th>
<th>PEDIATRIC DOSAGE</th>
<th>HALF-LIFE (HR)</th>
<th>SIDE EFFECTS</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Hydroxyzine HCl/Pamoate</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Atarax</td>
<td>Cap (as pamoate equivalent of HCl salt) 25, 50, 100 mg</td>
<td>PO (pruritus) 25 mg tid-qid; (seasonal allergic rhinitis) titrate up to 150 mg/day in 1–2 divided doses</td>
<td>IM 25–100 mg.</td>
<td>16–24</td>
<td>++</td>
</tr>
<tr>
<td>Vistaril</td>
<td>Susp (as pamoate equivalent of HCl salt) 5 mg/mL</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Syrup (as HCl) 2 mg/mL</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Tab (as HCl) 10, 25, 50, 100 mg</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Inj (as HCl) 25, 50 mg/mL.</td>
<td></td>
<td></td>
<td></td>
<td></td>
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<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Levocabastine HCl</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Livostin</td>
<td>Ophth Susp 0.05%</td>
<td></td>
<td></td>
<td>35–40</td>
<td>+</td>
</tr>
<tr>
<td></td>
<td>Nasal Spray (Investigational)</td>
<td></td>
<td></td>
<td></td>
<td>+</td>
</tr>
<tr>
<td><strong>Loratadine</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Claritin</td>
<td>Tab 10 mg</td>
<td>PO 10 mg/day.</td>
<td>PO (2–6 yr) 5 mg/day; (≥6 yr) 10 mg/day</td>
<td>8</td>
<td>±</td>
</tr>
<tr>
<td></td>
<td>Syrup 1 mg/mL</td>
<td></td>
<td>(metabolite: 28)</td>
<td></td>
<td>±</td>
</tr>
<tr>
<td><strong>Phenindamine</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Tartrate</td>
<td>Tab 25 mg.</td>
<td>PO 25 mg q 4–6 hr, to a maximum of 150 mg/day.</td>
<td>PO (6–11 yr) 12.5 mg q 4–6 hr, to a maximum of 75 mg/day.</td>
<td>—</td>
<td>+</td>
</tr>
<tr>
<td>Nolahist</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>++</td>
</tr>
</tbody>
</table>

(continued)
<table>
<thead>
<tr>
<th>DRUG</th>
<th>DOSAGE FORMS</th>
<th>ADULT DOSAGE</th>
<th>PEDIATRIC DOSAGE</th>
<th>HALF-LIFE (HR)</th>
<th>SIDE EFFECTS</th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>Promethazine HCl</td>
<td>Syrup 1.25, 5 mg/mL.</td>
<td>PO (allergy) 25 mg hs or 12.5 mg ac and hs; (nausea and vomiting) 25 mg initial dose, then 12.5–25 mg q 4–6 hr pm; (adjunctive preoperative use) 25–50 mg/dose IM, IV (IV maximum concentration 25 mg/mL, maximum rate 25 mg/min) or PR (allergy) 25 mg, may repeat in 2 hr; (nausea and vomiting) 12.5–25 mg q 4 hr pm; (adjunctive pre- and postoperative use) 25–50 mg/dose.</td>
<td>PO (allergy) 6.25–12.5 mg qid; (motion sickness or sedation) 12.5–25 mg/bid; IM, IV, or PR (PR not recommended &lt;2 yr); (allergy) 0.5 mg/kg/day in 4 divided doses; (adjunctive preoperative use) 1 mg/kg/dose, maximum dosage not to exceed one-half of adult dosage.</td>
<td>12</td>
<td>++++</td>
<td>++++</td>
</tr>
<tr>
<td>Phenergan</td>
<td>Tab 12.5, 25, 50 mg</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Various</td>
<td>Inj 25, 50 mg/mL.</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Supp 12.5, 25, 50 mg.</td>
<td></td>
<td></td>
<td></td>
<td></td>
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</tr>
</tbody>
</table>

(continued)
### ANTIHISTAMINES COMPARISON CHART (continued)

<table>
<thead>
<tr>
<th>DRUG</th>
<th>DOSAGE FORMS</th>
<th>ADULT DOSAGE</th>
<th>PEDIATRIC DOSAGE</th>
<th>HALF-LIFE (HR)</th>
<th>SIDE EFFECTS</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Tripelennamine HCl</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>PBZ</td>
<td>Tab 25, 50 mg</td>
<td>PO 25–50 mg q 4–6 hr, to a maximum of 600 mg/day.</td>
<td>PO 5 mg/kg/day in 4–6 divided doses, to a maximum of 300 mg/day; SR not recommended.</td>
<td>2–4</td>
<td>++</td>
</tr>
<tr>
<td>Various</td>
<td>SR Tab 100 mg.</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Triprolidine HCl</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Syrup 0.25 mg with pseudoephedrine 6 mg/mL</td>
<td>PO 2.5 mg q 4–6 hr, to a maximum of 10 mg/day.</td>
<td>PO (4 months–2 yr) 0.3 mg tid–qid; (2–5 yr) 0.625 mg tid–qid; (6–12 yr) 1.25 mg q 4–6 hr, to a maximum of 4 doses/day.</td>
<td>3</td>
<td>++</td>
</tr>
<tr>
<td></td>
<td>Tab 2.5 mg with pseudoephedrine 60 mg (Actifed)</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

++++ = very high; +++ = high; ++ = moderate; + = low; ± = low to none; — = not known

*Tolerance usually develops during long-term therapy.

From references 55, and 56–60 and product information.
Corticosteroids

**BECLOMETHASONE DIPROPIONATE**  Beconase, Beclovent, QVAR, Vancenase, Vanceril

**Pharmacology.** Potent topical glucocorticoid with little systemic activity because of low systemic bioavailability.

**Administration and Adult Dosage.** **Inhal for asthma** (Beclovent, Vanceril) 168-840 μg bid; (QVAR) 80–320 μg bid. (See Notes.) **Intranasal for nasal congestion** 42–84 μg/nostril bid–qid (168–336 μg/day total dosage) for several days, then decrease dosage (if symptoms do not recur) to minimum amount necessary to control stuffiness.

**Special Populations.** **Pediatric Dosage.** Titrate dosage to the lowest effective dosage. **Inhal for asthma** (Beclovent, Vanceril) (6–12 yr) 42–336 μg bid; (>12 yr) same as adult dosage. **Intranasal for nasal congestion** (<6 yr) not recommended; (6–12 yr) 42 μg/nostril bid or tid.

**Geriatric Dosage.** Same as adult dosage.

**Other Conditions.** During a severe asthma attack, patients require supplementary treatment with systemic steroids.

**Dosage Forms.** **Inhal** (Beclovent, Vanceril) 42, 84 μg/puff (80 and 200 doses/inhaler, and 40 and 120 doses/inhaler, respectively); (QVAR) 40, 80 μg/puff (see Notes); **Nasal Inhal** (Beconase, Vancenase) 42 μg/spray (80, 200 doses/inhaler); **Aq Susp** (Beconase AQ, Vancenase AQ) 42, 84 μg/spray (200 and 120 doses/bottle, respectively).

**Patient Instructions.** **Metered-dose Oral Inhaler.** (Aerosols) Remove inhaler cap and hold inhaler upright. Shake inhaler. Tilt your head back and breathe out slowly. To position the inhaler, open your mouth with the inhaler 1–2 inches away or in your mouth. (For young children and corticosteroid inhalers, use a spacer or holding chamber.) Press down on the inhaler to release medication as you start to breathe slowly. Breathe slowly for 3 to 5 seconds. Hold your breath for 10 seconds to allow the medication to reach deep into the lungs. Repeat as directed. (Dry Powder) close your mouth tightly around the mouthpiece and inhale rapidly. Hold the device horizontally (parallel to the ground) after it has been activated. Do not exhale into the device. Rinsing your mouth and gargling with water or mouthwash after administration may be beneficial. This medication is for preventive therapy and should not be used to treat acute asthma attacks. **Nasal Inhaler.** Blow your nose before use. Shake the container well. Remove the protective cap and hold the inhaler between your thumb and forefinger. Tilt your head back slightly and insert the end of the inhaler into one nostril. While holding the other nostril closed with one finger, press down once to release 1 dose and, at the same time, inhale gently. Hold your breath for a few seconds and then breathe out slowly through your mouth. Repeat the process in the other nostril. Avoid blowing your nose for the next 15 minutes.
**Missed Doses.** Take the missed dose as soon as possible. If it is almost time for the next dose, skip the missed dose and go back to regular dosage schedule. Do not double doses.

**Pharmacokinetics.**

**Onset and Duration.** Effect is usually evident within a few days but might take 2–4 weeks for maximum improvement.\(^6^2\)

**Fate.** Only \(\leq 10\%\) of an inhaled dose is deposited in the lung; 80% is deposited in the mouth and swallowed. Oral absorption is slow and incomplete (61–90%), and the drug undergoes extensive first-pass metabolism, resulting in oral bioavailability of less than 5\%.\(^6^3\) Well absorbed from the lung and extensively metabolized, with 65\% excreted in the bile and \(< 10\%\) of unchanged drug and metabolites excreted in urine.\(^6^3\)

\(t_{1/2}\) 15 hr.

**Adverse Reactions.** After oral use, localized growth of *Candida* in the mouth occurs frequently, but clinically apparent infections occur only occasionally. Hoarseness and dry mouth occur occasionally; minimal to no suppression of the pituitary–adrenal axis occurs at the recommended dosage; however, dose-dependent suppression occurs at higher dosages.\(^6^2,^6^4–^6^7\) After intranasal use, irritation and burning of the nasal mucosa and sneezing occur occasionally; intranasal and pharyngeal *Candida* infections, nasal ulceration, and epistaxis occur rarely. Cases of growth suppression unrelated to suppression of the pituitary–adrenal axis have been reported after use of intranasally or orally inhaled corticosteroids in children. With oral inhalation, the mean reduction in growth velocity is 1 cm/yr (range 0.3–1.8 cm/yr). The long-term implications for ultimate adult height are unknown.

**Contraindications.** Status asthmaticus or other acute episodes of asthma in which intensive measures are required; beclomethasone-exacerbated symptoms.

**Precautions.** During stress or severe asthmatic attacks, patients withdrawn from systemic corticosteroid should contact their physician immediately. Use the lowest effective dosage possible in children. The potential growth effects of inhaled corticosteroids in children should be weighed against the clinical benefits of the corticosteroids and the availability of nonsteroid alternatives.

**Drug Interactions.** None known.

**Parameters to Monitor.** For treatment of asthma, frequency of daytime asthmatic symptoms, and nocturnal use of prn sympathomimetic inhaler. For nasal congestion, relief of symptoms. Routinely monitor the growth of children receiving inhaled corticosteroids (eg, via stadiometry).

**Notes.** Patients needing long-term use of an orally inhaled corticosteroid should be continued on therapeutic doses of a bronchodilator. Before use, a patient should be as free of symptoms as possible, which can be achieved with a 1-week course of oral prednisone. The nasal inhalation provides effective, prompt relief of nasal congestion when the maximally tolerated dosage of oral sympathomimetics is inadequate. (See also Inhaled Corticosteroids Comparison Chart.)
<table>
<thead>
<tr>
<th>DRUG</th>
<th>DOSAGE FORMSb</th>
<th>DAILY DOSAGEc (Step 2)</th>
<th>DAILY DOSAGEc (Step 3)</th>
<th>DAILY DOSAGEc (Step 4)</th>
<th>RECEPTOR BINDING HALF-LIFE</th>
<th>TOPICAL POTENCYE</th>
<th>ORAL BIOAVAILABILITYd</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>SINGLE-INGREDIENT PRODUCTS</strong></td>
<td></td>
<td>Low</td>
<td>Medium</td>
<td>High</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Beclomethasone</td>
<td>MDI: 42, 84 µg/puff.</td>
<td>Adult: 168–504 µg</td>
<td>504–840 µg</td>
<td>&gt;840 µg</td>
<td>7.5 hr</td>
<td>600</td>
<td>20%</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Child: 84–336 µg</td>
<td>336–672 µg</td>
<td>&gt;672 µg</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Dipropionate HFA</td>
<td>MDI: 40, 80 µg/puff.</td>
<td>Adult: 80–160 µg</td>
<td>160–320 mg</td>
<td>&gt;320 µg</td>
<td>7.5 hr</td>
<td>600</td>
<td>20%</td>
</tr>
<tr>
<td>Vanceril</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Budesonide</td>
<td>DPI: 200 µg/inhal Neb Susp: 125, 250 µg/mL.</td>
<td>Adult: 200–400 µg</td>
<td>400–600 µg</td>
<td>&gt;600 µg</td>
<td>5.1 hr</td>
<td>980</td>
<td>11%</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Child: 100–200 µg</td>
<td>200–400 µg</td>
<td>&gt;400 µg</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Flunisolide</td>
<td>MDI: 250 µg/puff.</td>
<td>Adult: 500–1000 µg</td>
<td>1000–2000 µg</td>
<td>&gt;2000 µg</td>
<td>3.5 hr</td>
<td>330</td>
<td>21%</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Child: 500–750 µg</td>
<td>750–1250 µg</td>
<td>&gt;1250 µg</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>AeroBid</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
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<td></td>
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<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Fluticasone</td>
<td>MDI: 44, 110, 220 µg/puff. DPI: 50, 100, 250 µg/inhal.</td>
<td>Adult: 88–264 µg</td>
<td>264–660 µg</td>
<td>&gt;660 µg</td>
<td>10.5 hr</td>
<td>1200</td>
<td>1%</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Child: 88–176 µg</td>
<td>176–440 µg</td>
<td>&gt;440 µg</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

(continued)
<table>
<thead>
<tr>
<th>DRUG</th>
<th>DOSAGE FORMS</th>
<th>DAILY DOSAGE</th>
<th>RECEPTOR BINDING</th>
<th>TOPICAL POTENCY</th>
<th>ORAL BIOAVAILABILITY</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>Low (Step 2)</td>
<td>Medium (Step 3)</td>
<td>High (Step 4)</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>Adult: 400–1000 µg</td>
<td>1000–2000 µg</td>
<td>&gt;2000 µg</td>
<td>3.9 hr</td>
</tr>
<tr>
<td>Triamcinolone</td>
<td>MDI: Adult: 1000–2000 µg</td>
<td>3.9 hr</td>
<td>330</td>
<td>11%</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Child: 400–800 µg</td>
<td>800–1200 µg</td>
<td>&gt;1200 µg</td>
<td>3.9 hr</td>
<td>330</td>
</tr>
<tr>
<td>Acetonide</td>
<td>100 µg/puff.</td>
<td>800–1200 µg</td>
<td>&gt;1200 µg</td>
<td>3.9 hr</td>
<td>330</td>
</tr>
<tr>
<td>Azmacort</td>
<td></td>
<td>1000–2000 µg</td>
<td>&gt;2000 µg</td>
<td>3.9 hr</td>
<td>330</td>
</tr>
<tr>
<td>COMBINATION PRODUCTS</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Fluticasone</td>
<td>DPI: Fluticasone 100 µg, salmeterol 50 µg/inhal; Fluticasone 250 µg, salmeterol 50 µg/inhal; Fluticasone 500 µg, salmeterol 50 µg/inhal.</td>
<td>Adult: 100–50 bid</td>
<td>250–50 bid</td>
<td>500–50 bid</td>
<td>—</td>
</tr>
<tr>
<td>Propionate and Salmeterol</td>
<td>Advair Diskus</td>
<td>Fluticasone 100 µg, salmeterol 50 µg/inhal; Fluticasone 250 µg, salmeterol 50 µg/inhal; Fluticasone 500 µg, salmeterol 50 µg/inhal.</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

DPI = dry powder inhaler; MDI = metered-dose inhaler. Neb = nebulizer.

*Dosage ranges correspond to recommended treatment intensities for steps 2–4 of the NIH guidelines for diagnosis and management of asthma: step 1 = mild intermittent; step 2 = mild persistent; step 3 = moderate persistent; step 4 = severe persistent. The most important determinant of appropriate dosage is the clinician’s judgment of the patient’s response to therapy; the clinician must monitor the patient’s response on several clinical parameters and adjust the dosage accordingly. The stepwise approach to therapy emphasizes that once control of symptoms is achieved, the dosage of medication should be carefully titrated to the minimum dosage required to maintain control, thereby reducing the potential for adverse effects.

MDI dosages are expressed as the actuator dose (the amount of drug leaving the actuator and delivered to the patient), which is the labeling required in the United States. This is different from the dosage expressed as the valve dose (the amount of drug leaving the valve, not all of which is available to the patient), which is used in many European countries and in some of the scientific literature. DPI doses are expressed as the amount of drug in the inhaler following activation.

Potency determined from skin blanching; dexamethasone is the reference drug and has a value of 1 in this assay.

Bioavailability of the swallowed portion of the dose received by the patient. About 80% of the dose from an MDI without a spacer is swallowed. Nearly all of the drug delivered to the lungs is bioavailable. From 10–30% of an MDI dose is delivered to the lungs, depending on the product and device. Both the relative potency and the total bioavailability (inhaled + swallowed) determine the systemic activity of the product.

From references 20 and 68–70.
<table>
<thead>
<tr>
<th>DRUG</th>
<th>DOSAGE FORMS</th>
<th>ADULT DOSAGE</th>
<th>PEDIATRIC DOSAGE(^a)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Beclomethasone</td>
<td>Aerosol, Metered-Dose 42</td>
<td>1–2 sprays into each nostril bid–qid.</td>
<td>1 spray into each nostril bid–tid.</td>
</tr>
<tr>
<td>Dipropionate</td>
<td>µg/spray Beconase, Aqueous 42, 84 µg/spray.</td>
<td>2 sprays into each nostril bid or 4 sprays into each nostril q AM, to a maximum of 800 µg/day.</td>
<td>2 sprays into each nostril bid or 4 sprays into each nostril q AM, to a maximum of 400 µg/day.</td>
</tr>
<tr>
<td>Budesonide</td>
<td>Aerosol, Metered-Dose 32 µg/spray Rhinocort</td>
<td>2 sprays into each nostril bid or 4 sprays into each nostril q AM, to a maximum of 800 µg/day.</td>
<td>2 sprays into each nostril bid or 4 sprays into each nostril q AM, to a maximum of 400 µg/day.</td>
</tr>
<tr>
<td>Flunisolide</td>
<td>Spray, Aqueous 25 µg/spray Nasalide Nasarel Fluticasone Propionate</td>
<td>2 sprays into each nostril bid, to a maximum of 8 sprays/day into each nostril.</td>
<td>1 spray into each nostril tid–qid.</td>
</tr>
<tr>
<td>Fluticasone</td>
<td>Spray, Aqueous 50 µg/spray</td>
<td>2 sprays into each nostril daily or 1 spray into each nostril bid; maintenance 1 spray into each nostril daily, to a maximum of 200 µg/day.</td>
<td>(≥4 yr) 1 spray in each nostril daily (100 µg/day); for nonresponders, 2 sprays in each nostril daily or 1 spray in each nostril bid, decrease to 100 µg/day once a response is achieved.</td>
</tr>
<tr>
<td>Mometasone</td>
<td>Spray, Aqueous 50 µg/spray</td>
<td>2 sprays into each nostril once daily.</td>
<td>(&lt;12 yr) not established.</td>
</tr>
<tr>
<td>Furoate</td>
<td>Spray, Aqueous 55 µg/spray Nasonex Triamcinolone Acetonide Nasacort Nasarel Fluticasone Propionate</td>
<td>2 sprays into each nostril daily; adjust to a maximum of 4 sprays/day in 1–4 divided doses; maintenance as low as 1 spray/day.</td>
<td>Same as adult dosage.</td>
</tr>
</tbody>
</table>

\(^a\)Unless otherwise stated, pediatric dosage is for patients 6–12 yr; dosages for patients <6 yr have generally not been established. From references 68–70.
Cough and Cold

DEXTROMETHORPHAN HYDROBROMIDE Various

Pharmacology. Dextromethorphan is the nonanalgesic, nonaddictive D-isomer of the codeine analogue of levorphanol. With usual antitussive doses, the cough threshold is elevated centrally with little effect on the respiratory, cardiovascular, or GI systems.

Administration and Adult Dosage. PO as cough suppressant 10–30 mg q 4–8 hr, to a maximum of 120 mg/day; SR 60 mg q 12 hr.

Special Populations. Pediatric Dosage. PO as cough suppressant (<2 yr) not recommended; (2–6 yr) 2.5–7.5 mg q 4–8 hr, to a maximum of 30 mg/day (as syrup); (6–12 yr) 5–10 mg q 4 hr or 15 mg q 6–8 hr, to a maximum of 60 mg/day; (>12 yr) same as adult dosage. SR (2–5 yr) 15 mg q 12 hr; (6–12 yr) 30 mg q 12 hr. (See Notes.)

Geriatric Dosage. Same as adult dosage.

Dosage Forms. Cap 30 mg; Lozenge 2.5, 5, 7.5, 15 mg; Syrup 0.66, 0.7, 1, 1.5, 2, 3 mg/mL; SR Susp 6 mg/mL; (available in many combination products in different concentrations).

Patient Instructions. Do not use this drug to suppress productive cough or chronic cough that occurs with smoking, asthma, or emphysema. Report if your cough persists.

Pharmacokinetics. Onset and Duration. PO onset 1–2 hr; duration up to 6–8 hr with non-SR, 12 hr for SR suspension.71

Fate. Extensively metabolized, including appreciable first-pass effect, mainly to the active metabolite dextrorphan. Genetically determined polymorphic metabolism primarily by CYP2D6 with extensive (93%) and poor (7%) metabolizers.72 (See Notes.)

$t\text{_{1/2}}$. (Extensive metabolizers) <4 to about 9 hr; (poor metabolizers) 17–138 hr.73

Adverse Reactions. Occasional mild drowsiness and GI upset. Intoxication, bizarre behavior, CNS depression, and respiratory depression can occur with extremely high dosages. Naloxone might be effective in reversing these effects.74–77 Reports of dextromethorphan abuse have increased, especially among teenagers.78,79

Contraindications. MAOI therapy.80

Precautions. Generally, do not use in patients with chronic cough or cough associated with excessive secretions.

Drug Interactions. Concurrent MAOIs can cause hypotension, hyperpyrexia, nausea, and coma. Drugs that inhibit CYP2D6 can inhibit dextromethorphan metabolism, but serious effects are not reported.

Parameters to Monitor. Observe for relief of cough and CNS side effects.

Notes. Approximately equipotent with codeine in antitussive effectiveness in adults.71,74 One trial of dextromethorphan and codeine for night cough in children
found neither superior to placebo, and their efficacies have been questioned for this or any other use in children.\textsuperscript{75,81} Used commonly for CYP2D6 phenotyping.\textsuperscript{82} Dextromethorphan is currently being investigated for its analgesic-sparing effect.\textsuperscript{83} \textit{(See also Codeine Salts.)}

**Pharmacology.** Guaifenesin is proposed to have an expectorant action through an increased output of respiratory tract fluid, enhancing the flow of less viscous secretions, promoting ciliary action, and facilitating the removal of inspissated mucus. Evidence of the effectiveness of guaifenesin is largely subjective and not well established clinically.\textsuperscript{74,84–87}

**Administration and Adult Dosage.** PO as an expectorant 100–400 mg q 4 hr; SR 600–1200 mg q 12 hr, to a maximum of 2.4 g/day.\textsuperscript{85}

**Special Populations.** Pediatric Dosage. PO as an expectorant (2–6 yr) 50–100 mg q 4 hr, to a maximum of 600 mg/day; (6–12 yr) 100–200 mg q 4 hr, to a maximum of 1200 mg/day; (≥12 yr) same as adult dosage. SR (2–6 yr) 300 mg q 12 hr; (6–12 yr) 600 mg q 12 hr.

**Geriatric Dosage.** Same as adult dosage.

**Dosage Forms.** Cap 200 mg; Syrup 20, 40 mg/mL; Tab 100, 200, 1200 mg; SR Cap 300 mg; SR Tab 600, 1200 mg. SR Tab 600 mg with pseudoephedrine 120 mg (Entex PSE, various).

**Patient Instructions.** Take this drug with a large quantity of fluid to ensure proper drug action. Report if your cough persists for more than 1 week, recurs, or is accompanied by a high fever, rash, or persistent headache. Excessive dosage can cause nausea and vomiting.

**Adverse Reactions.** Occasional nausea and vomiting, especially with excessive dosage; dizziness; headache.

**Precautions.** Generally, do not use in patients with chronic cough or cough associated with excessive secretions.

**Drug Interactions.** None known.

**Notes.** May interfere with certain laboratory determinations of 5-hydroxyindoleacetic acid and vanillylmandelic acid but does not cause a positive stool guaiac reaction in normal subjects.\textsuperscript{86}

**PSEUDOEPHEDRINE HYDROCHLORIDE** Efidac/24, Sudafed, Various

**Pharmacology.** Pseudoephedrine is an indirect-acting agent that stimulates α-, β\textsubscript{1}, and β\textsubscript{2}-adrenergic receptors via release of endogenous adrenergic amines. It is used primarily for decongestion of nasal mucosa.

**Administration and Adult Dosage.** PO as a decongestant 60 mg q 4–6 hr, to a maximum of 240 mg/day. PO SR Cap/Tab 120 mg q 12 hr; (Efidac/24) 240 mg once daily.

**Special Populations.** Pediatric Dosage. PO (3–12 months) 3 drops/kg q 4–6 hr, to a maximum of 4 doses/day; (1–2 yr) 7 drops (0.2 mL)/kg q 4–6 hr, to a maximum of 4 doses/day; (2–5 yr) 15 mg (as syrup) q 4–6 hr prn, to a maximum of
60 mg/day; (6–12 yr) 30 mg q 4–6 hr prn, to a maximum of 120 mg/day; (>12 yr)
same as adult dosage. Do not give SR Cap/Tab 120 or 240 mg to patients <12 yr.

**Geriatric Dosage.** Demonstrate safe use of short-acting formulation before using
an SR product.

**Dosage Forms.** **Cap** 60 mg; **Drp** 9.4 mg/mL; **Syrup** 3, 6 mg/mL; **Tab** 30, 60 mg;
**SR Tab** (12-hr) 120 mg; (24-hr) 240 mg (Efidac/24). **Tab** 60 mg with triprolidine
HCl 2.5 mg (Actifed, various). **SR Cap** 120 mg with chlorpheniramine maleate
8 mg (Deconamine SR, various).

**Patient Instructions.** Avoid taking the last dose of the day near bedtime if you
have difficulty sleeping. Do not crush or chew sustained-release preparations.

**Pharmacokinetics. Onset and Duration.** Onset within 30 min on an empty stom-
ach, within 1 hr for SR forms; duration ≥3 hr, 8–12 hr for most SR forms, 24 hr
for Efidac/24.88,89

**Fate.** Solution and immediate-release tablets are rapidly and completely absorbed
orally. SR dosage forms attain peak serum levels in (12-hr product) 4–6 hr or (24-
hr product) 12 hr. Food appears to delay absorption of non-SR forms, but not the
SR forms.90 Vd is 2.7 ± 0.2 L/kg; Cl averages 0.44 L/hr/kg. Partly metabolized
to inactive metabolite(s), and 6% metabolized to active metabolite, norpseu-
doephedrine; 45–90% excreted unchanged in urine depending on urinary pH and
flow.92,93

\( t_{1/2} \) Urinary flow and pH dependent; 13 ± 3 hr at pH 8; 6.9 ± 1.2 hr at pH 5.5–6;
4.7 ± 1.4 hr at pH 5.92,93

**Adverse Reactions.** Frequent mild transient nervousness, insomnia, irritability, or
headache. Usually negligible pressor effect in normotensive patients.94,95

**Contraindications.** Severe hypertension; coronary artery disease; MAOI therapy.

**Precautions.** Use with caution in patients with renal failure,96 hypertension, dia-
betes mellitus, ischemic heart disease, increased intraocular pressure, prostatic hy-
pertrophy, urinary retention, or thyroid disease. Elderly patients might be particu-
larly sensitive to CNS effects. If use is necessary in infants with phenylketonuria,
reduce dosage to avoid possible increased agitation.97

**Drug Interactions.** Concurrent MAOIs can increase pressor response. Urinary al-
kalinizers can decrease pseudoephedrine clearance.

**Parameters to Monitor.** Nasal stuffiness, CNS stimulation, blood pressure in hy-
pertensive patients.

**Notes.** Combination with an antihistamine can provide additive benefit in sea-
sonal allergic rhinitis because antihistamines do not relieve nasal stuffiness.98,99
Neither these combinations nor decongestants alone provide consistent long-term
benefit for reduction of middle ear effusion in children with otitis media and are
not recommended for this use.100,101

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12. Shuk S et al. Efficacy of frequent nebulized ipratropium bromide added to frequent high-dose albuterol therapy.

11. Milner AD. Ipratropium bromide in airways obstruction in childhood.

14. Chung KF. Leukotriene receptor antagonists and biosynthesis inhibitors: potential breakthrough in asthma research and clinical practice.


15. Larsen JS et al. Antileukotriene therapy for asthma.


PART II

Clinical Information

Principal Editor: William G. Troutman, PharmD

- Drug-Induced Diseases
- Drug Use in Special Populations
- Immunization
- Medical Emergencies
- Drug Interactions and Interferences
- Nutrition Support
Drug-Induced Blood Dyscrasias

This table does not include all drugs capable of causing the specified dyscrasias and excludes cancer chemotherapeutic agents, which are known for producing dose-related bone marrow suppression. Five major types of blood dyscrasias have been selected for inclusion in this table; the following abbreviations indicate specific blood dyscrasias:

- AA — Aplastic Anemia
- AGN — Agranulocytosis, Granulocytopenia, or Neutropenia
- HA — Hemolytic Anemia
- MA — Macrocytic Anemia
- Th — Thrombocytopenia

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<tr>
<th>DRUG AND DYSRASIA</th>
<th>NATURE OF DYSRASIA</th>
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<tr>
<td><strong>Abciximab</strong></td>
<td></td>
</tr>
<tr>
<td>Th</td>
<td>The combination of abciximab and heparin presents twice the risk of mild and severe thrombocytopenia as the combination of placebo and heparin. (See also Heparin.)¹</td>
</tr>
<tr>
<td><strong>Acetaminophen</strong></td>
<td></td>
</tr>
<tr>
<td>Th</td>
<td>Scattered reports only; observed in 6 of 174 overdose patients in one report; might be an immune reaction.²³</td>
</tr>
<tr>
<td><strong>Alcohol</strong></td>
<td></td>
</tr>
<tr>
<td>HA</td>
<td>Most commonly encountered in chronic alcoholism.⁴</td>
</tr>
<tr>
<td>MA</td>
<td>Results from malnutrition and decreased folate absorption and/or utilization. Responds rapidly to folic acid administration.⁴</td>
</tr>
<tr>
<td>Th</td>
<td>Transient in many drinkers; persistent thrombocytopenia can accompany advanced alcoholic liver disease.⁴</td>
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<tr>
<th>DRUG AND DYSCRASIA</th>
<th>NATURE OF DYSCRASIA</th>
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</table>
| **Amphotericin B** | AGN: Scattered reports only.\(^{4,5}\)  
                      Th: Scattered reports only.\(^{5,6}\) |
| **Antidepressants, Heterocyclic** | AGN: Idiosyncratic reaction, probably resulting from a direct toxic effect rather than allergy. Most commonly occurs between the 2nd and 8th weeks of therapy.\(^{4,10,11}\) |
| **Ascorbic Acid** | HA: In G-6-PD deficiency with large doses.\(^{4}\) |
| **Aspirin** | HA: Almost always encountered in patients with G-6-PD deficiency, usually in conjunction with infection or other complicating factors.\(^{4,12}\)  
                      Th: Can occur in addition to the drug’s effects on platelet adhesiveness. Some evidence for an immune reaction.\(^{2,4,13}\) |
| **Azathioprine** | AGN: WBC counts <2500/\(\mu\)L occur in about 3% of rheumatoid arthritis patients treated with azathioprine; an additional 15% develop some lesser degree of leukopenia.\(^{14}\) |
| **Captopril** | AGN: Prevalence estimated at 1/5000 patients. The prevalence increases greatly in patients with reduced renal function or collagen–vascular diseases and reaches 7% in patients with renal impairment and a collagen–vascular disease. Most common during the first 3 weeks of therapy.\(^{15}\) |
| **Carbamazepine** | AA: 27 cases reported from 1964–1988; onset can be delayed until weeks or months after the initiation of therapy.\(^{4,16}\)  
                      AGN: Transient leukopenia occurs in about 10% of patients, usually during the first month of therapy. Recovery usually occurs within a week of drug withdrawal. Persistent leukopenia occurs in 2%.\(^{16,17}\)  
                      Th: Prevalence estimated at 2%.\(^{16,18}\) |
| **Cephalosporins** | AGN: Rare; possibly the result of an immune reaction but occurs most often with high dosages and parenteral therapy lasting >2 weeks.\(^{4,19,20}\)  
                      HA: Positive direct Coombs’ test occurs frequently and can persist for up to 2 months after discontinuation of therapy. Hemolysis is rare.\(^{4,19}\)  
                      Th: Rare; possibly the result of an immune reaction. Usually occurs late in the course of therapy.\(^{4,19}\) |

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<tr>
<th>DRUG AND DYSCRASIA</th>
<th>NATURE OF DYSCRASIA</th>
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<tbody>
<tr>
<td><strong>Chloramphenicol</strong></td>
<td></td>
</tr>
<tr>
<td>AA</td>
<td>Prevalence estimated at 1/12,000 to 1/50,000 patients. Most cases develop with oral administration and after discontinuation of therapy, suggesting the development of a toxic metabolite. An association between the ophthalmic use of chloramphenicol and the development of aplastic anemia is weak, if it exists at all. Blacks might be more susceptible than whites. Do not confuse with the dose-related anemia seen with chloramphenicol. (Note: One case report suggests that a patient’s dose-related anemia might have progressed to aplastic anemia, but most sources separate the two dyscrasias.)</td>
</tr>
<tr>
<td>AGN</td>
<td>Rare when compared with the prevalence of aplastic anemia.</td>
</tr>
<tr>
<td>HA</td>
<td>In G-6-PD deficiency.</td>
</tr>
<tr>
<td><strong>Chloroquine</strong></td>
<td></td>
</tr>
<tr>
<td>AGN</td>
<td>Scattered reports only; might be dose related.</td>
</tr>
<tr>
<td>HA</td>
<td>Only a few cases have been reported; some association with G-6-PD deficiency is suspected.</td>
</tr>
<tr>
<td><strong>Cimetidine</strong></td>
<td></td>
</tr>
<tr>
<td>AA</td>
<td>Scattered reports only; however, at least two fatalities reported (one fatality also was receiving chloramphenicol).</td>
</tr>
<tr>
<td>AGN</td>
<td>Usually occurs in patients with systemic disease or other drug therapy that might have contributed to the dyscrasia.</td>
</tr>
<tr>
<td><strong>Clopidogrel</strong></td>
<td></td>
</tr>
<tr>
<td>Th</td>
<td>At least 11 cases of clopidogrel-associated thrombotic thrombocytopenic purpura have been reported. Most cases occurred during the first 2 weeks of treatment.</td>
</tr>
<tr>
<td><strong>Clozapine</strong></td>
<td></td>
</tr>
<tr>
<td>AGN</td>
<td>Frequency of granulocytopenia is calculated to be 0.4–0.8% in closely monitored patients. Mild to moderate neutropenia occurs in 3–20%. Most cases occur in the first 4 months. Asians are more than twice as susceptible as whites. Recovery usually occurs 2–3 weeks after drug withdrawal. Frequent WBC counts are mandated.</td>
</tr>
<tr>
<td><strong>Cocaine</strong></td>
<td></td>
</tr>
<tr>
<td>Th</td>
<td>Reported with IV and inhalational use.</td>
</tr>
<tr>
<td><strong>Contraceptives, Oral</strong></td>
<td></td>
</tr>
<tr>
<td>MA</td>
<td>Results from impaired folate absorption and/or activity; of consequence only if the patient’s folate status is markedly impaired.</td>
</tr>
<tr>
<td><strong>Dapsone</strong></td>
<td></td>
</tr>
<tr>
<td>AGN</td>
<td>Many cases have occurred during combination therapy, so it is difficult to determine if dapsone alone is the causative agent.</td>
</tr>
<tr>
<td>HA</td>
<td>In G-6-PD deficiency; might have other mechanism(s). Might be dose related; uncommon at 100 mg/day but frequent at 200–300 mg/day.</td>
</tr>
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<tr>
<th>DRUG AND DYSCRASIA</th>
<th>NATURE OF DYSCRASIA</th>
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<tbody>
<tr>
<td>Digoxin</td>
<td>Scattered reports only; evidence of an immune mechanism.²,³²,³³</td>
</tr>
<tr>
<td>Dimercaprol</td>
<td>In G-6-PD deficiency.⁴</td>
</tr>
<tr>
<td>Dipyridamole</td>
<td>Relative risk of thrombocytopenia calculated to be 14 times higher than in untreated individuals, but needs confirmation.³⁴</td>
</tr>
<tr>
<td>Diuretics, Thiazide</td>
<td>Exact mechanism is unclear; might be an immune reaction.⁴,³⁵</td>
</tr>
<tr>
<td>Diuretics, Thiazide</td>
<td>Mild thrombocytopenia occurs frequently, but severe cases are rare. Might be caused by an immune reaction.²,⁴,³⁶</td>
</tr>
<tr>
<td>Eflornithine</td>
<td>Deaths caused by aplastic anemia have been reported.³⁷</td>
</tr>
<tr>
<td>Eflornithine</td>
<td>Leukopenia is reported in 18–37% of patients.³⁷</td>
</tr>
<tr>
<td>Eflornithine</td>
<td>Megaloblastic anemia is frequently reported.³⁷</td>
</tr>
<tr>
<td>Eflornithine</td>
<td>Thrombocytopenia is frequently reported.³⁷</td>
</tr>
<tr>
<td>Etanercept</td>
<td>Although the causal relationship is unclear, some cases of aplastic anemia, including fatalities, have been associated with etanercept.¹¹³</td>
</tr>
<tr>
<td>Felbamate</td>
<td>More than 30 cases were reported shortly after the introduction of felbamate, resulting in the manufacturer and FDA urging withdrawal of patients from therapy. When a strict definition of aplastic anemia is applied and confounding factors are accounted for, the risk of aplastic anemia from felbamate might not be markedly different from the risk posed by carbamazepine. Most cases developed 2–6 months after initiation of therapy. Monitoring has not been effective for early identification of cases.³⁸,³⁹</td>
</tr>
<tr>
<td>Fluconazole</td>
<td>Scattered reports only.⁴⁰</td>
</tr>
<tr>
<td>Flucytosine</td>
<td>Dose-related; usually requires plasma concentrations ≥125 mg/L.⁴¹</td>
</tr>
<tr>
<td>Foscarnet</td>
<td>Neutropenia occurs in 14% of patients treated for cytomegalovirus retinitis.⁴²</td>
</tr>
<tr>
<td>Furosemide</td>
<td>Uncommon, mild, and asymptomatic.³</td>
</tr>
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<tr>
<th>DRUG AND DYSCRASIA</th>
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<tbody>
<tr>
<td><strong>Ganciclovir</strong></td>
<td></td>
</tr>
<tr>
<td>AGN</td>
<td>Granulocytopenia occurs in about 40% of patients; it is usually reversible with drug discontinuation, but irreversible neutropenia and deaths have occurred.42,43</td>
</tr>
<tr>
<td>Th</td>
<td>Thrombocytopenia occurs in about 20% of patients.43</td>
</tr>
<tr>
<td><strong>Gold Salts</strong></td>
<td></td>
</tr>
<tr>
<td>AA</td>
<td>Not dose-dependent; although this reaction is not common, numerous fatalities have been reported.14,44</td>
</tr>
<tr>
<td>AGN</td>
<td>Often brief and self-limiting; usually responds to withdrawal of therapy.45,46</td>
</tr>
<tr>
<td>Th</td>
<td>Not dose- or duration-dependent; prevalence estimated at 1–3%; onset usually during the loading phase (first 1000 mg) but can be delayed until after the drug has been discontinued. Mechanism is unclear, but it often appears to be immunologically mediated. Up to 85% of patients with gold-induced thrombocytopenia have HLA-DR3 phenotype compared with 30% of all rheumatoid arthritis patients.2,4,47,48</td>
</tr>
<tr>
<td><strong>Heparin</strong></td>
<td></td>
</tr>
<tr>
<td>Th</td>
<td>Many patients demonstrate a mild to moderate transient decrease in platelets after only a few days of heparin therapy. Up to 3% experience immune-mediated, persistent thrombocytopenia, which is associated with increased thrombin generation and development of serious thrombotic complications in 30–60%. Intermittent, continuous infusion and &quot;minidose&quot; regimens have been implicated; this is uncommon with SC administration. Prompt cessation of heparin minimizes serious complications; platelet count usually returns to normal within 7–10 days. Low-molecular-weight heparins (e.g., dalteparin, enoxaparin, tinzaparin) are much less likely than unfractionated heparin to stimulate the formation of immune complexes, leading to thrombocytopenia. Low-molecular-weight heparins offer very little protection from thrombocytopenia in patients who have already formed heparin-associated antibodies.49–52</td>
</tr>
<tr>
<td><strong>Immune Globulin</strong></td>
<td></td>
</tr>
<tr>
<td>AGN</td>
<td>Transient neutropenia frequently accompanies IV use.53</td>
</tr>
<tr>
<td>HA</td>
<td>Acute Coombs’ positive hemolysis has been reported in patients receiving high-dose therapy.53</td>
</tr>
<tr>
<td><strong>Inamrinone</strong></td>
<td></td>
</tr>
<tr>
<td>Th</td>
<td>18.6% prevalence in one study of oral therapy (oral form not marketed in the United States); the prevalence during parenteral therapy has been estimated at 2.4%, although 8 of 16 children receiving parenteral inamrinone developed thrombocytopenia in one report. Thrombocytopenia might be caused by nonimmune peripheral platelet destruction.7–9</td>
</tr>
<tr>
<td><strong>Indomethacin</strong></td>
<td></td>
</tr>
<tr>
<td>AA</td>
<td>Although rare, indomethacin has been associated with a risk 12.7 times higher than in untreated individuals, especially when used regularly and for a long duration.54</td>
</tr>
<tr>
<td>AGN</td>
<td>Although rare, risk can be 8.9 times higher than in untreated individuals.54</td>
</tr>
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<tr>
<th>DRUG AND DYSCRASIA</th>
<th>NATURE OF DYSCRASIA</th>
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<tbody>
<tr>
<td>Interferon Alfa</td>
<td>Scattered reports only.</td>
</tr>
<tr>
<td>Isoniazid</td>
<td>Scattered reports; some evidence of an immune reaction.</td>
</tr>
<tr>
<td>Lamotrigine</td>
<td>Scattered reports; too early to establish a pattern of risk.</td>
</tr>
<tr>
<td>Levalisole</td>
<td>Might be the result of an autoimmune reaction, with a prevalence of ≥4% in some series. Presence of the HLA-B27 phenotype in seropositive rheumatoid arthritis might be an important predisposing factor.</td>
</tr>
<tr>
<td>Levodopa</td>
<td>Autoimmune reaction; positive direct and indirect Coombs’ tests are frequent, but hemolysis is rare. Carbipoda–levodopa combinations also have produced hemolysis.</td>
</tr>
<tr>
<td>Mefenamic Acid</td>
<td>Thought to be autoimmune.</td>
</tr>
<tr>
<td>Methimazole</td>
<td>Scattered reports only, but some increased risk is present. Most cases occur during the first 3 months of therapy.</td>
</tr>
<tr>
<td>Methyldopa</td>
<td>Autoimmune reaction; positive direct Coombs’ test occurs in 5–25% of patients, depending on dosage; hemolysis occurs in &lt;1%, and its onset is gradual after ≥4 months of therapy. Recovery is rapid after discontinuation of the drug.</td>
</tr>
<tr>
<td>Methylene Blue</td>
<td>In G-6-PD deficiency.</td>
</tr>
<tr>
<td>Nalidixic Acid</td>
<td>In G-6-PD deficiency; might have other mechanisms.</td>
</tr>
<tr>
<td>Nitrofurantoin</td>
<td>In G-6-PD deficiency; also encountered with enolase deficiency (mechanism unknown).</td>
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<tr>
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<tbody>
<tr>
<td><strong>Penicillamine</strong></td>
<td></td>
</tr>
<tr>
<td>AA Rare; develops after several months of therapy; due to direct marrow toxicity.</td>
<td></td>
</tr>
<tr>
<td>AGN Rare; most cases occur during the first month of therapy.</td>
<td></td>
</tr>
<tr>
<td>HA Scattered reports only; might be caused by G-6-PD deficiency or fluctuations in copper levels during therapy of Wilson’s disease.</td>
<td></td>
</tr>
<tr>
<td>Th Prevalence estimated at 10%; some decrease in platelet counts occurs in 75% of penicillamine-treated patients. Might be the result of an immune reaction; most commonly occurs during the first 6 months of therapy.</td>
<td></td>
</tr>
<tr>
<td><strong>Penicillins</strong></td>
<td></td>
</tr>
<tr>
<td>AA Prevalence very low when extent of use is considered.</td>
<td></td>
</tr>
<tr>
<td>AGN Uncommon with most penicillins but frequent with methicillin; in one report, neutropenia developed in 23 of 68 methicillin-treated patients; resolution occurred within 3–7 days after drug withdrawal. The risk of penicillin-induced neutropenia is increased with parenteral treatment lasting &gt;2 weeks.</td>
<td></td>
</tr>
<tr>
<td>HA Positive direct Coombs’ test occurs with large IV doses; hemolysis is rare.</td>
<td></td>
</tr>
<tr>
<td><strong>Phenazopyridine</strong></td>
<td></td>
</tr>
<tr>
<td>HA Prevalence and mechanism unknown; renal insufficiency and overdose might be contributing factors. Often accompanied by methemoglobinemia.</td>
<td></td>
</tr>
<tr>
<td><strong>Phenobarbital</strong></td>
<td></td>
</tr>
<tr>
<td>MA More than 100 cases reported; usually responds to folic acid.</td>
<td></td>
</tr>
<tr>
<td><strong>Phenothiazines</strong></td>
<td></td>
</tr>
<tr>
<td>AGN Most common during the first 2 months of therapy and in older patients (&gt;85% are &gt;40 yr). Rapid onset and general lack of dose dependence suggest an idiosyncratic mechanism. Prevalence estimated as high as 1/1200.</td>
<td></td>
</tr>
<tr>
<td><strong>Phenytoin</strong></td>
<td></td>
</tr>
<tr>
<td>AA Fewer than 25 reported cases, but the association with phenytoin is strong.</td>
<td></td>
</tr>
<tr>
<td>AGN Scattered reports only; onset after days to years of therapy.</td>
<td></td>
</tr>
<tr>
<td>MA Caused by impaired absorption and/or utilization of folate and responds to folic acid therapy (although folate replacement can lower phenytoin levels). Mild macrocytosis is very common (&gt;25%); onset is unpredictable but usually appears after &gt;6 months of therapy.</td>
<td></td>
</tr>
<tr>
<td>Th Scattered reports only; might be the result of an immune reaction.</td>
<td></td>
</tr>
<tr>
<td><strong>Primaquine</strong></td>
<td></td>
</tr>
<tr>
<td>HA In G-6-PD deficiency.</td>
<td></td>
</tr>
<tr>
<td><strong>Primidone</strong></td>
<td></td>
</tr>
<tr>
<td>MA Similar to phenobarbital, but prevalence might be lower; onset is unpredictable and can be delayed for several years during therapy. Some cases have responded to folic acid.</td>
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</tr>
</tbody>
</table>

(continued)
<table>
<thead>
<tr>
<th>DRUG AND DYSCRASIA</th>
<th>NATURE OF DYSCRASIA</th>
</tr>
</thead>
</table>
| **Procainamide**   | AGN Prevalence usually estimated at <1%, but with a 25% fatal outcome. Occurs with conventional and sustained-release products; usually occurs within the first 90 days of use. No relationship with daily or total dosage.  
AA Scattered reports only, but some increased risk is present. Most cases occur within the first 3 months of therapy.  
AGN Prevalence estimated at 0.55%. Occurs overwhelmingly in women and appears to increase with age. Most cases occur in the first 3 months of therapy, and monitoring during this time might detect agranulocytosis before it becomes clinically apparent. Some evidence for an immune reaction.  
HA In G-6-PD deficiency; usually requires concurrent infection or other complicating factors.  
Th Caused by quinidine-specific antibodies; little or no cross-reactivity with quinine. Accounts for a large portion of drug-induced thrombocytopenia. |
| **Propylthiouracil** | AA Scattered reports only.  
AGN Prevalence estimated at 0.55%. Occurs overwhelmingly in women and appears to increase with age. Most cases occur in the first 3 months of therapy, and monitoring during this time might detect agranulocytosis before it becomes clinically apparent. Some evidence for an immune reaction.  
|
| **Quinacrine** | AA About one-half of reported cases were preceded by a rash or lichenoid eruption; prevalence estimated at 3/100,000.  
HA In G-6-PD deficiency; usually requires concurrent infection or other complicating factors.  
|
| **Quinidine** | AGN Scattered reports only; an immune mechanism has been described.  
HA In G-6-PD deficiency (but not in blacks). A rapid onset immune mechanism has also been described.  
Th Caused by quinidine-specific antibodies; little or no cross-reactivity with quinine. |
| **Quinine** | AGN Scattered reports only.  
HA In G-6-PD deficiency (but not in blacks). An immune mechanism is also suspected because quinine-dependent antibodies to RBCs have been demonstrated in cases of quinine-induced hemolytic-uremic syndrome.  
Th Caused by quinidine-specific antibodies; little or no cross-reactivity with quinidine. Fatalities have been reported. It has occurred in people drinking quinine-containing tonic water.  
|
| **Rifabutin** | AGN In a study of the pharmacokinetic interactions between rifabutin and azithromycin or clarithromycin, rifabutin, alone or in combination with either of those drugs, produced neutropenia in most of the patients. Neutropenia was not seen when either of the other drugs was used without rifabutin.  
|
| **Rifampin** | HA Rare but many patients develop a positive Coombs’ test; onset in hours in some sensitized patients.  
Th Peripheral destruction of platelets appears to result from an immune reaction; difficult to separate rifampin contribution from that of other drugs because it is usually used in combination therapy. (continued) |
### Sulfasalazine

| AGN | Leukopenia reported in 5.6% of patients receiving the drug for rheumatoid arthritis and agranulocytosis/neutropenia in 4/1000 patients; prevalence of agranulocytosis/neutropenia among inflammatory bowel disease patients is considerably lower (0.3/1000 patients). Onset is usually during the first 3 months of therapy; recovery takes 2 weeks after drug discontinuation. |
| HA | In G-6-PD deficiency but also occurs in nondeficient patients. Hemolysis might be more common in slow acetylators. |
| MA | One series of 130 arthritis patients reported macrocytosis in 21% and macrocytic anemia in 3%. |

### Sulfonamides

| AA | Historically an important cause of aplastic anemia, but most cases were reported after use of older sulfonamides; rarely occurs with products currently in use. |
| AGN | Occurs mostly with older products; rarely occurs with products currently in use. Most current cases are in combined use with trimethoprim; also reported with silver sulfadiazine. Onset is usually rapid. |
| HA | In G-6-PD deficiency but also occurs in nondeficient patients. |
| Th | Scattered reports only; probably an immune reaction. (See also Trimethoprim.) |

### Ticlopidine

| AA | The growing number of cases of aplastic anemia associated with ticlopidine is disturbing; the incidence cannot be estimated. |
| AGN | Incidence of neutropenia estimated at 2.4% of treated patients with severe neutropenia or agranulocytosis in 0.85%. Obtain CBC every 2 weeks during the first 3 months of treatment. Discontinue ticlopidine if the ANC is <1200/µL. |
| HA | Thrombotic thrombocytopenia purpura occurs in 1 of every 1600–5000 exposed. Mean time to onset is 22 days. Plasmapheresis reduces the death rate from 60% to 21%. |

### Tocainide

| AGN | Prevalence estimated at 0.07–0.18% of patients. |

### Triamterene

| MA | Few cases reported, but it is a potent inhibitor of dihydrofolate reductase; greatest risk in those with folate deficiency before therapy (eg, alcoholics). |

### Trimethoprim

| AGN | Rare; occurs when used alone and in combination with sulfonamides, with the latter numerically more common. |
| MA | Most cases occur after 1–2 weeks of therapy; this drug can have weak antifolate action in humans that becomes important only in those with folate deficiency before therapy (eg, alcoholics). |
| Th | Thrombocytopenia is common, but severe cases are rare. Most commonly occurs in combination therapy with sulfonamides. Relative risk calculated at 124 times that of untreated individuals. |

(continued)
### DRUG AND
dyscrasia

<table>
<thead>
<tr>
<th>Vaccines</th>
<th>NATURE OF DYSCRASIA</th>
</tr>
</thead>
<tbody>
<tr>
<td>Th</td>
<td>A study of 9 million doses of measles, rubella, and mumps vaccines administered to children determined that the prevalence of thrombocytopenia was 0.17 cases/100,000 doses for measles vaccine and 0.23, 0.87, and 0.95 cases/100,000 doses for rubella, measles–rubella, and mumps–measles–rubella vaccines, respectively. These rates are similar to the rates of thrombocytopenia after the natural courses of the disease in unvaccinated children. Most of the cases had platelet counts &gt;10,000/µL.</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Valproic Acid</th>
<th>MA</th>
<th>Macrocystosis occurred in 11 of 60 patients in one report.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Th</td>
<td>Thrombocytopenia occurred in 12 of 60 patients in one report. Immune and dose-dependent mechanisms have been suggested.</td>
<td></td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Vancomycin</th>
<th>AGN</th>
<th>Scattered reports only, but prevalence might be as high as 2%; mechanism unknown.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Vesnarinone</td>
<td>AGN</td>
<td>Reversible neutropenia occurs in about 3%, mostly in the first 16–24 weeks of treatment. Absolute granulocyte count &lt;1 × 10^9/L occur in 0.85%, with counts &lt;0.1 × 10^9/L in 0.25%.</td>
</tr>
</tbody>
</table>

| Vitamin K | HA | In G-6-PD deficiency; usually requires concurrent infection or other complicating factors. Hemolysis from high doses can contribute to jaundice in neonates; rarely toxic in older children and adults. |

<table>
<thead>
<tr>
<th>Zidovudine</th>
<th>AGN</th>
<th>Most patients experience at least a 25% reduction in neutrophil count; ANC of &lt;500/µL occurs in 16% of patients. Usual onset is during the first 3 months of therapy.</th>
</tr>
</thead>
<tbody>
<tr>
<td>MA</td>
<td>Macrocytosis develops in most patients, usually beginning during the first few weeks of therapy. Zidovudine is the leading cause of drug-induced macrocytosis.</td>
<td></td>
</tr>
</tbody>
</table>

### REFERENCES


Drug-Induced Hepatotoxicity

This table includes only those drugs with well-established records of hepatotoxicity. A drug not listed in the table does not mean it cannot produce liver damage because virtually all drugs have been reported to elevate serum liver enzymes. Combining drugs that have hepatotoxic potential commonly results in greater than additive liver damage. In general, drug-induced hepatotoxicity is most prevalent in older patients, women, and those with pre-existing hepatic impairment.

**ACE Inhibitors**

Hepatic injury occurs occasionally with ACE inhibitors. Captopril and enalapril are implicated in most reported cases, but other ACE inhibitors likely have similar hepatotoxic potential. Most cases show cholestatic injury, but mixed and hepatocellular damage also are reported.1,2

**Acetaminophen**

Centrilobular hepatic necrosis can follow acute overdose with ≥140 mg/kg in children or ≥6 g in adults. These doses saturate the normal metabolic pathways, producing large quantities of a hepatotoxic metabolite. Children appear to have a lower risk than adults of developing acetaminophen-induced hepatitis. Laboratory evidence of hepatotoxicity peaks 3–4 days after the acute exposure. Therapy with acetylcysteine to bind the metabolite is indicated when the 4-hr postingestion serum acetaminophen level is >150 mg/L. Even without acetylcysteine, fatalities are uncommon after acetaminophen overdose. Nonfatal cases usually recover fully in a few weeks. Chronic alcohol ingestion increases acetaminophen toxicity, as does recent fasting. Acute alcohol ingestion is thought by some to have a protective action. Less destructive, but still detectable, hepatitis is reported in patients taking large acetaminophen doses for therapeutic purposes.1,3–5

**Alcohol**

Fatty infiltration of the liver occurs in 70–100% of alcoholics. Fatty liver is generally without clinical manifestation, but 30% of alcoholics develop alcoholic hepatitis and about 10% develop cirrhosis. Malnutrition can potentiate alcoholic liver disease, and alcohol can enhance the hepatotoxicity of other drugs.1

**Aldesleukin**

Increases in serum bilirubin, alkaline phosphatase, and transaminases occur frequently. These primarily cholestatic changes are rapidly reversible after drug discontinuation.1,6

**Allopurinol**

Hepatic granulomas, hepatitis and hepatic necrosis can accompany other symptoms (especially rash, fever, eosinophilia, and vasculitis) of allopurinol hypersensitivity. Damage is usually focal, but widespread damage also is reported. This reaction is rare but serious when it occurs. Onset is usually after 3–6 weeks of treatment. Renal impairment might be a predisposing factor for allopurinol-induced hepatitis. Cholestasis also has been attributed to allopurinol.1,7

**Aminoglutethimide**

Laboratory evidence of cholestasis is common, but clinical evidence is rare.1,8

**Aminosalicylic Acid**

Up to 5% of patients develop a generalized hypersensitivity reaction. About 25% of these patients have evidence of mixed cholestatic and hepatocellular injuries as part of their hypersensitivity reactions. Fatalities have been reported.1,9

(continued)
Amiodarone

Mild increases in transaminases and LDH levels occur in up to one-half of patients, whereas phospholipidosis occurs in virtually all; normal values often return despite continued therapy. Symptoms (e.g., jaundice, nausea and vomiting, hepatomegaly, or weight loss) occur in 1–4% of patients. Onset is typically after 2–4 months of therapy but can be delayed for ≥1 yr. Recovery after drug discontinuation can take from several months to ≥1 yr. The dose-related hepatotoxicity of amiodarone is reminiscent of alcoholic hepatitis. Cirrhosis and fatalities are also reported.1,10–12

Amoxicillin and Clavulanic Acid

Based on an extensive review of medical records, the frequency of acute hepatic injury with amoxicillin and clavulanic acid is 1.7 cases/10,000 prescriptions (compared with 0.3 for amoxicillin alone). In most cases, the hepatic injury is cholestatic. The risk of hepatic injury is increased by repeated prescriptions for amoxicillin and clavulanic acid and by advancing age.2,13

Androgens

(See Steroids, C-17-α-Alkyl.)

Antidepressants, Heterocyclic

The prevalence of hepatic injury is estimated at about 1%, with most of the cases presenting as cholestasis. This idiosyncratic reaction resembles the cholestasis associated with phenothiazines.1

Asparaginase

Slowly reversible steatosis occurs in 50–90% of patients, apparently due to the drug’s influence on protein synthesis. Daily administration might be more hepatotoxic than weekly administration.1,14–16

Azathioprine

This drug is less hepatotoxic than its metabolite, mercaptopurine. Azathioprine’s hepatotoxicity is predominantly cholestatic rather than hepatocellular. Vascular lesions, including venous occlusion and peliosis hepatis, have been reported, but their prevalence is unknown. Nodular regenerative hyperplasia has followed use of this drug in kidney and liver transplantations.1,14,17

Busulfan

Use in bone marrow transplant patients is associated with apparently dose-related veno-occlusive disease of the liver. Although the exact contribution of the drug is difficult to discern, this syndrome occurs in 20% of adults and 5% of children with total doses ≥16 mg/kg.1,14,18,19

Carbamazepine

Mild changes in liver function tests occur frequently. Hepatic necrosis, granulomas, and cholestasis have occurred, with some cases showing signs of hypersensitivity. Onset is most often in the first 4 weeks of therapy. Fatalities have been reported.1,20

Carmustine

Changes in liver function tests in 20–30% of patients, from a few days to several weeks after drug administration. Changes are usually mild and resolve quickly with drug discontinuation.1,14

Cephalosporins

Transient minor increases in AST, ALT, and alkaline phosphatase occur frequently. Ceftriaxone use is associated with development of “gallbladder sludge” in up to 25% of patients.1,21

Chlorpropamide

Most hepatotoxic reactions are cholestatic and probably are caused by an immune mechanism. Prevalence is estimated at 0.5–1.5%, with onset usually within the first 2 months of therapy.1

(continued)
Chlorzoxazone

Idiosyncratic hepatocellular damage occurs rarely, but fatalities have been reported. Discontinue the drug if elevated levels of transaminases or bilirubin are detected.\(^\text{1,23}\)

Cisplatin

Transient, dose-related elevations of hepatic enzymes occur frequently.\(^\text{1,14}\)

Clozapine

Transient elevations of hepatic enzymes occur frequently during the first 3 months of clozapine use. Although several cases of fulminant hepatitis have been reported, the risk of serious clozapine-induced hepatotoxicity remains small and some investigators recommend against routine testing.\(^\text{1,24,25}\)

Cocaine

Hepatic necrosis has been reported in cases of cocaine abuse, including at least one fatality. The prevalence of this reaction is not known.\(^\text{1,10,26}\)

Contraceptives, Oral

Data from two large, long-term cohort studies (about 33,000 users) did not detect any association between oral contraceptive use and serious liver disease. One study detected an increase in the frequency of mild liver disease among users of older, high-estrogen (>50 \(\mu\)g) products. Older combination oral contraceptives were associated with an increase in the annual incidence of hepatic adenomas (3.4/100,000 vs 1.3/100,000 in nonusers), especially after \(\geq 5\) yr of use. The frequency of gallbladder disease also was increased by older oral contraceptives.\(^\text{1,27,28}\)

Cyclosporine

Elevated serum levels of alkaline phosphatase and conjugated bilirubin consistent with cholestasis occur in 50–60% of patients. These changes are usually mild and pose little threat.\(^\text{1,29,30}\)

Dantrolene

At least 1.8% of patients develop laboratory evidence of hepatic dysfunction, with symptomatic hepatitis in about 0.6%; the fatality rate among jaundiced patients is about 25%. Predisposing factors seem to include dosage (>300 mg/day), sex (women more than men), age (>30 yr), and duration of therapy (\(\geq 2\) months).\(^\text{1,31,32}\)

Dapsone

Hepatitis can occur as part of the “dapsone syndrome,” a generalized hypersensitivity reaction that includes rash, fever, and lymphadenopathy. The true prevalence is unknown but might be as high as 5%. The onset is usually during the first 2 months of therapy. Although most dapsone-associated liver injury is hepatocellular, some cases of cholestasis have occurred.\(^\text{1,33–35}\)

Disulfiram

Small increases in serum transaminase levels occur frequently. Hepatitis is reported occasionally, which can be caused by hypersensitivity. Most cases develop during the first few months of treatment. The best estimate of the incidence of fatal hepatitis is about 1/30,000 users/year.\(^\text{1,36}\)

Erythromycin

Erythromycin was thought to be a frequent cause of jaundice, but recent studies indicate that jaundice occurs only occasionally. Cholestasis apparently results from hypersensitivity (60% have eosinophilia and 50% have fever), appearing after 10–14 days of initial therapy or after 1–2 days in patients with a history of erythromycin exposure. Despite extensive use in children, most cases are reported in adults. Rapid reversal of symptoms follows drug discontinuation, but laboratory (continued)
changes can persist for up to 6 months. Although most cases involve the estolate salt, hepatotoxicity has occurred with the ethylsuccinate, stearate, and propionate salts and with erythromycin base.\textsuperscript{1,37–39}

**Ethionamide**

Hepatitis can occur in 3–5% of patients, and serum enzyme elevations can occur in ≥30%. Onset of hepatitis is usually after several months of therapy.\textsuperscript{1,40}

**Felbamate**

Although the prevalence of hepatocellular destruction is unclear, it is of sufficient concern to limit the use of felbamate to carefully selected patients. At least 6 cases of fatal felbamate-induced hepatic necrosis have been reported.\textsuperscript{1,42}

**Ferrous Salts**

Hepatic necrosis can appear within 1–3 days of an acute overdose. The fatality rate is high if the patient is not treated promptly.\textsuperscript{1}

**Floxuridine**

Hepatic arterial infusion of floxuridine results in 9% sclerosing cholangitis at 9 months and 26% after 1 yr. Elevations of liver enzyme levels are common but not predictive of greater hepatotoxicity.\textsuperscript{2,16,43}

**Flutamide**

Through 1994, there were at least 20 reported deaths reasonably attributed to flutamide-induced hepatotoxicity. Those deaths, typically the result of massive hepatic necrosis, occurred between 5 days and 9 months (mean 3 months) after initiation of flutamide therapy. Further, the hospitalization rate for noninfectious liver disease in flutamide-treated patients was 10 times the expected rate. Monthly liver function testing is recommended for the first 4 months.\textsuperscript{1,44,45}

**Gold Salts**

Cholestasis occurs occasionally with normal doses of parenteral gold salts; hypersensitivity is the suspected mechanism. Onset is commonly within the first few weeks of therapy, and recovery usually occurs within 3 months after drug discontinuation. Lipogranulomas are frequently found in liver biopsies of parenteral gold-treated patients. These can persist long after drug withdrawal but do not seem to impair liver function. Hepatic necrosis can result from overdose.\textsuperscript{1,46,47}

**Halothane**

As many as 30% of patients have increased serum transaminases or other evidence of mild hepatic impairment. Despite extensive publicity, the actual frequency of severe halothane hepatitis is low, ranging from 1/3500 to 1/35,000, with reported case fatality rates of 14–67%. Susceptibility is greatest in adults, women, obese patients, and especially in patients with prior exposure to halothane. The mechanism of hepatitis is poorly understood, but hypersensitivity is most likely. Fever precedes jaundice in most patients. The onset of jaundice is usually 5–8.5 days after exposure but can occur 1–26 days after exposure; shorter latent periods are associated with prior halothane exposure. Methoxyflurane and enfurane produce similar hepatotoxic reactions, although less frequently.\textsuperscript{1,46,47}

**Histamine H$_2$-Receptor Antagonists**

Cimetidine and ranitidine are associated with increased liver enzymes. The risk of acute liver injury with cimetidine is about 1/5000, with most cases occurring during the first 2 months of use.\textsuperscript{1,50}

(continued)
Isoniazid
Elevated serum transaminase levels occur frequently, are presumed to be associated with subclinical hepatitis, resolve rapidly after drug discontinuation, and can resolve despite continued isoniazid therapy. A syndrome resembling viral hepatitis occurs in 1–2% of patients, with the onset usually during the first 20 weeks of therapy. The fatality rate from isoniazid hepatitis has fallen steadily over the past 2 decades, probably in response to more aggressive monitoring, and is now estimated to be 1–1.7/100,000 patients starting isoniazid and 1.5–2.9/100,000 patients completing a course of therapy. Most fatalities occur in women. Despite the widespread assumption that patients <35 yr are unlikely to develop isoniazid-induced fatal hepatotoxicity, reported deaths indicate otherwise. Alcohol consumption increases the risk of hepatotoxicity; the contribution of concomitant rifampin is poorly defined. The role of acetylator phenotype remains unclear, but a case-control study found that patients admitted to the hospital for suspected isoniazid-induced hepatotoxicity were significantly more likely to be slow acetylators than those who completed their courses of therapy without hepatotoxicity.1,10,51–53

Itraconazole
The FDA has received reports of liver failure and death apparently associated with itraconazole use, including some cases without predisposing risk factors.98

Ketoconazole
Elevated hepatic enzyme levels occur in about 20% of ketoconazole-treated patients, with overt hepatitis in 3%. The typical onset for overt hepatitis is 30–60 days after initiation of ketoconazole therapy. There have been a few deaths attributed to ketoconazole hepatotoxicity.1,54

Lamotrigine
At least 9 cases of lamotrigine-associated hepatotoxicity have been published, including at least 1 case of severe hepatic failure. Most of these cases were complicated by multiple-drug therapy.55

Mercaptopurine
Jaundice associated with cholestasis, hepatic necrosis, and mixed reactions occurs in 6–40% of patients, with the highest prevalence associated with doses ≥2 mg/kg/day. Onset is usually during the first 2 months of therapy.1,56

Methotrexate
Hepatic injury (macrovesicular steatosis, necrosis, and bridging fibrosis) occurs frequently, depends on dose and duration of therapy, and can progress to cirrhosis if the drug is not stopped. Intermittent high doses pose less risk than daily low doses. Cirrhosis is reported in up to 24% of patients receiving long-term daily doses; other contributing factors are alcoholism and pre-existing liver or kidney disease. Hepatic fibrosis is not detected by standard liver function tests and is best detected by biopsy. Biopsy has been recommended at intervals of up to 36 months, after every 1.5 g of methotrexate, if 6 of 12 monthly transaminase levels are elevated, or if the serum albumin level drops below normal. Isolated elevations of transaminase levels do not preclude continued methotrexate therapy.1,14,57–60

Methyldopa
Mild changes in liver function tests occur in up to 35% of patients taking methyldopa, but the prevalence of clinical hepatitis is probably <1%. Most cases occur during the first 3 months of therapy. Hepatitis is more common in women, and most patients have rapid recovery after drug discontinuation. The fatality rate is <10% among patients who develop hepatitis. There is evidence to support a hypersensitivity mechanism in some patients.1,61

(continued)
Minocycline
The long-term use of minocycline for acne or arthritis has resulted in at least 65 reported cases of minocycline-induced hepatitis. Autoimmune hepatitis associated with lupus-like symptoms occurs with a median onset of 1 yr, and an apparent hypersensitivity mechanism is responsible for other cases occurring during the first month of minocycline therapy.62,63

Nevirapine
Severe, life-threatening hepatotoxicity has been reported in patients taking nevirapine for HIV infection and health care workers taking the drug for postexposure prophylaxis. Fatalities have occurred in HIV-infected patients.64

Niacin
Elevations of hepatic enzyme and bilirubin levels occur in 30–50% of patients taking sustained-release niacin in therapeutic doses, with jaundice in 3% of patients taking 3 g/day for >1 yr. Symptomatic hepatic dysfunction occurs frequently and limits the use of the sustained-release product. Immediate-release niacin also is hepatotoxic but to a lesser extent than sustained-release.1,65

Nitrofurantoin
Hepatic damage occurs occasionally, usually during the first month of therapy. Cholestasis is the most common presentation; hepatic necrosis also is reported. Hypersensitivity is the suspected mechanism, and the onset is frequently associated with fever, rash, and eosinophilia. Several late-developing cases of chronic active hepatitis have been reported; almost all are in women and after >6 months of therapy.1,66

Nonsteroidal Anti-inflammatory Drugs
The incidence of clinically apparent hepatic injury from nonsalicylate NSAIDs is estimated to be about 1/10,000 patient–years. The incidence for sulindac may be 5–10 times higher than for the other nonsalicylate NSAIDs. Half of the reactions to sulindac are cholestatic and 25% are hepatocellular. Despite previous reports to the contrary, current data analysis does not support a higher incidence of hepatotoxicity with diclofenac.1,5,67

Octreotide
Most patients on long-term therapy develop cholelithiasis and/or gallbladder sludge; some require cholecystectomy. The prevalence and speed of onset of symptoms might be dosage related.68

Papaverine
Numerous reports of hepatocellular injury and elevated liver enzymes in 27–43% of patients indicate a marked hepatotoxic potential.1,69

Pemoline
Pemoline occasionally causes elevated liver enzymes. The prescribing information for pemoline includes a boxed warning describing 15 cases of acute hepatic failure reported to the FDA between 1975 and 1998; 12 cases resulted in death or required liver transplantation. The earliest onset of hepatic abnormalities in these cases occurred 6 months after the start of pemoline therapy. The few published reports of possible pemoline-induced fulminant hepatic failure do not hold up well under close scrutiny.1,70,71

Penicillamine
Cholestasis resulting from a hypersensitivity reaction occurs occasionally.1,72

Penicillins
Cloxacillin and flucloxacillin are rarely associated with cholestatic hepatitis. The effect is reversible but can persist for months after drug discontinuation.1,73–75 (continued)
Phenothiazines
Most reports of liver damage involve chlorpromazine. The prevalence of hepatic enzyme elevation with this drug has been estimated to be as high as 42%, although 10% is probably more realistic. Similarly, cholestatic jaundice has been projected to occur in up to 5% of patients receiving chlorpromazine, but the actual prevalence is closer to 1%. The onset of cholestasis is generally in the first month of therapy and usually follows a prodrome of GI or influenza-like symptoms. About 70% of affected patients show signs of hypersensitivity, most frequently fever and eosinophilia, and only 5% have rash. Cholestasis usually follows a benign course, and most patients recover 1–2 months after drug discontinuation. A syndrome resembling primary biliary cirrhosis occasionally can occur. Despite the dominance of chlorpromazine in the reported cases, other phenothiazines can produce similar hepatic damage.1,76

Phenytoin
Hepatocellular necrosis is occasionally associated with phenytoin therapy, usually accompanied by other signs of hypersensitivity (eg, eosinophilia, fever, rash, and lymphadenopathy). Onset is usually during the first 6 weeks of therapy. Reported fatality rates have been as high as 30%. Increasing age is an apparent risk factor, with <5% of cases occurring in patients <10 yr old.1,10,77,78

Plicamycin
Laboratory evidence of dose-related hepatotoxicity occurs in virtually all patients. A common lesion is perivenous necrosis.1,79

Progestins
(See Steroids, C-17-α-Alkyl.)

Propoxyphene
A small number of cases of propoxyphene-induced cholestasis have been reported; these are thought to be the result of hypersensitivity.1,80

Propylthiouracil
Increased ALT levels occur in up to 30% of patients. Onset is usually within the first 2 months of therapy, and ALT levels commonly return to normal with dosage reduction. Clinical hepatitis occurs rarely.1,81

Pyrazinamide
Pyrazinamide-induced hepatitis depends on dose and duration of therapy. Daily administration appears to present a greater risk than weekly administration.1,82,83

Riluzole
Elevated hepatic enzymes occur frequently; the prevalence appears to be dosage related.84

Ritonavir
Elevations of serum AST and ALT greater than 3.6 times base line occur in 30% of patients treated with ritonavir.85

Quinidine
Hepatic damage is rare and usually accompanied by other signs of hypersensitivity, especially fever. Most reactions occur in the first month of therapy. The pathology is usually a mixture of hepatocellular necrosis and cholestasis; granulomas also have been reported.1,86

(continued)
Salicylates

Up to 50% of patients taking antiarthritic dosages have laboratory evidence of liver damage. The risk of liver damage is greatest in patients with connective tissue disorders such as SLE or juvenile rheumatoid arthritis. Clinically apparent salicylate-induced hepatitis is uncommon, usually mild, and readily reversible. Hepatotoxicity most often occurs at serum salicylate concentrations >250 mg/L, and only 7% of cases have serum salicylate levels <150 mg/L. Salicylates can cause microvesicular steatosis after intentional overdose.1,5

Steroids, C-17-α-Alkyl

Canalicular cholestasis occurs with a minimal amount of hepatic inflammation. The prevalence appears to be dose related; although laboratory changes are common (occurring in almost all patients taking anabolic steroids), jaundice is not. Jaundice may or may not be preceded by other clinical signs and usually follows 1–6 months of therapy. Peliosis hepatis also has been associated with these compounds, especially the anabolic steroids. Examples are methyltestosterone, norethandrolone, methandrostenolone, fluoxymesterone, oxandrolone, oxymetholone, and stanozolol. C-17-α-ethinyl steroids such as ethinyl estradiol, mestranol, norethindrone, and norethynodrel can produce similar reactions. An association between C-17-α-alkyl steroids and an increase in the prevalence of hepatocellular carcinoma is unclear.1,87

Sulfasalazine

A small number of cases of sulfasalazine-associated hepatic damage, including fatalities, have been reported in children and adults. Hepatic necrosis is apparently part of a generalized hypersensitivity reaction that includes rash, fever, and lymphadenopathy. Onset is usually within the first 4 weeks of therapy.1,88

Sulfonamides, Antibacterial

The sulfonamides currently in use have a lower prevalence of hepatitis than their predecessors, with most reported cases appearing before 1947. Most cases of hepatotoxicity develop during the first 2 weeks of therapy and many are accompanied by other signs of hypersensitivity.1,39,89 (See also Trimethoprim-Sulfamethoxazole.)

Tacrine

In a study of 2446 patients receiving tacrine, 25% had serum ALT levels at least 3 times greater than the upper limit of normal (ULN), 6% had levels at least 10 times greater than the ULN, and 2% had levels at least 25 times greater than the ULN. Most increases were detected in the first week of therapy. Most patients’ ALT levels returned to no more than twice the ULN within 1 month after drug discontinuation, and no patients developed jaundice. Only 33% developed ALT levels more than 3 times the ULN on rechallenge.1,90

Terbinafine

The FDA has received reports of liver failure and death apparently associated with oral terbinafine use, including some cases without predisposing risk factors.98

Tetracycline

Microvesicular steatosis can occur in patients receiving large doses of tetracycline IV, usually >1.5 g/day. Contributing factors include pregnancy, malnutrition, and impaired renal function, but hepatotoxicity has been reported in patients with none of these factors. Onset is usually during the first 10 days of therapy. Most cases of overt liver disease have resulted in death. Oral therapy also can produce signs of hepatotoxicity, although far less frequently.1,39 (continued)
Tolcapone
ALT levels increase to >3 times the upper limit of normal in about 8% of tolcapone-treated patients. These elevations usually develop 6–12 weeks after the start of tolcapone use and can resolve despite continued therapy. At least 3 deaths from fulminant hepatic failure have been reported.91,92

Trimethoprim-Sulfamethoxazole
"Clinically important" liver disease occurs in at least 5.2/100,000 patients (3.8/100,000 with trimethoprim alone). Patients with AIDS are much more susceptible to hepatic injury. The available evidence supports hypersensitivity as the mechanism and cholestasis as the predominant form of injury. Fulminant hepatic failure has been reported.1,39,93

Troleandomycin
From 30 to 50% of patients receiving the drug show some laboratory evidence of abnormal liver function, and up to 4% develop jaundice.1

Valproic Acid
Hepatic enzyme elevations occur in 7–44% of patients, with clinically apparent liver disease in 0.05–1%. Fatal hepatotoxicity occurs most often in children ≤2 yr old on polydrug therapy (1/600) and 3–10 yr old on monotherapy (1/16,000) or polytherapy (1/8300). The diffuse hepatocellular injury, microvesicular steatosis, and hepatic necrosis do not appear to be dose related and most commonly occur in the first 2–3 months of therapy. Serial liver function tests in asymptomatic patients do not predict patients at risk but are commonly recommended because immediate discontinuation might reverse the condition.1,78,94,95

Vitamin A
Hepatomegaly, portal hypertension, and mild increases in liver enzyme levels are common features of chronic vitamin A toxicity. Central vein sclerosis and perisinusoidal fibrosis, which can progress to cirrhosis, have been reported in cases of chronic intoxication. These effects are associated with doses >50,000 IU/day (sometimes with doses as low as 25,000 IU/day). Hepatotoxicity also is possible with acute doses >600,000 IU.1,96

Zafirlukast
Asymptomatic hepatic enzyme elevations occur frequently. At least three cases of severe hepatitis have been reported including one that resulted in liver transplantation.97

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Drug-Induced Nephrotoxicity

This table includes agents that are associated with drug-induced nephrotoxicity but excludes drugs that produce nephrotoxicity as a result of damage to tissues other than the kidney (eg, liver or skeletal muscle). The following abbreviations are used in the table:

<table>
<thead>
<tr>
<th>Abbreviation</th>
<th>Definition</th>
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<tbody>
<tr>
<td>Clcr</td>
<td>Creatinine Clearance</td>
</tr>
<tr>
<td>Crs</td>
<td>Serum Creatinine</td>
</tr>
<tr>
<td>GFR</td>
<td>Glomerular Filtration Rate</td>
</tr>
<tr>
<td>mOsm</td>
<td>Milliosmole</td>
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</table>

**Acetaminophen**

Tubular necrosis has been reported, usually in association with hepatotoxicity from acute overdose. Whether nephrotoxicity is a direct effect of acetaminophen or the result of liver damage is the subject of controversy. There is a possible association between prolonged acetaminophen use (1–5 kg cumulative dosage) and the development of chronic renal failure. There is insufficient evidence to associate acetaminophen use alone with analgesic nephropathy. (See Analgesics.)

1-8
ACE Inhibitors
ACE inhibitors are frequently associated with proteinuria and renal insufficiency. The prevalence of proteinuria in captopril-treated patients is estimated at 1%. The risks of renal insufficiency are greater with long-acting ACE inhibitors such as enalapril or lisinopril than with captopril. Immune complex glomerulopathy is a major contributor to ACE inhibitor nephrotoxicity. Hyponatremia, diuretic therapy (and other causes of hypovolemia), pre-existing renal impairment, CHF, and diabetes mellitus contribute to an increased risk of nephrotoxicity. Recovery of renal function usually follows ACE inhibitor discontinuation.1,2,9–12

Acetazolamide
Glaucoma therapy with acetazolamide is associated with a 10-fold increase in the risk of renal stone formation. Calcium phosphate and calcium oxalate stones have been identified.13,14

Acyclovir
Acyclovir is concentrated in the urine, and its precipitation in the collecting tubules with subsequent obstructive nephropathy frequently accompanies high-dose (500 mg/m²) IV use; oral therapy is apparently free from this problem. Aggressive hydration (100–150 mL urine/hr) and administration over 1–2 hr should minimize the risk. Normal renal function usually returns within 6 weeks after drug withdrawal.2,9,15,16

Aldesleukin
Almost all patients receiving aldesleukin develop acute renal impairment marked by decreased Clcr, oliguria or anuria, and fluid retention. Most patients recover within 1 week after drug discontinuation, but some require ≥1 month.17

Allopurinol
Glomerulonephritis, interstitial nephritis, and interstitial fibrosis occur rarely in allopurinol-treated patients. Most cases are associated with generalized hypersensitivity reactions to allopurinol (allopurinol hypersensitivity syndrome).18,19

Aminoglycosides
Proximal tubular necrosis occurs in up to 30% of patients treated with aminoglycosides for >7 days. Because of slow clearance of these drugs from renal tissue, they still can be present in high concentrations in the kidney after serum levels are undetectable, but there does not appear to be a good correlation between renal tissue concentrations of individual aminoglycosides and their nephrotoxic potential. Aminoglycoside-induced acute renal failure is usually nonoliguric, which can delay its recognition. It is often first detected as an asymptomatic increase in Cr. Detectable changes in GFR usually occur at least 5 days after initiation of therapy and can progress after drug discontinuation. Aminoglycoside-induced renal damage is related to total dosage and duration of treatment. Administration of single daily doses does not markedly affect the frequency of nephrotoxicity. Recovery of some to all lost renal function can occur over several weeks after drug discontinuation. Monitoring of aminoglycoside plasma levels and serial renal function tests might be of value in recognizing nephrotoxicity. Neomycin has the greatest and streptomycin the least nephrotoxic potential of the aminoglycosides. All other currently marketed aminoglycosides have intermediate nephrotoxic potentials. Concomitant therapy with other nephrotoxic drugs should be avoided.1,10,20–23
Amphotericin B

Mild or moderate renal impairment occurs in 50% of patients treated with conventional amphotericin B, with severe renal impairment in 8%. The drug causes a reduction in renal plasma flow as well as glomerular and tubular damage. Most patients experience a rapid decline in GFR, which often stabilizes at 20–60% of normal and might not return to normal until several months after drug discontinuation. Distal tubular damage can lead to loss of concentrating ability, renal tubular acidosis, and electrolyte disturbances (most commonly hypokalemia but also hyponatremia and hypomagnesemia). These effects appear to be dosage related, and many patients respond favorably to temporary drug discontinuation or reduction in dosage. The prevalence of nephrotoxicity increases as the cumulative dose increases. Some investigators suggest that the total dosage of conventional amphotericin B should be kept below 3–5 g. Nephrotoxicity is increased by the co-administration of other nephrotoxic drugs, especially cyclosporine. Sodium loading (eg, 1 L normal saline IV daily) reduces the frequency and severity of amphotericin B–induced nephrotoxicity. Newer dosage forms (eg, liposomal amphotericin B) appear to be less nephrotoxic.1,2,9,10,24–26

Analgesics

Analgesic nephropathy is a syndrome of papillary necrosis, interstitial nephritis, and progressive renal medullary impairment that occurs in persons with long-term consumption of large quantities of oral analgesic products, especially combination products. Most reported patients are 30–70 yr old, and women greatly outnumber men. The syndrome is characterized by proteinuria, reduced renal concentrating ability, and RBCs and WBCs in the urine. Analgesic nephropathy has been clearly associated with products containing phenacetin, but the removal of phenacetin from non-prescription analgesic products has not been consistently associated with a decline in analgesic nephropathy mortality. Acetaminophen or salicylates taken alone or in combination do not seem to cause analgesic nephropathy. Historically, this syndrome has been responsible for a large percentage of chronic renal failure deaths, with considerable variation in prevalence among nations (high in Australia and Germany, low in the United States), apparently reflecting analgesic abuse patterns. Mild cases are reversible, but severe cases can continue to deteriorate after the discontinuation of analgesics. The prevalence of urinary tract cancer appears higher than normal among chronic analgesic abusers.1,2,5–8

Azacitidine

Proximal and distal tubular dysfunction, polyuria, glucosuria, and decreases in serum bicarbonate occur occasionally during azacitidine therapy.27

Carboplatin

Although apparently less nephrotoxic than cisplatin, carboplatin therapy is frequently associated with reductions in GFR and increased electrolyte losses (especially calcium and magnesium). Patients with pre-existing renal impairment and those who receive inadequate hydration during drug administration are at greatest risk.28

Cephalosporins

The cephalosporin (and cephamycin) antibiotics are capable of producing rare interstitial nephritis similar to the penicillins. Increases in BUN and Cr occur occasionally. The nephrotoxicity of the newer cephalosporins is minimal compared with older drugs such as cephalothin.29–31

Cidofovir

Proteinuria occurs frequently during cidofovir therapy. Probenecid decreases the prevalence and magnitude of proteinuria and must be given with cidofovir.32

(continued)
Cisplatin
Dosage-related proximal tubular impairment is the major limiting factor in cisplatin therapy and can occur in 50–75% of patients. Clcr is typically reduced to 60–80% of baseline with repeated courses of therapy. The greatest damage occurs in the first month of therapy, and it appears to be more likely when the drug is administered repetitively at close intervals. Forced hydration and mannitol diuresis can reduce renal toxicity, at least for the first cycle of therapy. Magnesium and calcium losses are common manifestations of cisplatin-induced nephrotoxicity. Cisplatin-induced renal effects can be detected as long as 6 months after the end of therapy.1,19,28,33

Contrast Media, Radiopaque
Increased Crs occurs frequently in patients receiving iodine-containing contrast media. In unselected patients, the prevalence of Crs >0.5 mg/dL or >50% above pretreatment is 2–7%. Renal lesions include medullary necrosis and proximal tubular vacuolation and necrosis as well as the deposition of urate and oxalate crystals. The most common pattern is acute oliguric renal failure developing within 24 hr after the administration of the contrast agent and lasting 2–5 days; nonoliguric renal failure also has been reported. Most patients recover fully, but permanent renal impairment has been reported. Crs usually peaks 3–5 days after exposure and returns to baseline in 10–14 days. Patients with pre-existing renal impairment are at much greater risk and constitute 60% of those experiencing nephrotoxicity. Vigorous hydration before, during, and after drug administration with hypotonic saline reduces the risk of nephrotoxicity, but mannitol or furosemide diuresis can increase the risk. High-osmolality ionic contrast media might be more nephrotoxic than low-osmolality ionic contrast media. Nonionic contrast agents might be less nephrotoxic than ionic agents.1,2,10,34–36

Cyclosporine
Dose-related nephrotoxicity occurs in 30–50% of cyclosporine-treated patients and frequently limits the usefulness of the drug. Reduction in dosage usually reduces the renal toxicity. The drug produces decreased GFR, impaired tubular function, interstitial nephritis, hypertension, fluid retention, and hyperkalemia. Cyclosporine causes vasoconstriction in preglomerular arterioles, which can lead to chronic arteriopathy and tubular atrophy if the dosage is not reduced. Cyclosporine nephrotoxicity is usually reversible during the first 6 months of therapy, but the risk of permanent renal impairment increases with time. Calcium-channel blockers appear to reduce the prevalence of cyclosporine-induced nephrotoxicity in renal transplant patients.1,2,9,20,37,38

Demeclocycline
This drug can produce nephrogenic diabetes insipidus, which is usually, but not always, dosage related. For this reason, it has been used in the management of the syndrome of inappropriate antidiuretic hormone secretion.20,39 (See also Tetracyclines.)

Diuretics, Thiazide
Occasional cases of interstitial nephritis have been reported, which might be the result of hypersensitivity reactions. Long-term use of diuretics might increase the risk of renal cell carcinoma, especially in women.2,40

Fluoroquinolones
Acute interstitial nephritis is associated with fluoroquinolone antibiotics; a hypersensitivity mechanism is suspected but remains to be confirmed. Most reported patients are >50 yr old.41

Foscarnet
Acute tubular necrosis occurs frequently with foscarnet. Crs increased during 35 of 56 courses of therapy in one retrospective study. Hydration with normal saline appears to markedly decrease the severity and frequency of nephrotoxicity.42

(continued)
Furosemide

Use of high-dose furosemide (5–10 mg/kg/day) in adults with refractory CHF is associated with a 40% decrease in Clcr. Nephrocalcinosis and nephrolithiasis occur in up to 64% of low-birthweight infants treated with furosemide. These effects usually resolve after drug discontinuation.43,44

Gallium Nitrate

Nephrotoxicity is the most frequent adverse effect of gallium, and elevations in BUN and Cr can occur after only 1 dose. At least 1 death has been associated with gallium-induced nephrotoxicity.45,46

Gold Salts

A lesion resembling membranous glomerulonephritis with proteinuria can occur in 3–10% of patients receiving parenteral gold therapy. Microhematuria and nephrotic syndrome are less frequent. One-half of the cases of proteinuria develop in the first 6 months of therapy. Occasionally, acute tubular necrosis and interstitial nephritis are reported. Although recovery can take up to 18 months, permanent renal impairment after drug withdrawal is uncommon. There is evidence for immune and direct toxic mechanisms for gold nephrotoxicity. Oral auranofin appears to be less nephrotoxic than parenteral gold products.1,2,9,10,47,48

Ifosamide

Reversible, subclinical nephrotoxicity occurs in almost all ifosamide-treated patients, with clinically important nephrotoxicity in many. Renal damage might correlate with total dosage, and cumulative doses >60 g/m² should be avoided, especially in children <5 yr old. Fanconi syndrome–like symptoms including renal loss of glucose, electrolytes, and small proteins occur in 4%.28,49,50

Immune Globulin

Intravenous administration of immune globulin can produce reversible acute renal failure after the first or repeated exposures. The origin of the acute renal failure is not the immune globulin but rather the large amount of sucrose used in some immune globulin products to reduce the formation of immunoglobulin aggregates. The damage is probably due to the delivery of a high-osmotic solute load to the kidneys. Maltose- and dextrose-stabilized products might have the same capacity, but there are no case reports in the literature.51

Indinavir

Crystalluria occurs in most indinavir-treated patients; many develop nephrolithiasis, back pain, or flank pain. The crystals contain indinavir; good hydration (2–3 L fluid/day) reduces their formation.16,52,53

Lithium

Lithium frequently produces nephrogenic diabetes insipidus, which is, at least in part, dosage related. This typically mild effect is usually reversible with drug withdrawal. Long-term therapy (10–15 yr) is associated with an increased prevalence of reduced Clcr and renal concentrating ability that are frequently not reversible, despite withdrawal of lithium. Interstitial nephritis and nephrotic syndrome also have been reported.1,2,9,10,54-56

Mannitol

High doses (>200 g/day or >400 g/2 days) are associated with acute oliguric renal failure. Although low doses act as renal vasodilators, high doses produce renal vasoconstriction. Keeping the osmolar gap to no more than 55 mOsm/kg should minimize the risk. Acute renal failure might require 7–10 days for recovery; dialysis shortens the recovery period to 1–2 days.57,58

(continued)
Methotrexate

This drug is directly toxic to the kidney in large doses, causing acute tubular necrosis. Acute renal impairment occurs in 30–50% of patients treated with high-dose methotrexate and leucovorin rescue. Most cases are reversible within 3 weeks. Methotrexate is eliminated primarily through the kidney, and its nephrotoxicity compounds itself by causing the serum level of the drug to rise. About 20% of deaths associated with methotrexate therapy are caused by acute renal failure. The drug and its metabolites precipitate in the distal tubule. Close monitoring of methotrexate serum concentrations and adjustment of dosage might minimize the risk of nephrotoxicity, as would vigorous hydration and alkalinization during drug administration.2,16,59,60

Methoxyflurane

Nephrogenic diabetes insipidus, proximal tubular damage, and interstitial nephritis are reported. The nephrotoxicity of methoxyflurane appears to be dose related and might be caused by increased circulating fluoride ion concentrations. Fluoride causes distal tubular dysfunction by inhibiting sodium and chloride transport in the ascending loop of Henle and reducing the response to antidiuretic hormone. Urinary oxalate crystallization also has been reported after methoxyflurane anesthesia.20,61,62

Mitomycin

Tubular necrosis occurs most frequently with daily therapy but is also reported with the intermittent therapy now recommended. Nephrotoxicity appears to be related to the total dosage administered, with the risk of renal impairment rising when the total dosage exceeds 30 mg/m². Onset can be delayed for many months.63,64

Nitrosoureas

The nitrosoureas can produce insidious nephrotoxicity in patients on long-term therapy. Lomustine seems to have the greatest nephrotoxic potential. Some cases of permanent renal function impairment have been reported.65

Nonsteroidal Anti-inflammatory Drugs

NSAIDs, including COX-2 inhibitors, can reduce Clcr and produce renal insufficiency as a result of renal circulatory changes caused by inhibition of prostaglandin synthesis. These effects tend to be relatively minor and usually reversible. The prevalence is usually low (0.5–1% of patients), but some patients are at increased risk; predisposing factors are advanced age, pre-existing renal impairment, and states of renal hypoperfusion (eg, sodium depletion, hypotension, diuretic use, hepatic cirrhosis, and CHF). Reversible acute interstitial nephritis and necrosis occur occasionally. It is not possible at this time to accurately categorize the prevalence associated with each NSAID. Fenoprofen is the NSAID most commonly associated with interstitial nephritis and nephrotic syndrome.1,2,6–10,20,66

Omeprazole

Interstitial nephritis occurs rarely during omeprazole therapy. At least 13 cases have been published, 10 with positive biopsies; 5 cases were rechallenged with recurrence of interstitial nephritis in all. Onset is usually after 2 weeks to 6 months of omeprazole therapy.67

Penicillamine

Slight to moderate proteinuria occurs in 7–30% of patients on long-term (≥6 months) therapy with penicillamine for rheumatoid arthritis. Most cases develop in the first year. Proteinuria is usually benign and slowly reversible over 6–12 months, but nephrotic syndrome is occasionally encountered. The lesions appear to be perimembranous glomerulonephritis resulting from the deposition of antigen–antibody complexes on the renal basement membrane.1,2,10,20,68

(continued)
Penicillins
Interstitial nephritis has been reported with most penicillins. Methicillin was by far the most frequently implicated penicillin (frequency 10–16%); the reason for its dominance is unknown. Penicillin-induced interstitial nephritis is an immune reaction that most commonly occurs during a long course of therapy. The reaction is usually accompanied by other signs of hypersensitivity such as fever, rash, and eosinophilia; hematuria also can occur. The reduction of renal function might not be oliguric; so urine volume is not a reliable parameter to monitor. Recovery usually occurs within weeks to months after drug discontinuation.1,2,20,69

Pentamidine
Prospective trials of IV pentamidine for the treatment of Pneumocystis carinii pneumonia show nephrotoxicity in 4–66% of patients. Onset is usually 8–12 days after the start of therapy.70

Plicamycin
High doses (50 µg/kg/day) produced renal impairment in 40% of patients, including some who died of acute renal failure. Nephrotoxicity is far less likely at the 25–30 µg/kg/day (or lower) dosage used most often.71

Polymyxins
Adverse reactions involving the kidney occur in about 20% of patients receiving colistimethate parenterally. Tubular necrosis is the most frequently described lesion, but interstitial nephritis is also reported. High dosage, long duration of therapy, and renal impairment are predisposing factors. Polymyxin-induced renal damage is usually reversible, but some patients continue to deteriorate after drug withdrawal.72

Rifampin
There are at least 49 published cases of rifampin-induced acute renal failure. Acute tubular necrosis is the most common lesion. This appears to be a hypersensitivity reaction and most often occurs with intermittent or interrupted dosage regimens but has accompanied continuous therapy.73

Streptozocin
Nephrotoxicity is the most common dosage-limiting side effect. The prevalence increases with prolonged administration until virtually all patients demonstrate renal impairment. Dosages <1.5 g/m²/week are less toxic. The damage is glomerular and tubular. The drug should be discontinued as soon as renal damage is detected.28,71

Sulfonamides, Antibacterial
Early sulfonamides were poorly soluble, and urinary crystallization was a common problem. Crystalluria occurs in 8–29% of sulfadiazine-treated patients; symptomatic renal impairment resulting largely from nephrolithiasis occurs in 2–8%. Crystallization occurs in <0.3% of patients receiving the more soluble sulfonamides and adequate hydration. Interstitial nephritis, glomerulonephritis, and tubular necrosis are reported rarely. These reactions are probably allergic in origin.7,9,16,74,75

Tacrolimus
Acute nephrotoxicity occurs with a prevalence similar to that of cyclosporine. Progressive nephrotoxicity is reported with long-term (>1 yr) therapy. The risk of nephrotoxicity can be greatly limited by keeping the tacrolimus whole blood concentration <20 µg/L.97,76,77

(continued)
Tetracyclines

Fanconi syndrome, characterized by tubular damage with proteinuria, glycosuria, aminoaciduria, and electrolyte disturbances, was associated with the use of outdated tetracycline products. Because of changes in the manufacturing process, this syndrome is now unlikely to occur. The antibacterial effects of tetracyclines can contribute to azotemia in patients with pre-existing renal impairment.78 (See also Demeclocycline.)

Topiramate

Nephrolithiasis occurs in 1.5% of topiramate-treated patients.79

Triamterene

Triamterene therapy is associated with an increase in urinary sediment, and the drug can be incorporated into existing renal calculi. One report suggests that 1/1500 users of the drug will develop triamterene-associated calculi during the course of 1 yr. As a precaution, the drug probably should not be used in patients with a history of renal calculi. Triamterene also might be associated with the development of interstitial nephritis.16,80,81

Vancomycin

Nephrotoxicity from vancomycin was commonly reported early in its history. Currently, the prevalence of vancomycin-induced renal impairment (usually mild) is 5–17%. It is usually reversible after discontinuation of the drug. Concomitant administration of aminoglycosides results in at least additive nephrotoxicity.2,82–85

REFERENCES


### Drug-Induced Oculotoxicity

Occasionally, nonspecific blurred vision occurs with almost all drugs. The agents in this table are associated with a specific pattern of drug-induced oculotoxicity when administered systemically.

#### Allopurinol

Despite the discovery of allopurinol in cataractous lenses taken from patients on long-term (>2 yr) therapy, there is no clinical evidence for an increased risk of cataracts in allopurinol-treated patients.1–3
Amantadine
At least 9 cases of diffuse, white, subendothelial corneal opacities have been reported. These opacities usually resolved within a few weeks after amantadine discontinuation.4

Amiodarone
Most patients treated with amiodarone develop bilateral corneal microdeposits (75% after 1 yr of therapy). Visual symptoms occur in 6–14%. Halo vision at night is most commonly reported, but patients also might complain of photophobia and blurred vision. The deposits are apparently dose related and reversible, disappearing 3–7 months after drug discontinuation. Minute lens opacities occurred in 7 of 14 amiodarone-treated patients in one study.5–7

Anticholinergic Agents
Blurring of vision can result from paralysis of accommodation (cycloplegia). These drugs also dilate the pupil (mydriasis), which can produce photophobia and precipitate narrow-angle glaucoma. With systemic administration, large doses are usually required to produce mydriasis, which is most commonly associated with potent anticholinergics such as atropine, scopolamine, or benztropine. Patients being treated for narrow-angle glaucoma can usually tolerate systemic anticholinergic therapy but nevertheless should avoid these drugs unless absolutely necessary. Patients with open-angle glaucoma, particularly if treated, can receive anticholinergic medications without much risk. Patients receiving nebulized ipratropium by face mask are at risk for developing increased intraocular pressure and precipitation of narrow-angle glaucoma, probably from the drug escaping from beneath ill-fitting masks and directly affecting the eyes. All of the ocular effects of anticholinergics are dose related and reversible.5,8–10

Anticonvulsants
Diplopia and nystagmus occur frequently. Blurred vision can be caused by mydriasis (phenytoin) or cycloplegia (carbamazepine). All of these effects are dose related.11

Antidepressants, Heterocyclic
These drugs have anticholinergic properties and can precipitate narrow-angle glaucoma and cycloplegia at usual doses. (See Anticholinergic Agents.) There is a 10–30% prevalence of blurred vision resulting from cycloplegia, but it is rarely troublesome and is reversible with drug discontinuation. Blurred vision usually resolves despite continued antidepressant use as the eye becomes tolerant to the drug’s effects. SSRIs do not seem to produce any important ocular effects.5,12,13

Antihistamine Drugs (H1-Blockers)
With the exception of loratadine and fexofenadine, these drugs have some anticholinergic properties and can precipitate narrow-angle glaucoma and cycloplegia. (See Anticholinergic Agents.) These effects are minor and reversible with drug discontinuation. Antihistamines (most notably diphenhydramine) can reduce night vision.5,8,14

β-Adrenergic Blocking Agents
A reduction in tear production occurs, which can produce a hot, dry, gritty sensation in the eyes. This is rapidly reversible with drug discontinuation.

Bromocriptine
Myopia is a frequent complication of long-term bromocriptine therapy and often goes unappreciated until the patient complains of blurred vision. The cause is not fully determined but might be due to lens swelling. Myopia is reversible within 1–2 weeks after drug discontinuation.5,15,16

Busulfan
Long-term therapy (usually ≥1 yr) with busulfan is associated with the development of posterior subcapsular cataracts in about 10% of patients.5,17–19

(continued)
Chloramphenicol

Optic neuritis, papilledema, and visual field defects are occasionally reported. These effects can occur after weeks or years of therapy but are most common after several months of chloramphenicol use. Most cases are reported in children with cystic fibrosis, but the association with this disorder is unclear and might only reflect the types of patients who received long-term chloramphenicol therapy. Permanent visual impairment and recovery are reported after drug discontinuation. There are anecdotal reports that large doses of vitamins B6 and B12 have beneficial effects on these adverse ocular effects.5,19–22

Chloroquine

The oculotoxicity of chloroquine limits its usefulness; two general types of ocular change occur: corneal deposits and retinopathy. About 50% of patients demonstrate corneal deposits, less than one-half of whom have visual impairment resulting from these deposits. Opacities present as punctate or whirling patterns. They can appear after 2 months and usually do not interfere with vision. They are usually reversible in 6–8 weeks after drug discontinuation. Early changes in the retina (deposition of pigment in the macula) are usually asymptomatic and reversible. More advanced damage includes hyperpigmentation of the macula surrounded by a depigmented ring and hyperpigmented retina (“bull’s-eye” retinopathy). Patients complain of reading difficulty, blurred vision, visual field defects, and photophobia; some also report defective color vision and light flashes. The prevalence ranges from 3% to 45% in various reports. The drug should be discontinued if these symptoms develop. Patients receiving long-term therapy with chloroquine 3 mg/kg/day should have ophthalmologic examinations at least every 6 months initially and then annually if their vision remains stable. Those receiving >3 mg/kg/day should be examined every 6 months.27 Daily dosage seems to be more important than the total dosage or duration of therapy for the development of retinopathy; limiting the daily dosage to 4 mg/kg up to a maximum of 250 mg in adults minimizes the risk. The prognosis of chloroquine-induced retinopathy is uncertain. Weekly use of chloroquine for malarial prophylaxis does not seem to cause retinopathy.15,23–27

Cidofovir

Anterior uveitis occurs in about one-third of AIDS patients receiving the drug intravenously for treatment of cytomegalovirus retinitis. The onset is usually after 4–5 days of treatment. Uveitis usually responds to topical cycloplegics and corticosteroids and does not require discontinuation of cidofovir.28

Cisplatin

Blurred vision and altered color perception are frequently associated with high-dose cisplatin. Blurred vision gradually improves after drug discontinuation, although altered color vision can persist. Pigmentary retinopathy is also reported.2,18,19

Clomiphene

Visual disturbances, most commonly blurred vision, occur frequently with clomiphene. These disturbances usually disappear after the drug is withdrawn, but one report of three patients describes prolonged afterimages, shimmering of the peripheral visual field, and photophobia.29

Contraceptives, Oral

A variety of retinal vascular disorders have been attributed to oral contraceptives, but the association remains unproved. It is purported that some oral contraceptive users cannot tolerate contact lenses, possibly because of ocular edema or dryness; however, a prospective study failed to show any differences in lens tolerance between oral contraceptive users and nonusers.30,31

(continued)
Corticosteroids

These drugs can produce a variety of ocular disorders with long-term therapy, most notably glaucoma and cataracts. Corticosteroid-induced increases in intraocular pressure occur in approximately 30% of long-term users and appear to be dose related. Glaucoma can persist for several months after drug discontinuation. Corticosteroid-induced cataracts (usually posterior subcapsular) are found in 10–40% of patients on long-term, systemic therapy and are correlated with total dosage and duration of therapy. Outcome is variable, ranging from improvement despite continued therapy to rare loss of sight. Most patients have no vision impairment. Although they most commonly occur with large oral doses, increased intraocular pressure and cataracts are reported in patients receiving corticosteroids by the topical ophthalmic, inhalation, and intranasal routes. Children develop cataracts more frequently than adults; Hispanics might be affected more often than blacks or non-Hispanic whites.2,3,5,18,19,32–35

Cyclophosphamide

Keratoconjunctivitis occurs in up to 50%. One report showed a 17% prevalence of transient reversible blurred vision during high-dose cyclophosphamide therapy. Recovery took from 1 hr to 14 days.3,18,19,96

Cyclosporine

Retinopathy occurs frequently with cyclosporine and severe visual disturbances, including cortical blindness, occur occasionally. Oculotoxicity appears to be dose related and resolves after drug discontinuation.19,37

Cytarabine

Keratoconjunctivitis, corneal damage, ocular pain, and photophobia are frequent, dose-related side effects of cytarabine. These symptoms usually resolve 1–2 weeks after drug discontinuation. Pretreatment with corticosteroid eye drops can be beneficial but should be used with caution in patients with corneal damage.3,18,19,38,39

Deferoxamine

Oculotoxicity, including blurred vision, impaired color vision, night blindness, and retinal deposits, occurs in 4–11% of patients receiving deferoxamine for chronic iron overload. These effects appear to be dose related and might be caused by the chelation of trace minerals.40–43

Digitalis Glycosides

The most unique ocular effect is the frosted or snowy appearance of objects or colored halos around them. These effects are most noticeable in bright light. Color vision might be affected such that objects appear yellow (green or other colors are reported, but far less frequently). With digoxin, color changes usually occur when the plasma level exceeds 1.5 µg/L. Digitalis glycosides also are reported to produce photophobia, blurred vision, central scotomas, and flickering or light flashes before the eyes. Reversible ocular side effects occur in up to 25% of patients with digitalis intoxication.5,44,45

Disopyramide

The anticholinergic effects of disopyramide frequently produce blurred vision.46

Disulfiram

A few cases of retrobulbar neuritis have occurred, manifested by a dramatic decline in visual acuity and impairment of color vision. In most patients, vision returns to normal after drug discontinuation.5,47

Doxorubicin

This drug stimulates excessive lacrimation shortly after administration in about 25% of patients. Conjunctivitis also has been reported.18,19,48

(continued)
Ethambutol

Retrobulbar neuritis is the primary ocular complication. Symptoms include blurred vision, scotoma, and reduction of the visual field. Color vision defects also occur, usually presenting as a reduction in green perception. Retrobulbar neuritis is dose related, occurring most frequently with dosages ≥25 mg/kg/day. Its onset is usually after 3–6 months of therapy, and it is slowly reversible after drug discontinuation. Dosages ≤15 mg/kg/day appear relatively free of ocular side effects.5

Fenoldopam

Treatment of hypertensive emergencies with fenoldopam results in dose-dependent, mild increases in intraocular pressure during the infusion. Increases in intraocular pressure occur in patients with and without ocular hypertension. The importance of these findings is not established.49,50

Fluorouracil

Adverse ocular effects occur in 25–50% of patients receiving fluorouracil systemically. Blurred vision, ocular irritation and pain, conjunctivitis, keratitis, and excessive lacrimation occur frequently. These effects resolve in 1–2 weeks after drug discontinuation. Some patients can develop erosion of the eyelid margin (cicatricial ectropion) or potentially irreversible fibrosis of the tear duct (dacryostenosis) with prolonged therapy.3,5,18,19,51,52

Gold Salts

Parenteral gold can produce microscopic crystalline deposits in the cornea, most commonly in the superficial layers. These deposits are dose related and rarely occur until the total dosage of parenteral gold exceeds 1 g. The deposits slowly resolve after drug discontinuation, do not appear to affect vision, and are not a reason to stop gold therapy. Auranofin does not seem to produce these ocular effects.5,53,54

Hydroxychloroquine

This drug can produce the same spectrum of ocular toxicity as chloroquine. (See Chloroquine.) Corneal deposits occur only with high daily doses. Limiting the daily dosage to 6.5 mg/kg up to a maximum of 400 mg in adults minimizes the risk of retinopathy.5,24,25,27,55,56

Interferon Alfa

Although the prevalence cannot be accurately determined, retinal vascular complications have been reported. Onset is usually after 2–3 months of treatment. These effects appear to be reversible after drug discontinuation.57

Iodine, Radioactive (131I)

Ophthalmopathy, including diplopia and changes in visual acuity, occurred or worsened in 15% of patients with Graves’ hyperthyroidism treated with 131I after a 3–4 month course of methimazole. Patients treated with a combination of 131I and prednisone or continued methimazole did not show any increased ophthalmopathy. All changes occurred during the first 6 months after 131I treatment. Ophthalmic changes persisted for 2–3 months in 65% of those affected, longer in the other 35%.58

Isoniazid

Optic neuritis occurs occasionally, most commonly in malnourished or alcoholic patients, and often manifests itself as impaired red–green perception. It responds to pyridoxine therapy.5

Methotrexate

Adverse ocular effects associated with systemic methotrexate occur in up to 25% and include conjunctivitis, increased or decreased lacrimation, photophobia, and eye pain. Onset is during the first week of therapy, and resolution usually occurs 1–2 weeks after drug discontinuation.13,18,19
**Minocycline**
Dark-blue discoloration of the sclera has been reported. Although the prevalence cannot be accurately estimated, the growing use of minocycline as an antiarthritic drug should increase the number of cases. It is not known if the discoloration is reversible.59

**Muromonab-CD3**
Conjunctivitis and photophobia occur frequently.60

**Oprelvekin**
Transient blurred vision and conjunctival injection occur frequently during oprelvekin therapy. Papilledema occurs in 1.5%.

**Oxygen**
Retrolental fibroplasia is an important complication of oxygen therapy in neonates, in particular premature or other low-birthweight neonates. The risk of retrolental fibroplasia in these patients increases whenever the concentration of inspired oxygen exceeds normal.61–63

**Paclitaxel**
Scintillating scotomas or photopsia occur frequently during paclitaxel infusions. The onset of these short-lived effects is usually during the last hour of the infusion. They do not always recur during subsequent infusions.19,64,65

**Pamidronate**
Reversible anterior uveitis and conjunctivitis are occasional complications of pamidronate therapy. Onset is usually 24–48 hr after IV infusion.28,66

**Pentostatin**
Conjunctivitis and keratitis frequently occur during pentostatin therapy. Whereas conjunctivitis is usually mild, keratitis can be severe.3

**Phenothiazines**
Lesions of the lens, cornea, and retina are the most important features of phenothiazine-induced oculotoxicity. White to yellow-brown deposits in the lens most frequently occur with long-term, high total-dose (>600 g) chlorpromazine therapy. Epithelial keratopathy, possibly resulting from a photosensitivity reaction, can occur after only a few months of high-dose therapy. It is characterized by diffuse opacification of the corneal epithelium. The consistent use of sunglasses can reduce the risk of keratopathy. Lens and corneal deposits usually do not interfere with vision, and all of these effects might be slowly reversible. Thioldiamide is most noted for producing pigmentary retinopathy. As with most phenothiazine-induced ocular effects, pigmentary retinopathy is dose related. Patients might complain of blurred vision, decreased night vision, brown discoloration of vision, and central scotoma. Vision might improve if the drug is withdrawn soon enough; however, some cases continue to deteriorate despite drug discontinuation. Other phenothiazines can cause pigmentary retinopathy, but the supporting data are limited to case reports. Phenothiazines (especially thioridazine) have anticholinergic effects and might precipitate narrow-angle glaucoma. Corneal edema is a rare, but dangerous, complication of phenothiazine use, requiring immediate discontinuation of therapy.12,67,68

**Psoralens**
The combination of psoralens and long-wave ultraviolet light (PUVA therapy) radiation is associated with the development of conjunctivitis, photophobia, and other signs of ocular irritation. The use of UVA protective lenses during therapy greatly reduces the prevalence. An experimentally demonstrated connection between PUVA therapy and cataracts has not been confirmed clinically.5,69,70

(continued)
Quinine
Loss of visual acuity and reduction of the visual field to the point of blindness can occur with quinine therapy or (especially) overdose. Other reported ocular effects are impaired color vision and night blindness. These effects are usually reversible, but permanent constriction of the visual field and blindness are reported. The ocular effects of quinine might be the result of changes in the retinal vasculature.5,71,72

Retinoids
Blepharoconjunctivitis occurs in >50% of patients receiving isotretinoin. This painful condition appears to be dose related, and its onset is usually during the first 2 months of therapy. Dry eyes can occur with or without blepharoconjunctivitis. Corneal opacities, which clear in 6–7 weeks after drug discontinuation, also are reported. Similar effects were reported with etretinate. Other effects associated with retinoid therapy are papilledema and night blindness. Resolution usually occurs within a week after retinoid discontinuation.5,19,73–76

Rifabutin
Uveitis occurs frequently during rifabutin treatment and prophylaxis of Mycobacterium avium complex infection in AIDS patients. Its onset is variable (2 weeks to 7 months after starting treatment). Uveitis can be unilateral or bilateral and responds to topical corticosteroid therapy.28,77,78

Rifampin
Exudative conjunctivitis, ocular pain, and orange staining of tears (and consequent staining of soft contact lenses) are occasionally reported with rifampin. These effects are rapidly reversible when the drug is withdrawn.79–81

Sympathomimetic Agents
These drugs can dilate the pupil and precipitate narrow-angle glaucoma. Sympathomimetics with marked α-adrenergic activity (eg, ephedrine, phenylpropanolamine, tetrahydrozoline) should be avoided. The risk of this reaction is slight unless large doses are taken orally or the drugs are applied topically.9

Tamoxifen
Fine, refractile retinal opacities and retinopathy occur frequently; corneal opacities also are reported. The prevalence of retinopathy has been 1.5–11.8% in prospective studies. Although these lesions can occur with any dosage, they occur most often with daily dosages >180 mg or cumulative dosages >100 g. They can result in reduced visual acuity and are slowly reversible after drug discontinuation.3,5,18,19,82

Vigabatrin
Visual field loss (concentric or bilateral nasal) occurs to some degree in 29–40% of vigabatrin-treated patients and is severe in 9%. Males are more susceptible than females.83,84

Vincabatrin
Various ocular disorders occur frequently. Most (ptosis, blurred vision, night blindness) are thought to be the result of cranial nerve impairment. Posis occurs in up to 50% of vincristine-treated patients. Time to onset ranges widely (2–44 weeks) as does resolution after drug discontinuation (2–24 weeks). Vincristine might be more oculotoxic than vinblastine.3,5,18,19,85

REFERENCES
Drug-induced ototoxicity can affect hearing (auditory or cochlear function), balance (vestibular function), or both, depending on the drug. Drugs of almost every class have been reported to produce tinnitus, as have placebos. The agents in this table are associated with measurable changes in hearing or vestibular defect when administered systemically.

**Aminoglycosides**

Aminoglycoside antibiotics can cause cochlear and vestibular toxicities. Cochlear toxicity presents as progressive hearing loss, starting with the highest tones and advancing to lower tones. Thus, considerable damage can occur before the patient is cognizant of it. Vestibular damage presents as dizziness, vertigo, or ataxia. Both forms of ototoxicity are usually bilateral and potentially reversible, but permanent damage is common and can progress after aminoglycoside discontinuation. Estimates of the prevalence of aminoglycoside-induced ototoxicity vary widely depending on the criteria applied. Clinically detectable ototoxicity probably occurs in as many as 5% of patients, with a much higher percentage demonstrating audiometrically detectable damage. Most aminoglycoside-induced ototoxicity is associated with parenteral therapy, but it has followed topical, oral, and irrigation use of these drugs, especially neomycin. A patient should receive dosages by these routes that are no greater than the dosages given by injection. Possible predisposing factors for ototoxicity are decreased renal function, long duration of therapy, large total dosage, plasma levels exceeding the therapeutic range, previous aminoglycoside use, concurrent use of other ototoxic drugs, dehydration, and old age. There is some evidence of an inherited susceptibility to aminoglycoside-induced ototoxicity. Hearing impairment is less common in neonates and children. Two meta-analyses found no difference in the effects on hearing of single daily dosing and multiple daily dosing of aminoglycosides. The comparative effects on vestibular function have not been adequately investigated. Serial audiometry might be useful in early detection of ototoxicity. Each aminoglycoside has a slightly different spectrum of ototoxicity; the table below serves as a general guide to their relative ototoxic potentials.1–10

<table>
<thead>
<tr>
<th>RELATIVE OTOTOXIC POTENTIAL</th>
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<tr>
<td><strong>DRUG</strong></td>
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<td>Gentamicin</td>
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**Antidepressants, Heterocyclic**

The prevalence of tricyclic antidepressant–associated tinnitus is estimated to be 1%. Tinnitus can subside despite continued therapy.\(^2,4,11\)

**Azithromycin**

In elderly patients or patients with AIDS treated with 600 mg/day for *Mycobacterium avium* complex or toxoplasmosis, hearing loss occurs in 15–25%. Hearing loss occurs at all frequencies, but lower frequencies, including the speech range, are affected most often. Drug withdrawal or reduction of the dose to 300 mg/day resolves the hearing loss. Tinnitus and vestibular disturbances also occur frequently.\(^12-14\)

**Carboplatin**

Although carboplatin is generally considered to be far less ototoxic than cisplatin, it can contribute to hearing loss when used in consolidation-phase treatment after cisplatin-containing induction. IV injection of 16–20 g/m\(^2\) of *sodium thiosulfate* 2 hr after IV carboplatin showed significant protection against hearing loss in patients with CNS malignancies.\(^4,15-17\)

**Chloroquine**

Nerve deafness is a rare but consistent feature of chloroquine therapy. Its onset is usually delayed and thought of as irreversible and accompanying long-term therapy. A partly reversible case and a case resulting from only 1 g of chloroquine have been reported.\(^2-4,18\)

**Cisplatin**

Tinnitus occurs frequently and usually subsides within 1 week of drug discontinuation. It cannot be relied on to predict further ototoxicity. Hearing loss occurs frequently in patients receiving cisplatin and can be dose limiting. Audiometric abnormalities can be detected in most patients and appear within a few days after the drug is started, although a delay of several months is common. High frequencies are lost first. If therapy continues despite early hearing loss, most patients experience hearing loss in the speech frequencies. Effects are cumulative, dose related, and probably irreversible. Prolonged, low-dose therapy might produce less ototoxicity than short-term, high-dose treatment. Ototoxicity occurs more frequently in children and the elderly, and those with pre-existing hearing loss appear to be at increased risk.\(^1-5,16,19-23\)

**Deferoxamine**

Dosage-related hearing impairment occurs during long-term deferoxamine therapy. The prevalence reported varies among studies from 6% to 57%. High-frequency hearing is affected first; reversible and irreversible hearing losses have been reported.\(^4,24-26\)
Diuretics, Loop

Rapid-onset hearing loss is a frequent feature of high-dose, rapid IV administration of furosemide. The onset might be more gradual with ethacrynic acid. Renal failure is usually listed as a predisposing factor, but only renal failure patients are likely to receive large IV doses. Co-administration with aminoglycoside antibiotics is often said to result in increased ototoxicity, but one study did not confirm this. The hearing loss is usually transient, but permanent loss has been reported, more often with ethacrynic acid than with furosemide. Hearing loss and vestibular toxicity after oral therapy have been reported. Bumetanide or torsemide produce less ototoxicity than ethacrynic acid or furosemide.1–5,27,28

Eflornithine

High- and low-frequency hearing impairments are reported frequently and dizziness occurs occasionally.23

Erythromycin

Hearing loss has occasionally followed high-dose (>4 g/day) parenteral or oral therapy and does not seem to be caused by any particular salt form. Impaired hepatic or renal function and advanced age can increase the risk. The loss occurs at speech frequencies and is usually reversible, but irreversible hearing loss has been reported. Recovery usually begins within 24 hr of drug discontinuation.1,3–5,28,30

Interferons

Tinnitus and hearing loss occur frequently during parenteral interferon therapy. These effects usually resolve 1–2 weeks after drug discontinuation. Interferon beta is more ototoxic than interferon alfa.4,31

Minocycline

Reversible vestibular toxicity, manifested primarily by dizziness, loss of balance, and lightheadedness, is a frequent occurrence. This adverse effect was noted in an average of 76% of patients in 6 studies and required 12–52% of affected patients to discontinue the drug or to stop working. Other studies have found lower, but still large, percentages of patients with vestibular toxicity. Women are more susceptible than men. Onset is often during the first 2 days of therapy, and recovery begins soon after minocycline discontinuation.1,4,32–34

Nonsteroidal Anti-inflammatory Drugs

Although not as common as with salicylates, NSAIDs have been associated with hearing impairment and deafness, including some cases of permanent damage. Tinnitus and vestibular dysfunction also have been reported.1–4,35

Quinine

Tinnitus and high-frequency hearing impairment occur frequently. Although these effects are usually reversible, permanent hearing impairment has occurred with long-term therapy. Vestibular effects also have been described.1,4,5,35

Salicylates

Tinnitus, high-frequency hearing loss, and occasional vertigo are common features of salicylate intoxication. Hearing loss appears to be related to the unbound plasma salicylate level, explaining the marked interpatient variability in the total salicylate serum level at which it is first detected. Most patients demonstrating ototoxicity from salicylates are receiving long-term, high-dose therapy, such as for rheumatoid arthritis. Salicylate ototoxicity, even if severe, is almost always reversible in 48–72 hr, but permanent hearing loss has been reported.1–5,35,36

(continued)
Vancomycin

Transient and permanent hearing loss, tinnitus, and dizziness have occurred. Hearing impairment is rare with plasma levels <30 mg/L (21 µmol/L). In many of the reported cases, the patients also had been exposed to other ototoxic drugs, especially aminoglycoside antibiotics. The prevalence of purely vancomycin-induced ototoxicity is unknown but probably low, especially with the current, highly purified vancomycin products.1,4,5,37–39

Zidovudine

Audiometry determined that hearing loss occurred in 29% of 99 patients receiving antiretroviral drugs, with most cases associated with zidovudine. The prevalence of hearing loss was marked for patients >35 yr.40

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**Drug-Induced Pancreatitis**

Pancreatitis can be acute or chronic, and most drug-induced cases are acute. The diagnosis of acute drug-induced pancreatitis requires laboratory (elevated serum amylase and lipase levels) and clinical (abdominal pain) evidence. The strongest associations are made when readministration of the drug results in a recurrence of pancreatitis (ie, a positive rechallenge). Pancreatitis has occurred during therapy with many drugs; the drugs included in this table are those that present sufficient evidence to establish themselves as probable causes of pancreatitis.

**ACE Inhibitors**

There are numerous cases of ACE inhibitor–induced pancreatitis in the literature and the files of manufacturers. Captopril, enalapril, and lisinopril have been implicated. It is not possible to estimate a prevalence. A few cases have been confirmed by rechallenge.1,3–5

**Alcohol**

Alcohol is the greatest cause of drug-induced pancreatitis, easily exceeding the number of cases caused by all other drugs. Acute pancreatitis occurs in about 5% of alcoholics and usually develops after several years of alcohol abuse. It probably represents an acute flare of chronic pancreatitis.5

**Asparaginase**

The estimated prevalence of asparaginase-induced acute pancreatitis is 1–26%, with fatalities in 1.8–4.6% of cases. Many patients who develop pancreatitis during asparaginase therapy are in poor condition and receiving other chemotherapeutic agents. Asparaginase inhibits amylase and lipase production, complicating the diagnosis and evaluation of asparaginase-induced pancreatitis.1,2,7–9

(continued)
Azathioprine

There are many published cases of azathioprine-induced pancreatitis including at least 11 with positive rechallenge. Most cases occur in transplant recipients who are receiving other drugs implicated in causing pancreatitis.1,2

Calcium Salts

Pancreatitis is associated with hypercalcemia from pathologic causes, and it is likely that hypercalcemia resulting from the administration of exogenous calcium also can produce pancreatitis. There are at least 6 published cases of pancreatitis from parenteral nutrition–induced hypercalcemia.1,10

Contrast Media

Up to 11% of patients receiving contrast media through endoscopic retrograde cholangiopancreatography develop pancreatitis. Use of lower-osmolarity agents reduces the prevalence of pancreatitis.2,11

Corticosteroids

Although corticosteroids are commonly implicated as causes of pancreatitis, most of the reported cases involve disease states that predispose to pancreatitis. The weak evidence against corticosteroids is further complicated by data supporting the use of corticosteroids in the treatment of acute pancreatitis.1,2,12

Cyclosporine

Cyclosporine-induced pancreatitis was identified in 5 of 143 heart and heart–lung transplant recipients in one study. In another, 4 of 105 cyclosporine-treated renal transplant recipients developed pancreatitis, compared with only 2 of 180 azathioprine-treated patients. All cases occurred within 4 months of the start of cyclosporine therapy.3,2,13,14

Didanosine

Estimates of the prevalence of pancreatitis in didanosine-treated patients are 3–26%. A published report of didanosine treatment of 51 adult males with AIDS (10–12 mg/kg/day) found clinical pancreatitis in 12 (24%) and asymptomatic elevations of amylase and lipase levels in 10 others. Two patients died from fulminant pancreatitis. Pancreatitis might be dose related because in one study pancreatitis developed in 7 of 60 HIV-infected children receiving doses ≥360 mg/m²/day but not in any of the 35 patients receiving ≤270 mg/m²/day.1,2,15,16

Diuretics, Thiazide

Although there are at least 25 published case reports of thiazide-associated pancreatitis, the quality of the evidence is poor. Some of the cases are complicated by hypercalcemia, a known risk factor for pancreatitis.1,2

Estrogens

Estrogen therapy increases the risk of pancreatitis in patients with pre-existing hyperlipidemia, especially hypertriglyceridemia. Hypertriglyceridemia is a known cause of pancreatitis, and estrogen therapy raises serum triglyceride levels. In one report, 4 of 7 women with serum triglycerides >1500 mg/dL (17 mmol/L) while receiving postmenopausal estrogen replacement therapy (ERT) developed pancreatitis. Cases also have been reported in younger patients taking oral contraceptives. ERT is relatively contraindicated when serum triglycerides are >350 mg/dL (4 mmol/L) and absolutely contraindicated at >750 mg/dL (8.5 mmol/L).1,2,17,18

Furosemide

There are few published cases of furosemide-associated pancreatitis. The dosage range for these cases is 40–160 mg/day and most cases occurred during the first few weeks of treatment. The evidence for furosemide-associated pancreatitis is weakened by the small number of positive rechallenges.1,2,19

(continued)
Interferon Alfa

Although few cases have been reported, the association with the administration of interferon alfa is strong.20

Mercaptopurine

Inflammatory bowel disease is associated with pancreatitis. In one study of 400 patients with inflammatory bowel disease, 13 (4.25%) developed pancreatitis while receiving mercaptopurine (50–100 mg/day). Seven of the 13 were rechallenged and all developed recurrent pancreatitis, thereby establishing a strong cause-and-effect relationship. Pancreatitis developed during the first month of initial treatment in all patients and within 24 hr for 4 of the 7 rechallenges.1,2,21

Mesalamine Derivatives

Inflammatory bowel disease is associated with pancreatitis, but mesalamine, sulfasalazine, and olsalazine have been implicated in cases of acute pancreatitis confirmed by rechallenge. Positive rechallenge can occur after rectal administration.1,2,22,23

Metronidazole

Pancreatitis occurs occasionally with metronidazole. One study of 6485 HMO patients found a rate of pancreatitis requiring hospitalization of 3.9–4.6/10,000 in patients receiving metronidazole. The study did not report on nonhospitalized cases.1,2,24

Nonsteroidal Anti-inflammatory Drugs

There are isolated case reports of pancreatitis associated with most NSAIDs, but sulindac is clearly the most commonly reported. Many cases have positive rechallenges. The onset of symptoms is from 2 weeks to 9 months after initiation of therapy.1,2

Octreotide

When placebo or 100 µg doses of octreotide were administered before and immediately after endoscopic retrograde cholangiopancreatography in 84 patients, the frequencies of pancreatitis within the first 24 hr after the procedure were 11% in the placebo group and 35% in the octreotide group. Despite the higher frequency of pancreatitis, the octreotide patients were NPO for fewer days.1,2,25

Pegaspargase

The risk of pancreatitis with pegaspargase is similar to or greater than that of asparaginase. Onset is usually within a few days to 2 weeks after the start of therapy but has occurred up to 6 weeks after the start of therapy.26 (See also Asparaginase.)

Pentamidine

Injected and aerosolized pentamidine have been implicated in causing pancreatitis; a few fatalities have been reported. Most of the patients had AIDS, which might have contributed to their pancreatitis.2

Propofol

At least 25 cases of pancreatitis associated with propofol have been reported. Many, but not all, of the cases were patients who developed hypertriglyceridemia that was attributed to the lipid-containing vehicle for injectable propofol.27,28

Valproic Acid

Five of 72 valproate-treated, mentally retarded patients in one series developed pancreatitis. There are at least 50 other published cases, including some fatalities. The prevalence of asymptomatic elevations of serum amylase might be as high as 20%. There is no obvious connection with dosage or duration of therapy, although one-half of the cases occur in the first 3 months of therapy and two-thirds occur during the first year. When detected, pancreatitis is rapidly reversible after drug withdrawal.1,2,29–31

(continued)
Vinca Alkaloids

Pancreatitis with vinca alkaloid (i.e., vincristine) therapy occurs primarily in patients receiving multiple-drug therapy, making the establishment of a cause-and-effect association difficult. Animal data show that vinca alkaloids can severely disrupt pancreatic architecture.1

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Drug-Induced Sexual Dysfunction

The large subjective component of human sexual response makes the evaluation of drug-induced sexual dysfunction difficult. Variations in study design have produced widely divergent reported rates of sexual dysfunction in the “normal” or control populations. Common drug-induced sexual dysfunctions are decreased libido or sexual drive, impotence (failure to achieve or maintain an erection in men), priapism (persistent and often painful erection), delayed ejaculation or failure of ejaculation, retrograde ejaculation (into the urinary bladder), and, in women, failure to achieve orgasm and decreased vaginal lubrication. Gynecomastia (enlargement of the male breast) has been included in this table. Although not life-threatening, drug-induced sexual dysfunction has a negative effect on quality of life and is an important contributor to noncompliance with prescribed drug regimens.

Alcohol
Low doses result in behavioral disinhibition. With higher doses, sexual response is impaired, frequently resulting in failure of erection in men and reduced vaginal vasodilation and delayed orgasm in women. In chronic alcoholics, sexual dysfunction frequently persists long after alcohol withdrawal and is permanent in some. The long-term effects are probably neurologic and endocrine in origin; alcohol reduces testosterone levels and increases luteinizing hormone levels. Long-term effects are independent of liver disease.1–8

Alprostadil
Intracavernous injection of alprostadil produces penile pain in 44% of patients, prolonged erection in 6%, and priapism in 1%. Fibrotic nodules or scarring occur frequently. Intraurethral administration does not appear to cause priapism or fibrosis, but 36% experience penile pain.9–11

Aminocaproic Acid
This drug can inhibit ejaculation without affecting libido and has produced “dry” ejaculation. Effects are rapidly reversible with drug discontinuation.1,2,12,13

Amphetamines
Low doses can increase libido and delay male orgasm. High doses have been associated with failure to achieve an erection in men and loss of orgasm in both sexes.2,14–16

Anabolic Steroids
Impotence and gynecomastia occur frequently in men and might be the result of reduction in the circulating levels of natural testosterone.1,3,17

Anticonvulsants
Female and male libido can be reduced. Self-reported sexual dysfunction has been described in a widely varying percentage of patients. Social and psychological aspects of epilepsy probably play important roles in these findings. Some effects might be caused by a reduction in the level of free testosterone, resulting from hepatic enzyme induction and higher concentrations of sex hormone–binding globulins.1,4,18–20

Antidepressants, Heterocyclic
Impotence, delayed ejaculation, and painful ejaculation have been reported in men. Women and men have reported delayed orgasm and anorgasmia. Clomipramine is the worst offender. Increased and, more commonly, decreased libido have been reported in men and women. The frequency of these effects varies considerably among published reports, perhaps reflecting the influence of the underlying depressive illness.1–4,16,21–24 (See also Selective Serotonin Reuptake Inhibitors and Trazodone.)

(continued)
\(\beta\)-Adrenergic Blocking Agents

These drugs are associated with a variety of sexual problems, most commonly impotence. In a study of 46 men taking propranolol, 7 experienced “complete” impotence, 13 noted reduced potency, and 2 complained of reduced libido. In a larger trial, the frequencies of impotence during propranolol therapy were 13.8% and 13.2% after 12 weeks and 2 years, respectively. However, these figures did not differ significantly from placebo. Most of the published reports implicate propranolol; other more cardioselective \(\beta\)-blockers are less frequently associated with complaints of adverse sexual effects. There have been at least 25 reported patients who complained of sexual dysfunction (18 impotence, 9 decreased libido) while receiving topical ophthalmic treatment with timolol. Some of these patients were rechallenged, with positive results.\(^{1,4,12,25–31}\)

Calcium-Channel Blockers

Although these drugs are generally thought to be free of adverse effects on sexual function, they are associated with gynecomastia. Verapamil is the most commonly implicated calcium-channel blocker, but nifedipine and diltiazem also can produce gynecomastia. Other calcium-channel blockers seem less likely to cause gynecomastia.\(^{1,32}\)

Carbonic Anhydrase Inhibitors

Many patients receiving carbonic anhydrase inhibitors (eg, acetazolamide, methazolamide) develop a syndrome of malaise, fatigue, weight loss, and depression that often includes loss of libido. These patients appear to be more acidic than those without the syndrome and some respond to therapy with sodium bicarbonate. Decreased libido has occurred in men and women and usually requires 2 weeks of carbonic anhydrase inhibitor therapy to develop.\(^{1,2}\)

Cimetidine

In a group of 22 men treated with high dosages of cimetidine for hypersecretory states, 11 developed gynecomastia and 9 experienced impotence. These effects appear to be dose related and readily reversible and are not an important problem at dosages used for peptic ulcers. Cimetidine has some antiandrogenic effects, possibly the result of hyperprolactinemia, which are thought to be responsible for sexual dysfunction. Displacement of androgens from breast androgen receptors might contribute to the development of gynecomastia. Ranitidine does not appear to be associated with as high a prevalence of sexual dysfunction, and famotidine is not antiandrogenic.\(^{1–4,12,17,32,33}\)

Clofibrate

In large multicenter trials, impotence has been reported more frequently than with placebo.\(^{1–4,12,34,35}\)

Clonidine

Although some reports have indicated no sexual problems, others have indicated problems in up to 24% of patients. Impotence is the most frequently noted effect, but delayed or retrograde ejaculation in men and failure of arousal and orgasm in women have been described.\(^{1–4,12,18,36,37}\)

Cocaine

Although cocaine is often perceived as a sexual stimulant, its use is associated with difficulty in establishing an erection and delayed ejaculation.\(^{5,16,38,39}\)

Cypromeone

Gynecomastia results from the antiandrogen effects of cypromeone.\(^{17}\)

Danazol

Most women treated with danazol for endometriosis experience reversible decreased libido.\(^{40}\)

(continued)
Digoxin

Digitalis glycosides have some estrogen-like activity, and digoxin has been associated with decreased libido, impotence, and gynecomastia in men. In one study, digoxin use was associated with a 60% decrease in testosterone and a similar increase in estrogen in men.\textsuperscript{1–4,13,41}

Diuretics, Thiazide

In one large study, the prevalence of impotence was reported to be significantly higher with ben­droflumethiazide than with placebo (23% after 2 yr compared with 10% for placebo), and in another, hydrochlorothiazide was reported to produce more impotence and loss of libido than propranolol. In a well-designed study, 14% of men taking chlorthalidone complained of impotence, as did 14% of placebo-treated men. In three studies, chlorthalidone therapy resulted in more impotence than placebo (17% vs 8% in one).\textsuperscript{1–4,12,27–29,36,42–44}

Estrogens

Impotence and gynecomastia occur frequently in men taking estrogens for prostate cancer. Estrogens have been used to reduce libido and sexual activity of male sex offenders.\textsuperscript{1–4,45}

Finasteride

Gynecomastia occurs in 0.4% of finasteride-treated men. Onset is usually delayed until after 5–6 months of treatment.\textsuperscript{46}

Flutamide

Gynecomastia might result from the antiandrogen effects of flutamide.\textsuperscript{17}

Gonadotropin-Releasing Hormone Analogues

Most men and women treated with goserelin experience reversible decreased libido. Leuprolide-treated patients likely react similarly.\textsuperscript{40}

Growth Hormone

(See Somatropin.)

Guanethidine

Up to 54% of men have reported impotence and up to 71% have reported ejaculatory impairment. Guanethidine does not affect parasympathetic function and would not be expected to produce impotence, leading some to suggest that the impotence is secondary to the inhibition of ejaculation. Retrograde ejaculation occurs as a result of the failure of the internal urethral sphincter to close; this action is sympathetically mediated. Although not well characterized, decreased libido in women taking guanethidine has been reported. Guanethidine effects are reversible with drug discontinuation and can be alleviated by a reduction in dosage.\textsuperscript{1–4,12,36}

HMG-CoA Reductase Inhibitors

At least 47 cases of simvastatin-associated impotence have been reported, including some with positive rechallenge. There are scattered reports of impotence with lovastatin and pravastatin.\textsuperscript{47,48}

Ketoconazole

Gynecomastia has been reported, apparently the result of the inhibition of testosterone synthesis.\textsuperscript{1,17,32}

Marijuana

Positive and negative effects on sexual function are possible. Low doses can have a disinhibiting effect, whereas large doses have been associated with decreased libido and impotence. Long-term use also can result in gynecomastia.\textsuperscript{2,32,49}

(continued)
Methyldopa
Impotence and ejaculatory failure in men and reduced libido in both sexes have been described. The frequency of sexual dysfunction varies from quite low in some reports to >50% in response to direct questioning. These effects are dose related and reversible. They might be the result of drug-induced sympathetic inhibition and mild CNS depression. Gynecomastia in men and painful breast enlargement in women have occurred.1–4,12,29,36

Metoclopramide
Gynecomastia and galactorrhea have been reported in adults and children receiving metoclopramide. These effects are probably due to metoclopramide-induced hyperprolactinemia.50

Monoamine Oxidase Inhibitors
Reported adverse sexual effects of MAOIs are highly variable. Impotence, spontaneous erections, and ejaculatory delay in men and orgasmic failure in men and women have been described. The true prevalence of these effects cannot be determined from available data, but MAOIs might be associated with more sexual dysfunction than heterocyclic antidepressants.1,2,4,16

Narcotics
Long-term narcotic use (especially abuse) is frequently associated with decreased libido and orgasmic failure in both sexes and impotence in men. These effects are dose related, with the highest frequency of impotence reported in narcotic addicts (80–90% in some series), and are reversible with drug discontinuation.2,51–53

Nitrates and Nitrites
These vasodilators have been used (primarily by inhalation) to enhance the perception of orgasm. When they are used too soon before orgasm, however, the vasodilation rapidly produces loss of erection. This effect has been used therapeutically to reduce spontaneous erections in men undergoing urologic procedures.2,54,55

Omeprazole
Although the prevalence is unclear, impotence and gynecomastia in men and breast enlargement in women have been described.1,56

Papaverine
Intracavernous injection of papaverine resulted in priapism (defined as an erection lasting >3 hr) in 17% of 400 patients. Those with psychogenic or neurogenic impotence were more likely to experience priapism than those with vasculogenic impotence.9,57

Phenothiazines
These drugs have been implicated in producing a wide variety of adverse sexual effects such as impotence and priapism, absent and spontaneous ejaculation, painful ejaculation, retrograde ejaculation, menstrual irregularities, and decreased libido. These effects result from the complex actions of the drugs on the patient’s hormonal balance and central sympathetic and parasympathetic pathways. With the exception of priapism, these effects are usually benign and respond to drug discontinuation. Thioridazine is the most commonly implicated drug. The possible contribution of the underlying disease state cannot be overlooked.1–4,16,25,58

Phenoxybenzamine
This α-adrenergic blocker is associated with dosage-related failure of ejaculation but not interference with orgasm. This effect was present in all 19 patients in one study and reversed 24–48 hr after drug discontinuation.1–3,59

(continued)
Progestins

Impotence has been reported in 25–70% of men receiving progestins for prostatic hypertrophy. Progestins have been used to reduce libido and sexual activity of male sex offenders.2,4,60

Reserpine

Impotence (33%) and failure of ejaculation (14%) in men and reduced libido in both sexes occur frequently.1,2,4,12,36

Sedative-Hypnotics

In a manner similar to alcohol, low doses can produce some disinhibition, whereas large doses can reduce sexual performance.1–4

Selective Serotonin Reuptake Inhibitors

Anorgasmia and delay of orgasm are frequent adverse effects of SSRIs, affecting the majority of patients in some studies. These effects have been confirmed in patients without depression. Fluvoxamine, fluoxetine, paroxetine, and sertraline are the most frequently mentioned. Some patients have benefited from dosage reduction. Bupropion, mirtazapine, and nefazodone have limited, if any, adverse effects on sexual function and should be considered in patients with SSRI-induced dysfunction.1,3,24,61–63

Somatropin

Benign gynecomastia can occur in prepubertal and adult males receiving somatropin. Onset might not occur until after months or years of treatment.64

Spironolactone

Gynecomastia in men and painful breast enlargement or menstrual irregularities in women are frequent with large dosages. Less frequently reported effects are impotence, inhibition of vaginal lubrication, and loss of libido. The structural similarity of the drug to estrogens and progestins is thought to be a key factor in the genesis of adverse sexual effects. Spironolactone might inhibit the formation of testosterone and its breast receptor binding. It also might increase the metabolic clearance of testosterone and its rate of peripheral conversion to estradiol. These effects appear to be dosage related.1,2,4,12,17,32,36

Tamoxifen

Use of tamoxifen increases the prevalence of vaginal dryness and painful intercourse.65

Trazodone

Numerous cases of priapism have been reported, usually during the first month of therapy.1,4,12,57,66

REFERENCES


(continued)


Drug-Induced Skin Disorders

Most drugs occasionally have been associated with rashes or other dermatologic reactions. The difficulty of determining a correct diagnosis of a skin disorder and the complexity of establishing a causal relationship with drug therapy make estimating the frequency of occurrence of these reactions virtually impossible. Only skin disorders resulting from systemic administration of drugs are represented in this table. Drugs believed to be among the most common causes of a particular drug-induced skin disorder are designated by “XX” in the table. Stevens–Johnson syndrome and toxic epidermal necrolysis have been combined into a single column because of their similarity in histopathology and because they are usually caused by the same drugs. The following abbreviations are used to indicate specific skin disorders:

- **AE** — Acneiforms Eruptions
- **AL** — Alopecia
- **ED** — Exfoliative Dermatitis
## Drug-Induced Diseases

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**REFERENCES**

In the United States, fetal malformations occur in 3 to 6% of pregnancies. These include major and minor malformations from any cause, be it drug, infection, maternal disease state, genetic defect, or pollutant.\textsuperscript{1,2} Drug use during pregnancy can be associated with risk to the developing fetus and the pregnant woman. Drugs are probably responsible for only about 1 to 5% of fetal malformations; 60 to 70% of malformations have unknown causes.\textsuperscript{2–4}

The genetic makeups of the fetus and the mother influence the extent to which an agent affects the developing fetus. For example, the rates of absorption, metabolism, and elimination of an agent by the mother, its rate of placental transfer, or the way it interacts with cells and tissues of the embryo are genetically determined factors. Thus, human teratogenicity cannot be predicted based only on animal data or extrapolated from one pregnancy to another.

### PHYSIOLOGIC AND DEVELOPMENTAL FACTORS

Teratogenic substances rarely cause a single defect. Most often, a spectrum of defects occurs that corresponds with the systems undergoing major development at the time of exposure. Major malformations are usually the result of first-trimester exposure during critical periods of organogenesis. Exposures during the second and third trimesters can result in alterations or damage in fine structure and function. Intrauterine growth retardation is perhaps the most reliable indicator that a teratogen was present during the second and third trimesters of fetal development. Several organs and systems continue to develop after birth. Therefore, exposure to agents late in pregnancy carries some risk and can result in debilitating alterations in development such as mental retardation. Figure 2–1 shows the stages of human structural development in relation to teratogenic potential.\textsuperscript{5}

### DRUG FACTORS

Most chemicals in the maternal bloodstream cross the placenta. Movement of compounds across the placenta is generally bidirectional, although the net transfer occurs from mother to fetus in most instances.\textsuperscript{6,7} Although active and facilitated transport of some substances across the placenta have been demonstrated, the
Figure 2–1. Variation in teratogenic susceptibility of organ systems during stages of human intrauterine development. (Reproduced with permission from Pagliaro LA, Pagliaro AM. Problems in pediatric drug therapy. 3rd ed. Hamilton, IL: Drug Intelligence Publications; 1995.)

aAverage time for fertilization to parturition is 38 weeks.
bDrugs administered during this period can cause neonatal depression at birth (or other effects directly related to the pharmacologic effect of the administered drug).
transplacental passage of most agents occurs primarily by simple diffusion. Only the unbound (free) fraction of a drug is subject to placental transfer; therefore, the greater the degree of protein binding of a drug, the less will be transferred to the fetus. Early in pregnancy, the placental membrane is relatively thick, and this characteristic tends to reduce permeability. The thickness of the trophoblastic epithelium decreases and surface area increases in the last trimester. The passage of drugs is increased during this stage of pregnancy.

The rate-limiting factors in placental transfer of drugs are the same as those that govern membrane diffusion by molecules in general. Thus, the rate of diffusion across the placental barrier is directly proportional to the maternal–fetal concentration gradient and the surface area of the placenta. Higher concentrations are generally attained in fetal serum and amniotic fluid after bolus injection than after continuous infusion of drug into the mother and by multiple-dose rather than single-dose therapy. Certain physicochemical properties of drugs or chemicals favor transport to the fetus, including low molecular weight, lipid solubility, and nonionization at pH 7.4.

Each drug has a threshold above which fetal defects can occur and below which no effects are discernible. Whether an agent reaches a “threshold concentration” in the fetus depends on maternal factors (eg, rates of absorption and clearance) and the chemical nature of the agent.

Administration of drugs near term poses another potential threat to the fetus. Before birth, the fetus relies on maternal systems for drug elimination. After birth, the infant must rely on its own metabolic and excretory capabilities, which have not yet fully developed. Drugs given near term or during birth, especially those with long half-lives, can have an even more prolonged action in the neonate. Drugs that cause maternal addiction also are known to cause fetal addiction. Neonatal withdrawal symptoms can occur when mothers have been addicted to drugs during pregnancy or when they have taken addicting drugs near term, even though the mothers themselves are not addicted.

**EFFECTS OF PREGNANCY ON THE MOTHER**

Maternal physiology changes as pregnancy progresses and can have an effect on drug disposition and clearance. Maternal plasma volume increases by about 20% at midgestation and 50% at term and then falls toward prepregnancy levels postpartum. The volume of distribution for many drugs increases as the fetal compartment enlarges, causing changes in maternal serum drug concentrations. Drugs with narrow therapeutic ranges require careful monitoring during pregnancy and possibly dosage increases. As postpartum maternal plasma volume returns to normal, dosages of many drugs require reduction. Changes in plasma protein concentrations during pregnancy can affect the degree of binding and thus the amount of unbound drug. Despite an increase in production of serum albumin, the increased intracellular and intravascular volumes cause serum albumin concentrations to decline. A decrease in total plasma protein concentrations of about 10 g/L occurs during pregnancy. Body fat increases by 3–4 kg during pregnancy and can act as a depot for fat-soluble drugs, thereby increasing their volume of distribution. Renal blood flow and glomerular filtration rate increase by almost 50%
during pregnancy because of increased cardiac output. Renally excreted drugs therefore can have increased rates of clearance.9

**INTERPRETATION OF STUDIES**

There are few controlled, prospective studies of drug use in pregnancy. Most of the available information comes from case reports or case-control studies. Cause-and-effect relationships between drugs and teratogenicity are difficult to establish retrospectively because of the numerous variables in each report. These include maternal drug dosage, time of ingestion relative to the date of conception, duration of therapy, concomitant exposure to other potential teratogens, and questionable study design or methodology. Because studies cannot disprove that a slight teratogenic risk might occur with in utero exposure to drugs, drugs should be used during pregnancy only when absolutely necessary. The following table provides information concerning the effects of drugs used during pregnancy on the pregnant woman and on pregnancy outcome. For a more thorough discussion of the principles of teratology, the reader should consult reference 1.

The following abbreviation is used in the table:

IUGR—intrauterine growth retardation, less than the 10th percentile (of an appropriate standard) birth weight for gestational age.12

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**DRUGS AND PREGNANCY**

**ANALGESIC AND ANTI-INFLAMMATORY DRUGS**

**ANTIMIGRAINE DRUGS**

**Ergotamine**

Ergotamine can stimulate uterine contraction and potentially cause abortion.13

**Sumatriptan**

Evidence collected through the Swedish Medical Birth Registry indicates no increased risk of birth defects in 658 pregnancies with drug exposure.14 A prospective study of 86 women showed no increased risk of birth defects.15 Sumatriptan does not have the oxytocic effect of the ergot alkaloids.

**NONSTEROIDAL ANTI-INFLAMMATORY DRUGS**

**Acetaminophen**

Acetaminophen does not cause congenital malformations and is the analgesic-antipyretic of choice for use near term because it does not affect platelet function or peripheral prostaglandin synthesis.16,17 In maternal acetaminophen overdose, most infants are normal at birth,18–21 but there have been a few cases of neonatal liver toxicity.18,22 Acetaminophen might prevent fetal distress in laboring women with chorioamnionitis and fever.23

(continued)
ANALGESIC AND ANTI-INFLAMMATORY DRUGS

Nonsteroidal Anti-inflammatory Drugs

Early case reports implicating indomethacin as a cause of prenatal closure of the ductus arteriosus are inconclusive. Indomethacin might cause oligohydramnios because of decreased fetal urine output, which places the fetus at risk for pulmonary hypoplasia and umbilical cord compromise. Indomethacin inhibits uterine contractions and has been used as a tocolytic agent. Echocardiographic surveillance of the fetus might be indicated to monitor effects on the ductus arteriosus.

A large prospective study of more than 50,000 pregnancies did not show an increased risk of birth defects, altered birth weight, or perinatal deaths associated with exposure to aspirin. First-trimester aspirin use does not increase the risk of congenital heart defects compared with other structural malformations. Repeated third-trimester administration of aspirin 325 mg can result in prolonged constriction of the ductus arteriosus and pulmonary hypertension. Maternal ingestion of aspirin 325 mg during the third trimester can interfere with uterine contractility and prolong gestation and labor. Maternal and neonatal platelet function can be affected, resulting in increased maternal blood loss at delivery and abnormal platelet function tests and clinical bleeding in newborns, including intracranial hemorrhage. Second- or third-trimester use of low-dose (20–100 mg/day) aspirin in mothers at risk of developing pregnancy-related hypertension decreases the frequency of this disorder and its complications. A study of more than 9000 women found that 60 mg/day was not protective. A follow-up 18 months after delivery of infants exposed in utero to aspirin 50 mg/day showed no increase in malformations and normal physical and neurologic development. Newborns exposed to low-dose aspirin have not been found to have bleeding abnormalities. Systematic evaluations of other commonly used NSAIDs have not been conducted in humans, but no substantive reports of NSAID teratogenicity exist. However, caution is warranted because of their similarity to indomethacin and aspirin. Ibuprofen can cause mild oligohydramnios and mild constriction of the fetal ductus arteriosus. Persistent pulmonary hypertension occurred in a neonate whose mother had ingested 5 g naproxen 8 hr before delivery. There are no reports of adverse effects of ketorolac; it is usually avoided during pregnancy.

OPIOIDS

Narcotics

Narcotic analgesics do not cause fetal malformations, but narcotic abuse during pregnancy or use near term can lead to fetal tolerance and neonatal withdrawal. Meconium might be present in the amniotic fluid, caused most likely by increased bowel activity during periods of fetal withdrawal and/or hypoxia, putting the fetus at risk for meconium aspiration. (See Heroin.) Withdrawing symptoms such as irritability, increased muscle tone, sleep disturbances, vague autonomic nervous system symptoms, tremulousness, high-pitched crying, frantic and uncoordinated sucking, and seizures can occur in neonates born to narcotic-addicted women and nonaddicted women using narcotics near term. Neonatal respiratory depression can occur when narcotic analgesics are given during labor and is dependent on the drug, dose, dosing interval, and route of administration (IV > IM). Epidural alfentanil can cause neonatal hypotonus. Meperidine crosses the placenta rapidly and can cause a sinusoidal fetal heart rate pattern. It is eliminated by the fetus at a rate much slower than the mother’s; its metabolite, normeperidine, is long acting. Meperidine given during delivery can interfere with the early establishment of breastfeeding because of infant sedation. Meperidine by patient-controlled analgesia (PCA) for postcesarean pain causes a much greater decrease in neonatal alertness and sucking than an equivalent dosage of PCA morphine. Infants born to narcotic-dependent women maintained on methadone during (continued)
ANALGESIC AND ANTI-INFLAMMATORY DRUGS

Pregnancy is reported to have lower birth weights, jaundice, thrombocytosis, and withdrawal. Divided doses of methadone better stabilize the fetal activity pattern (which might indicate fetal withdrawal) before and after drug administration than do single daily doses. It is not known if opioids can cause alterations in the neurobehavioral function of infants exposed in utero.

Narcotic Partial Agonists

All narcotics can cause respiratory depression and possibly some behavioral abnormalities in the newborn if used at the time of delivery. Buprenorphine used by a pregnant woman daily near term resulted in a mild narcotic-like withdrawal syndrome in her newborn. Butorphanol and nalbuphine during labor can cause a sinusoidal fetal heart rate pattern. Nalbuphine offers no advantage over pure narcotics and can cause more abnormal Apgar scores (<7) at 1 min. Nalbuphine with pentazocine use throughout pregnancy has caused infant withdrawal symptoms similar to those reported in offspring of heroin and methadone addicts. Small-for-gestational-age infants, prematurity, and fetal distress also have been observed. Tramadol withdrawal has been reported in a neonate exposed in utero throughout pregnancy.

OTHER ANTI-INFLAMMATORY DRUGS

Gold Salts

Although teratogenic in animals, reports have described normal children born to women using gold salts during pregnancy. A few cases of musculoskeletal problems and one case of IUGR occurred among 128 infants exposed to parenterally administered gold during pregnancy. Six women given auranofin during pregnancy delivered normal infants.

Penicillamine

Data on the teratogenicity of penicillamine are contradictory. Most pregnant women taking penicillamine deliver normal healthy babies, even at the high dosages used in treating Wilson’s disease. However, there have been cases of fetal connective tissue abnormalities. Penicillamine is best avoided during pregnancy.

ANTIMICROBIAL DRUGS

AMINOGLYCOSIDES

There is no evidence that aminoglycosides are teratogenic. Streptomycin can cause congenital hearing loss, ranging from minor high-frequency loss to total deafness, when given to pregnant women for the treatment of tuberculosis. Prevalence is low, especially with careful dosage calculation and limited duration of therapy. There is a theoretical risk of nephrotoxicity and ototoxicity for all aminoglycosides. There is no evidence that aminoglycosides are teratogenic. Streptomycin can cause congenital hearing loss, ranging from minor high-frequency loss to total deafness, when given to pregnant women for the treatment of tuberculosis. Prevalence is low, especially with careful dosage calculation and limited duration of therapy. There is a theoretical risk of nephrotoxicity and ototoxicity for all aminoglycosides, including streptomycin. There is no evidence that aminoglycosides are teratogenic.

ANTIMYCOBACTERIAL DRUGS

Antituberculars

The treatment of choice for tuberculosis during pregnancy is isoniazid and rifampin with ethambutol added if isoniazid resistance is suspected. Isoniazid is the safest and most effective antitubercular during pregnancy, although there can be an increased risk of hepatotoxicity in pregnant women. Of the reported isoniazid exposures during pregnancy, only 1% demonstrated any malformation. No pattern of malformation was shown, but several abnormalities involved the CNS. Exposures were confounded by concomitant ethambutol therapy. Rifampin safety during pregnancy is less well established; however, it has not been associated with an increased risk of fetal malfor- (continued)
ANTIMICROBIAL DRUGS

In most reports, rifampin was taken with isoniazid or ethambutol. Neonatal hypoprothrombinemia has been reported and raises some concern about the use of rifampin, especially near term. If rifampin is given during pregnancy, maternal oral prophylaxis with vitamin K 20 mg/day for 2 weeks before delivery is recommended. Infants should receive 0.5–1 mg of vitamin K IM or SC immediately after delivery and again 6–8 hr later. Ethambutol does not appear to cause malformations, but several anomalies involving the CNS occurred in 655 reported exposures.

Sulfones

Dapsone does not appear to increase the risk of fetal abnormalities. There are, however, some reports of hemolytic anemia in mothers and their infants after dapsone use. Because dapsone is similar to sulfonamides, it might displace bilirubin from albumin binding sites and increase the risk of kernicterus in the infant due to hyperbilirubinemia. This risk is minimized if the drug is discontinued 1 month before the expected date of delivery. (See Sulfonamides.)

Thalidomide

Thalidomide causes bilateral limb reduction defects, facial hemangioma, esophageal or duodenal atresia, and anomalies of the kidneys, heart, and external ears. The time of greatest risk is between gestational days 22 and 32. If exposure occurs between days 27 and 30, the arms are most often affected; with exposure between days 30 and 33, the legs and arms are affected.2

ANTIPARASITIC DRUGS

Antimalarials

Chloroquine is the drug of choice for prophylaxis and treatment of malaria during pregnancy. Chloroquine or hydroxychloroquine malaria prophylaxis does not cause adverse fetal effects; however, larger anti-inflammatory doses have resulted in spontaneous abortion and fetal retinal and vestibular damage. These drugs are best avoided in anti-inflammatory dosages during pregnancy. There is a risk of hemolysis in fetuses that are G-6-PD deficient. There are no reports of primarquine teratogenicity. It might induce hemolysis in neonates by the same mechanism as in adults. Pyrimethamine is a microbial folate antagonist and should be used cautiously because the mammalian folate antagonist methotrexate is teratogenic; folic acid supplementation might be warranted during treatment. In a study of 210 women exposed during pregnancy, however, no increased risk of birth defects was observed. Quinine has been used as a folk medicine abortifacient, despite its poor efficacy. Maternal deaths have been reported. Fetal anomalies include blindness, optic nerve hypoplasia, deafness, and hearing impairment.

ANTIVIRAL DRUGS

Acyclovir

Maternal acyclovir has not been shown to cause malformations and is recommended for various infections during pregnancy.

Zidovudine

Use of zidovudine in pregnant women has not been shown to cause an increased rate of birth defects. Concentrations of zidovudine are 2.5–7 times higher in amniotic fluid than in cord blood; concentrations in cord blood are higher (113 to 140%) than those in maternal blood. Transmission of HIV from mother to fetus is substantially reduced with zidovudine treatment, and treatment is recommended for HIV-infected pregnant women. In a long-term study of almost 200 children prenatally exposed to zidovudine, no adverse effects were observed on lymphocyte function, height, weight, or malignancy.

(continued)
ANTIMICROBIAL DRUGS

β-LACTAMS

Cephalosporins and penicillins are thought to be without teratogenic risk. Treatment of early syphilis with penicillin (or other drugs) during pregnancy might produce the Jarisch–Herxheimer reaction, resulting in uterine cramping, decreased fetal movement, and, in some cases, fetal death.

MACROLIDES

There is no evidence that erythromycin is harmful to the fetus. About 10–15% of women treated with erythromycin estolate in the second trimester develop elevated serum AST levels that normalize when therapy is discontinued; other derivatives might be preferred. Preliminary studies of women exposed to clarithromycin during pregnancy show no increased risk of anomalies. Clarithromycin currently is not recommended during pregnancy because of limited data.

QUINOLONES

Nalidixic acid use during pregnancy can cause increased intracranial pressure, papilledema, and bulging fontanelles in the newborn. Avoid first-trimester use. Fluoroquinolones (eg, ciprofloxacin, norfloxacin, ofloxacin) cause arthropathy in immature animals, but preliminary data have not shown this effect in humans.

SULFONAMIDES

There are occasional reports of abnormalities, but no malformation pattern has emerged. Evidence associating sulfonamide use near term with neonatal kernicterus is lacking, despite sulfonamide displacement of bilirubin from albumin binding sites. There is a theoretical risk for hemolysis in the fetus or neonate because of its relative deficiencies in G-6-PD and glutathione.

TETRACYCLINES

Tetracyclines can cause depression of fetal bone growth and permanent staining of the teeth when taken after the 12th week of pregnancy. Rebound bone growth can follow tetracycline discontinuation. The risk of enamel discoloration increases with dose, duration of therapy, and advancing pregnancy; one-third to one-half of third-trimester exposures can be affected. Pregnant women with pyelonephritis or underlying renal disease or after an overdose are at risk for developing acute fatty necrosis of the liver and azotemia. Doxycycline might be less likely to discolor enamel.

MISCELLANEOUS ANTIMICROBIALS

Chloramphenicol

Although reports of fetal abnormality or toxicity from maternal chloramphenicol are lacking, there is a theoretical risk of blood dyscrasias. Particular caution should be exercised near term because of the “gray baby” syndrome, the result of toxic accumulation of chloramphenicol in neonates caused by their slow elimination of the drug.

Metronidazole

Twenty years’ experience and several studies demonstrate no association of metronidazole with congenital malformations, abortions, or stillbirths. However, a few cases of facial clefting have been reported. Metronidazole has not been demonstrated to be carcinogenic in humans.
ANTIMICROBIAL DRUGS

Nitrofurantoin

No fetal abnormalities or neonatal hemolytic anemia have been observed with nitrofurantoin use during pregnancy.6,55,98–100 A theoretical risk for the development of hemolysis in neonates exists if the drug is taken by mothers near term because of infants’ relative G-6-PD and glutathione deficiencies.100

Trimethoprim

Because it is a folate antagonist, caution is advised in pregnancy.6,64 (See Methotrexate and Pyrimethamine.) However, data suggest a lack of teratogenicity.6,55,81,88,89,100,101–103 Neither single-dose trimethoprim 600 mg nor the usual 5-day course (300 mg/day) for asymptomatic bacteriuria caused any detrimental effects on pregnancy outcome when given between the 16th and 30th weeks of gestation.103

Vancomycin

There is a theoretical risk for auditory and renal toxicity in the fetus.6,53 However, in one small prospective trial, IV vancomycin during the second or third trimester produced no cases of fetal renal toxicity or hearing impairment.104

ANTINEOPLASTICS AND IMMUNOSUPPRESSANTS

ANTINEOPLASTICS

Antineoplastic agents have teratogenic and mutagenic potential, and reports of infertility and congenital defects exist. Nevertheless, several studies indicate that fertility is preserved, with normal pregnancy outcome, among women and men treated for cancer before conception. Although aggressive treatment of malignancy is necessary on occasion, avoidance or minimum use of these drugs, especially during the first trimester, is recommended because of reports of teratogenicity and spontaneous abortion. However, normal pregnancy outcome has occurred, particularly if exposure is early in gestation (first 2 weeks after conception).105 Chemotherapy generally produces a decrease in birth weight but not IU GR.106 Use of antineoplastics near delivery might cause neonatal bone marrow suppression107 because the drugs might not have been eliminated before delivery. Chemotherapy should be avoided within 3 weeks of delivery, if possible.105 Chemotherapy-induced tumors in the infant are a theoretical possibility.

ALKYLATING AGENTS

Busulfan

Use during pregnancy has been associated with IUGR and multiple malformations, although no specific pattern is evident.

Chlorambucil

Treatment in the first and second trimesters can cause spontaneous abortion, cleft palate, renal aplasia, or skeletal abnormalities.105 There also have been reports of normal pregnancy.108

Cyclophosphamide

Use in the first trimester has resulted in fetal malformations, particularly of the toes; syndactyly; cleft palate; facial anomalies; IUGR; and possible developmental delay. The overall risk of malformation is estimated to be 16–22%. No malformations have been reported with second- or third-trimester use, but pancytopenia occurred in neonates exposed late in pregnancy. A report of malignancies in a child that occurred 10 and 14 yr after exposure in utero raises the question of

(continued)
ANTINEOPLASTICS AND IMMUNOSUPPRESSANTS

whether intrauterine exposure to cyclophosphamide can cause iatrogenic or secondary cancers.\textsuperscript{109} Cyclophosphamide should be avoided in pregnancy.

\textbf{Procarbazine}

Use in pregnancy is limited, but several fetal abnormalities have occurred, all with first-trimester exposure.\textsuperscript{40}

\textbf{ANTIMETABOLITES}

\textbf{Fluorouracil}

Use can cause malformations consistent with inhibition of cell division and cell growth.\textsuperscript{40} Inadvertent first-trimester topical administration of 5\% fluorouracil to the lower genital tract did not result in abnormal-appearing infants in 10 pregnancies.\textsuperscript{110,111}

\textbf{Methotrexate}

Use in the first trimester causes spontaneous abortion and congenital abnormalities such as cranial anomalies, cleft palate, syndactyly, growth retardation, and developmental abnormalities.\textsuperscript{2} One study of 10 pregnancies in 8 women taking methotrexate 7.5–10 mg/week during the first trimester for arthritis resulted in three spontaneous abortions, two elective abortions, and the birth of five full-term normal infants who had no medical illnesses or learning disabilities. Normal pregnancy outcome has occurred after use in the second and third trimesters.\textsuperscript{2,112} It is recommended that methotrexate be discontinued for at least three menstrual cycles before conception and that the mother be taking folic acid.

\textbf{Thioguanine}

Use in the first and second trimesters has been associated with abnormalities.\textsuperscript{40,113}

\textbf{DNA INTERCALATING DRUGS}

\textbf{Daunorubicin} and \textbf{doxorubicin} have been given without resultant fetal malformation, although some spontaneous abortions have occurred.\textsuperscript{109} Premature delivery and transient bone marrow suppression have been reported.\textsuperscript{114}

\textbf{MITOTIC INHIBITORS}

Most reports of \textbf{vinblastine} and \textbf{vincristine} use during pregnancy describe a lack of congenital malformations.\textsuperscript{109}

\textbf{IMMUNOSUPPRESSANTS}

\textbf{Azathioprine}

Normal pregnancy outcome has occurred with azathioprine taken for renal transplantation, SLE, or acute or chronic leukemias.\textsuperscript{115} However, IUGR, neonatal lymphopenia, hypogammaglobulinemia, thymic hypoplasia, fetal bone marrow suppression, leukopenia, and thrombocytopenia have occurred.\textsuperscript{116} Azathioprine for inflammatory bowel disease with or without a corticosteroid plus sulfasalazine resulted in no congenital anomalies or developmental problems.\textsuperscript{116} Older studies showed some chromosomal aberrations; however, there is no evidence of permanent genomal or gonadal damage.

\textbf{Cyclosporine}

Use of cyclosporine throughout pregnancy after renal or hepatic transplant does not appear to cause malformations, although experience is limited.\textsuperscript{52} IUGR occurred in 11 of 20 infants, and
ANTINEOPLASTICS AND IMMUNOSUPPRESSANTS

6 had severe IUGR (below the 3rd percentile). However, normal-size-for-gestational-age infants are often delivered. No neonatal distress or increased mortality has been reported.\textsuperscript{117,118}

\textbf{Mycophenolate}

There are no data on the safety of mycophenolate taken during pregnancy.

\textbf{Tacrolimus}

Use in pregnancy after liver transplant resulted in preterm deliveries of 27 infants of 21 mothers.\textsuperscript{119} However, prenatal and postnatal growth were appropriate for age. One infant was born with unilateral polycystic kidney disease.

CARDIOVASCULAR DRUGS

ANTIARRHYTHMIC DRUGS

\textbf{Amiodarone}

Neonatal hypothyroidism with and without goiter and hyperthyroidism have occurred with amiodarone use.\textsuperscript{120} A child with transient congenital hypothyroidism showed some delay in motor development and impaired speech performance at age 5 yr.\textsuperscript{120}

\textbf{Digoxin}

Digoxin is not a teratogen. It may be given to pregnant women to treat fetal CHF and supraventricular tachycardia. Maternal digitalis toxicity has caused fetal toxicity and miscarriage in one case and neonatal ECG changes with subsequent infant death in another. As pregnancy progresses, renal clearance of digoxin increases and its bioavailability can increase. Therefore, maternal serum concentrations might decrease or increase and should be monitored.\textsuperscript{121,122}

\textbf{Disopyramide}

Use during pregnancy has not been well studied, but in one report, uterine contractions precipitated by disopyramide subsided when the drug was discontinued.\textsuperscript{121}

\textbf{Procainamide}

Limited data indicate that procainamide use during pregnancy is probably safe. However, because of the potential for SLE, caution is advised.\textsuperscript{121}

\textbf{Quinidine}

Quinidine is not teratogenic.\textsuperscript{40,121} Neonatal thrombocytopenia has occurred after maternal use of quinidine.\textsuperscript{40}

ANTIHYPERTENSIVE DRUGS

\textbf{ACE Inhibitors}

Several cases of IUGR and prolonged neonatal anuria and hypotension with resultant renal failure have occurred after maternal \textit{captopril} or \textit{enalapril} use.\textsuperscript{123–126} Oligohydramnios or anhydramnios was present in 7 of 9 cases (not recorded in the other 2 cases) and led to pulmonary hypoplasia and death in some. Respiratory problems occurred in several neonates. Some infants had altered or absent skull formation with dysmorphic facial features, microcephaly, and occipital encephalocele. One infant exposed during the second and third trimesters had prolonged anuria and hypotension requiring dialysis, and one infant exposed throughout pregnancy had hypoglycemia; others have had persistent ductus arteriosus (this might have been caused by low birth weight).\textsuperscript{125–127}

(continued)
CARDIOVASCULAR DRUGS

There might be an increased frequency of fetal loss with ACE inhibitors. The FDA warns against their use in the second and third trimesters.

Diazoxide

When given by rapid IV bolus, diazoxide can cause excessive maternal hypotension and fetal distress. Slow IV infusion or minibus administration might prevent these effects. Other reported effects are inhibition of labor, neonatal hyperglycemia when exposure preceded delivery, alopecia, hypertrichosis lanuginosa, and decreased bone age after exposure in the last 19–70 days of gestation. Other investigators report no problems after long-term oral administration of diazoxide.

Hydralazine

Hydralazine is not associated with congenital anomalies. Use in pregnancy can cause reduced uteroplacental blood flow, fetal heart rate changes after acute administration, and neonatal hypothermia and thrombocytopenia.

Methyldopa

Methyldopa in pregnancy has been studied more extensively than any other antihypertensive. Available data show no teratogenicity. Transient reduction in neonatal blood pressure can occur after maternal methyldopa ingestion. There is a questionable association of IUGR and maternal treatment with methyldopa. One study suggested IUGR was caused by chronic hypertension rather than by methyldopa.

Reserpine

When given to mothers within 24 hr of delivery, reserpine produces edema of the neonatal nasal mucosa. This effect is especially important because newborns are obligate nose breathers. Lethargy, hypothermia, and bradycardia also have been reported in infants whose mothers received antenatal reserpine.

β-ADRENERGIC BLOCKING DRUGS

β-Blockers such as atenolol, pindolol, metoprolol, and propranolol are not associated with congenital anomalies and are generally safe in pregnancy. Maternal hypertension can cause IUGR, decreased placental size, neonatal respiratory depression, and hypoglycemia. IUGR and neonatal hypoglycemia and hypotension have occurred after taking β-blockers. Whether these effects are caused by the drugs or maternal disease has not been established. Neonatal bradycardia might be caused by these agents; it is usually mild but was severe after an intravenous infusion in one case. β-Blockers can adversely affect fetal adaptation to intrauterine hypoxia such as that associated with umbilical cord compression. Although generally regarded as safe during pregnancy, labetalol given intravenously to control severe hypertension has caused neonatal bradycardia, weak femoral pulses, inadequate breathing, hypotonia, hypotension, and hypoglycemia.

CALCIUM-CHANNEL BLOCKING DRUGS

No malformations have been associated with the use of calcium-channel blocking agents during pregnancy. Bolus doses of IV isradipine during labor decrease maternal blood pressure and increase fetal heart rate. Nifedipine does not appear to alter uterine arterial resistance when given at 17–22 weeks or 26–35 weeks of gestational age.
CARDOVASCULAR DRUGS

INOTROPIC DRUGS
The treatment of hypotension during pregnancy with sympathomimetic agents (eg, dopamine, dobutamine, norepinephrine) is complicated by the fact that the uterine vasculature is supplied solely with α-adrenergic receptors and maximally dilated under basal conditions. Pure α-adrenergic agents markedly constrict uterine vessels and decrease blood flow, thereby compromising the fetus. β-Adrenergic agents cause peripheral vasodilation and tend to shunt blood away from the uterus and also can cause fetal compromise. Volume-expanding agents seem to be the most prudent treatment for sudden hypotension in pregnancy.

CENTRAL NERVOUS SYSTEM DRUGS

ANTICONVULSANTS
Many congenital malformations have been reported in children of epileptic mothers, and all anticonvulsants for which data are adequate have been implicated as possible causes of malformations. Epileptic mothers have increased frequencies of fetal malformation compared with nonepileptic mothers, and anticonvulsant drugs appear to increase these frequencies. Major malformations seem to be more common after combination therapy than with monotherapy. Fetal deficiency of epoxide hydrolase, a major enzyme in the metabolic pathway of many anticonvulsants (eg, carbamazepine, phenytoin, and valproic acid) that helps eliminate toxic intermediates, might mediate teratogenic effects. Deficient function of this enzyme might be inherited as an autosomal recessive trait. Total concentrations of carbamazepine, phenytoin, phenobarbital, and valproic acid decline as pregnancy progresses, caused mainly by changes in plasma protein binding. However, free or unbound drug concentrations fall appreciably only for phenobarbital. Free concentrations of valproic acid increase. Measurement of free anticonvulsant drug concentrations allows for appropriate dosage adjustment. Decreased fetal folic acid concentrations have been reported with several anticonvulsants (eg, carbamazepine, phenobarbital, phenytoin, and valproic acid). Folic acid supplementation during pregnancy might decrease the risk of abnormal offspring.

Carbamazepine
Carbamazepine is associated with a 1% risk of spina bifida. Malformations similar to those ascribed to other anticonvulsants also have been reported: specific facial features, nail hypoplasia, and small head circumference. Data concerning developmental delay or impairment require substantiation. Higher serum concentrations of carbamazepine were found in mothers of abnormal offspring than in mothers of normal offspring. Mothers receiving carbamazepine should receive supplemental folic acid.

Gabapentin
There is little information about gabapentin in pregnancy. Of 9 exposed women reported to the manufacturer, 4 had elective abortions, 4 had normal infants, and 1 infant had pyloric stenosis and an inguinal hernia.

Lamotrigine
There are no studies of the safety of lamotrigine in pregnancy. The manufacturer maintains a pregnancy registry with birth outcomes reported. As of 1994, no constellation of defects was observed.

Oxazolidinones
Long-term use of trimethadione or paramethadione during pregnancy has resulted in children with abnormalities such as mental deficiency, prominent forehead with V-shaped eyebrows, epi-

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canthal folds, microcephaly, low-set ears with anteriorly folded helices, short stature, and hypospadias. This syndrome of abnormalities is called fetal trimethadione syndrome. The frequency of spontaneous abortion also might be increased.2,40,141

Phenobarbital

Abnormalities have been reported with phenobarbital alone and in combination, but causality is not established. Malformations similar to those of phenytoin and alcohol have been reported with phenobarbital and might be related to the folate deficiency that each of these can cause.141 These effects might be more likely when maternal serum concentrations exceed usual therapeutic concentrations. Barbiturates can cause a decrease in vitamin K-dependent clotting factors, leading to bleeding in the newborn.40,141 Neonatal withdrawal can occur after phenobarbital use during pregnancy.

Phenytoin

Phenytoin is a teratogen that causes a number of anomalies such as heart defects and facial clefts. It also can cause a cluster of anomalies called the fetal hydantoin syndrome (FHS), the principal features of which are craniofacial anomalies (eg, bowed upper lip, ocular hypertelorism, broad nasal bridge, short nose, epicanthal folds), digital hypoplasia with small or absent nails, and pre- and postnatal growth deficiency.2,141,151 The risk of developing the full-blown FHS is about 5 to 10% when phenytoin is taken throughout pregnancy; the risk of a less serious effect is 30%.152 Serum anticonvulsant concentrations were higher in mothers of malformed infants than in mothers of normal infants in some studies.149 Maternal phenytoin use also can result in developmental delay and intellectual impairment.148 Phenytoin can cause a decrease in vitamin K-dependent clotting factors, leading to bleeding in the newborn.40,141 Phenytoin serum concentrations often decrease during pregnancy because of increased plasma clearance. Adjustment of phenytoin dosage to maintain serum free drug concentrations seems to improve seizure control.153

Primidone

The teratogenicity of primidone alone is difficult to assess because few cases have been reported and primidone is often taken with other anticonvulsants, primarily phenytoin. Although no specific pattern is established, reported malformations include craniofacial alterations, hirsute forehead, cardiac defects, pre- and postnatal growth retardation, digital hypoplasia with small or flat nails, inguinal hernias, and hypospadias.40,141 Some cases of developmental delay also have been reported. These effects might be more likely when maternal serum concentrations exceed usual therapeutic concentrations. Phenobarbital is one metabolite of primidone. (See Phenobarbital.)

Valproic Acid

Neural tube defects (eg, spina bifida) occur in 1 to 2% of valproic acid–exposed fetuses compared with 0.4% in fetuses exposed to other anticonvulsants.141,152 The risk for neural tube defects might be 5- to 10-fold higher with valproic acid compared with the background frequency.142 The rate of spina bifida appears to be related to valproate serum concentration. Serum concentrations should be kept as low as possible during pregnancy and the mother should be given supplemental folic acid.141 External ear anomalies, congenital heart defects, hypospadias, craniofacial anomalies, low birth weight, and small head circumference have been reported.141,150,152,154–156 Cases of limb reduction defects, radial ray aplasia, talipes equinovarus, developmental delay, neurologic abnormality, and brain atrophy have been reported.150,156 Congenital liver damage has been described.157

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CENTRAL NERVOUS SYSTEM DRUGS

ANTIDEPRESSANTS

Selective Serotonin Reuptake Inhibitors

Prospective follow-up of exposed pregnancies, including first-trimester exposure and spontaneous reports to the manufacturer, do not implicate fluoxetine as a teratogen. However, a prospective study found increased risks of multiple minor malformations, prematurity, large-for-gestational-age size, admission to a special care nursery, and poor neonatal adaptation. Several case reports of neonatal withdrawal after maternal fluoxetine or sertraline use during the third trimester have been reported. Symptoms are restlessness, irritability, crying, tremors, increased muscle tone, poor feeding, and sleep disturbance. Neurodevelopmental assessments in offspring exposed to fluoxetine did not substantiate behavioral teratogenicity.

Tricyclic Antidepressants

Although there are several case reports of different fetal anomalies after maternal use of tricyclic antidepressants (TCAs) during pregnancy, no consistent pattern of malformation has been observed and they are not considered to pose a teratogenic risk. Maternal use of TCAs during pregnancy occasionally has produced neonatal symptoms of breathlessness, respiratory distress, hypertonia with tremor, clonus, spasm, cyanosis, irritability, and feeding difficulties. In one case, neonatal urinary retention occurred after maternal nortriptyline use. Neurodevelopmental assessments of offspring exposed in utero did not show abnormalities.

ANTIPSYCHOTIC DRUGS

There is some evidence that women with psychoses have twice the rate of fetal malformation as the general population. Most data do not implicate phenothiazines, haloperidol, or clozapine as teratogens. Phenothiazine use near term can result in extrapyramidal effects and withdrawal reactions in the neonate. If drug therapy is necessary during pregnancy, high-potency agents (eg, haloperidol, fluphenazine) are preferred to low-potency agents (eg, chlorpromazine, thioridazine) because the latter can cause maternal hypotension.

ANXIOLYtics, SEDATIVES, AND HYPNOTICS

High dosages of any sedative-hypnotic close to or during delivery can result in neonatal CNS and respiratory depression.

Anesthetics, General

Single exposures to general anesthesia in early pregnancy have not been associated with an increased risk of birth defects. There is, however, a possible association between general anesthesia for surgery performed between gestational weeks 4 and 5 and neural tube defects, and first-trimester exposure to general anesthesia is associated with hydrocephalus and eye defects. However, not all patients in these studies received the same preoperative medications or inhalation anesthetics, although nitrous oxide was received by >98% in one study and a causal relationship between other factors (eg, underlying disease) could not be ruled out. Although a 2- to 4-fold increase in the rate of spontaneous abortion in pregnant women with long-term exposure to inhalation anesthetics (eg, operating room and dental personnel) has been suggested, poor study design makes these conclusions suspect. CNS and respiratory depression in the neonate can occur after inhalation anesthetic use during labor. During labor, halothane is not recommended because of its low analgiesic activity. Nitrous oxide, methoxyflurane, and enflurane are commonly used. Halogenated anesthetics can cause uterine relaxation and, theoretically, increase maternal blood loss. Neonatal EEG recordings have documented that thiopental

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CENTRAL NERVOUS SYSTEM DRUGS

general anesthesia has more neonatal depressant effects than 0.5% bupivacaine epidural anesthesia when used for cesarean section.170

Barbiturates

Long-term use of barbiturates during pregnancy for control of seizures might be associated with an increased risk of birth defects. First-trimester use of thiopental or pentobarbital has not been associated with defects.171 Barbiturate addiction during pregnancy can result in neonatal withdrawal. Symptoms include tremors, irritability, restlessness, high-pitched crying, increased tone, hyperphagia, and overreaction to stimuli, which can persist for up to 6 months.40 (See Anticonvulsants and Phenobarbital.)

Benzodiazepines

Recent data do not support a previously reported association between diazepam or chlor-diazepoxide and oral clefting.12,40,161,172–174 Infants of mothers using benzodiazepines near term might exhibit withdrawal symptoms (including tremors, irritability, and hypertonicity) and cardiovascular, respiratory, and CNS effects consistent with benzodiazepine pharmacology. Many exhibit the “floppy baby syndrome” characterized by muscular relaxation, poor sucking, disturbances in thermoregulation, and regurgitation.13,40,169,172,175–177 Oxazepam, lorazepam, and temazepam are short acting and predominantly metabolized into inactive glucuronides and subsequently excreted by the kidneys; therefore, they might be preferred to diazepam, although irritability, feeding difficulties, and muscle tone disorders lasting 2 months were reported in two infants of mothers who used lorazepam 2–3 mg/day throughout pregnancy.176,178 It is not known if benzodiazepines cause behavioral abnormalities after prolonged intrauterine exposure.

Meprobamate

In one study, meprobamate ingestion during the first 42 days of pregnancy resulted in 8 of 66 exposed children having abnormalities, 5 of whom had cardiovascular malformations. Other large-scale investigations have not confirmed these findings.40 Alternative agents are advised, especially during the first trimester.

LITHIUM

Lithium use during the first trimester of pregnancy, particularly from the 3rd to 8th weeks, might increase the risk of cardiovascular abnormalities, including Ebstein’s anomaly.179–183 Anomalies of the external ear, ureters, CNS, and endocrine system, as well as macrosomia also have been reported.181,182 A case of polyhydranmios possibly caused by lithium-induced fetal polyuria has occurred.184 Of lithium-exposed pregnancies followed by the Lithium Registry, 39% delivered prematurely, 36% had macrosomia (>90th percentile body weight for gestational age), and there was an 8.3% perinatal mortality rate. Symptoms of lithium toxicity, including lethargy, hypotonia, poor sucking reflex, respiratory distress, cyanosis, arrhythmias, and thyroid depression with goiter and hypothermia, have been reported in newborns of women on long-term lithium therapy. Newborn blood concentrations were 0.6–1 mEq/L. Monitor maternal serum concentrations frequently during pregnancy because lithium clearance changes as pregnancy progresses.161,179–181

GASTROINTESTINAL DRUGS

ACID-PEPTIC THERAPY

Antacids

Available data do not suggest teratogenicity for commonly used antacids (aluminum, magnesium, calcium salts).185–187 Long-term administration, however, is not recommended because of the potential toxicity.188

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GASTROINTESTINAL DRUGS

Histamine H2-Receptor Antagonists

No reports link cimetidine or ranitidine to adverse pregnancy outcome.189,190 Manufacturer data on cimetidine use in 50 pregnant women showed no increased risk to the developing fetus, although time of exposure was not noted. No reported adverse effects on the course of delivery or on the neonate have been reported following the use of H2-antagonists as preanesthetic agents to prevent aspiration of acidic stomach contents.87,191,192

Sucralfate

No adequate studies exist on the use of sucralfate in pregnant women. Because little drug is absorbed, little risk is expected.87

ANTIEMETICS

Studies of women with hyperemesis gravidarum treated with metoclopramide, prochlorperazine, or placebo found no malformations or neonatal problems.171,193,194 However, a reanalysis of some early data questions the initial negative findings and suggests an increased risk of malformations with antinauseant phenothiazine use during the 4th to 10th weeks of gestation.163 Antihistamines used for pregnancy-induced emesis (dimenhydrinate, diphenhydramine, doxylamine, meclizine) are not considered teratogens.195,196 (See Antihistamines.)

GASTROINTESTINAL MOTILITY

Metoclopramide

There are no reports of adverse fetal outcome with use of metoclopramide during pregnancy; however, the drug has not been used widely in this setting.87,193 The use of metoclopramide during labor, anesthesia, or cesarean section to prevent acid aspiration has not resulted in adverse maternal, fetal, or neonatal outcomes.87,193

MISCELLANEOUS GASTROINTESTINAL DRUGS

Mesalamine Derivatives

Mesalamine derivatives including sulfasalazine have not been associated with teratogenic risk. It is not clear, however, if the drugs or the disease for which they are used (eg, Crohn’s disease) contribute adverse effects such as low birth weight.52,87,197–202 (See also Antimicrobial Drugs, Sulfonamides.)

HEMATOLOGIC DRUGS

COAGULANTS AND ANTICOAGULANTS

Heparin

Heparin has not been associated with an increased risk for structural or functional defects or with IUGR. Its large molecular size prevents it from crossing into the fetal circulation. Maternal thrombocytopenia and hemorrhage can occur.203–206

Low-Molecular-Weight Heparins

Low-molecular-weight heparins do not cross the placenta. Limited data do not suggest a teratogenic risk.207,208

Warfarin

Warfarin and related anticoagulants can produce the fetal warfarin syndrome or warfarin embryopathy. The critical period of risk appears to be between 6 and 12 weeks of gestation. Features include nasal hypoplasia, neonatal respiratory distress from upper airway obstruction, stippled

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HEMATOLOGIC DRUGS

epiphyses, IUGR, and different degrees of hypoplasia of the extremities. Eye abnormalities, including blindness, have also been reported. About one-third of exposed cases result in adverse pregnancy outcomes. One-half of these are spontaneous abortions or stillbirths, and the other half exhibit some type of congenital abnormality.2,152,203–206,209–212 A few cases of diaphragmatic malformation have been reported when warfarin was used early in pregnancy.213 CNS defects occur in about 3% of those exposed and appear to occur independent of the fetal warfarin syndrome. Critical periods of risk for CNS effects appear to be during the second and third trimesters.212 Warfarin also increases the risk for fetal and maternal hemorrhage, especially when used near term.203,209

HORMONAL DRUGS

ADRENAL HORMONES

Corticosteroid use during pregnancy can increase the risk for maternal gestational diabetes, hypertension, and excessive weight gain.118 Although corticosteroids do not represent a major teratogenic risk, there might be a small increased risk of oral clefts, which is consistent with animal studies.214 Large prospective investigations and cases of pregnant women with inflammatory bowel disease, asthma, or rheumatoid arthritis do not show an increase in spontaneous abortions or congenital defects in infants exposed prenatally. Some studies show decreased birth weight and increased rates of prematurity, although these effects might have been due to underlying maternal disease.118,215 Women who had status asthmaticus during pregnancy were more likely to have infants with low birth weight or IUGR.215 A placental enzyme (11β-OH-dehydrogenase) inactivates certain corticosteroids, leading to low fetal concentrations. This inactivation is greater with hydrocortisone, prednisone, or prednisolone than with betamethasone or beclomethasone. The fetal liver is relatively ineffective in converting prednisone into the active prednisolone, even at term. Therefore, prednisone or hydrocortisone might be preferred during pregnancy. When appropriate, corticosteroids can be administered by aerosol inhalation or intrasynovial injection to minimize systemic availability.215 Neonates born to women taking long-term corticosteroids should be monitored for adrenal insufficiency, although this rarely occurs. Maternal betamethasone therapy is used to prevent respiratory distress in infants born between 28 and 34 weeks of gestation, with no apparent adverse effects.216,217 No differences in psychological or physical development, pulmonary function, ophthalmologic findings, or neurologic function were noted between exposed and nonexposed children 10 to 12 yr old.218 However, there was a significantly increased frequency of hospitalizations for infections during early childhood in the exposed group.

ANDROGENS

Androgen (eg, testosterone, methyltestosterone, and danazol) exposure during the first week of gestation can cause masculinization of the female embryo. Clitoromegaly, with or without fusion of the labia minora, can occur. In some cases, a urogenital sinus opening at the base of the clitoris has been observed. The extent of the defect is correlated with the time of exposure and dosage.2,219,220 Although danazol dosages <400 mg/day can carry a lower risk, virilization was observed with 200 mg/day in one case.220 Because differentiation of external genitalia occurs 8–12 weeks postconception, exposure beyond the 12th week of gestation is expected to produce only clitoral hypertrophy. If danazol is discontinued by the 8th week of gestation, the risk is substantially lower. Internal genitalia are unaffected because they are not androgen responsive. Male fetuses do not appear to be adversely affected.

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HORMONAL DRUGS

ANTIDIABETIC DRUGS

Maternal diabetes increases the rate of malformations and perinatal mortality.\textsuperscript{221,222} Pregnancy in 20 women exposed to an oral hypoglycemic (16 to a sulfonylurea, 3 to a biguanide, 1 unknown) during organogenesis resulted in a higher proportion of major and minor malformations than a similar, unexposed control group.\textsuperscript{222} A retrospective case review did not find a similar increase in malformations.\textsuperscript{223} Malformations reported included auricular, vertebral, cardiac, neural tube, and limb defects.\textsuperscript{222–224} Complicating the issue is a study that shows \textit{glyburide} does not cross the human placenta \textit{ex vivo} in appreciable amounts.\textsuperscript{225} Sulfonylureas (eg, \textit{chlorpropamide, glyburide}, and \textit{glipizide}) and biguanides (eg, \textit{metformin}) can cause prolonged hypoglycemia or hyperinsulinism in the newborn.\textsuperscript{2,40,152,222,226} Pregnant diabetics should be treated with \textit{insulin}.

FEMALE SEX HORMONES

\textit{Clomiphene}

Clomiphene-induced ovulation results in an increased frequency of multiple ovulations (primarily twinning), delayed follicular rupture, ectopic pregnancy, and female fetuses. There also is an increase in abnormal karyotypes in oocytes and abortuses. The high proportion of abnormal karyotypes might be related to the increase in spontaneous abortions and low pregnancy rate noted after ovulation induction and in vitro fertilization.\textsuperscript{227,228} The rate of birth defects reported after ovulation induction in most studies has been within the expected range.\textsuperscript{227,229,230} A pooled analysis of available studies concluded that, if clomiphene causes neural tube defects, the elevated risk is no greater than twice the baseline rate.\textsuperscript{231}

\textit{Diethylstilbestrol (DES)}

Daughters of DES-exposed mothers have an increased frequency of vaginal adenosis; structural defects of the cervix, vagina, uterus, and fallopian tubes; and reproductive complications such as infertility, spontaneous abortion, ectopic pregnancy, premature deliveries, and perinatal deaths.\textsuperscript{2,232–235} Two large cohort studies do not confirm the relation of DES to clear cell carcinoma. Early studies showing such an association were retrospective and had serious methodologic flaws.\textsuperscript{236} If an association exists, the risk of developing this cancer in exposed females is very low. Data on DES-exposed males suggest an increased risk for infertility, various urogenital abnormalities, and testicular cancer.\textsuperscript{237}

\textit{Progestins}

High doses of progestins (eg, \textit{medroxyprogesterone, norethindrone}, and \textit{norethynodrel}) during pregnancy can cause masculinization of female external genitalia (clitoral hypertrophy or labioscrotal fusion). Hypospadias has occurred in males.\textsuperscript{238,239} This most likely occurs during the period of external genitalia development (8–12 weeks postconception).\textsuperscript{2,40,240–242} Doses used in oral or implantable contraceptives do not appear to carry these risks.\textsuperscript{243}

THYROID AND ANTITHYROID DRUGS

\textit{Propylthiouracil} and \textit{methimazole} are not considered major human teratogens.\textsuperscript{244} However, both drugs were associated with neonatal goiter in early reports, possibly because of the use of iodine in addition to unnecessarily high dosages of antithyroid medication. However, even conservative management does not completely eliminate the risk of congenital goiter. Transient neonatal hypothyroidism and neonatal thyrotoxicosis can occur. Hypothyroidism can be prevented by discontinuing the drug 4–6 weeks before delivery if the mother has remained euthyroid. Monitor in-
HORMONAL DRUGS

Fants for thyrotoxicosis because it can be masked by the antithyroid agent. Cases of infants with ulcer-like midline scalp defects have been associated with maternal methimazole. Another report, however, did not confirm an association of methimazole with skin defects.245 Mothers with moderate to severe hyperthyroidism not receiving treatment have an increased risk of complications including toxemia, small-for-gestational-age babies, and neonatal morbidity.246–249 Excessive iodine use during pregnancy can cause congenital goiter and hypothyroidism. Goiters might be large enough to cause tracheal compression and interfere with delivery. Severe maternal iodine deficiency produces hypothyroidism and cretinism.246,247

No adverse effects have been reported after inadvertent exposure to 131I before 10 weeks of gestation. After 10 weeks of gestation, the fetal thyroid actively concentrates iodine; any radioactive iodine ingested by the mother will cross the placenta and destroy fetal thyroid tissue.2,246

Thyroid hormones (levothyroxine, liothyronine) may be used during pregnancy for the treatment of hypothyroidism.171

MISCELLANEOUS HORMONAL AGENTS

Oxytocin

Oxytocin given for induction of labor can cause tetanic uterine contractions, resulting in decreased uterine blood flow and fetal distress. The risk of neonatal hyperbilirubinemia is increased 1.6-fold after oxytocin-induced labor compared with spontaneous labor, and neonatal jaundice has occurred.249

Prostaglandins

Misoprostol has uterotonic effects and should not be used during pregnancy. In 56 pregnant women who requested first-trimester abortions, 1 or 2 oral doses of misoprostol 400 g resulted in spontaneous fetal expulsions.250 Twenty-five women had uterine bleeding. High doses of misoprostol in early pregnancy, such as those used to induce abortion, cause defects. Anomalies reported include paralysis of cranial nerves VI and VII and limb, orofacial, and musculoskeletal defects.251–254 Dinoprostone, when administered as vaginal tablets to mildly hypertensive women at term to induce labor, produces significant decreases in the percentage of time occupied by fetal body and breathing movements 3–4 hr later.255 Fetal outcome is normal. Dinoprostone might shorten labor more than oxytocin.256,257

RENALEND ELECTROLYTES

DIURETICS

Diuretics should be used with great caution during pregnancy because they can decrease maternal intravascular volume and consequently diminish uteroplacental perfusion, thereby compromising fetal oxygenation. This effect is most rapid and severe with loop diuretics (eg, bumetanide, furosemide, and torsemide). IV furosemide administration to the pregnant woman has enabled ultrasonic imaging of the fetal bladder because of increased fetal urine output. Thiazide diuretic use is not associated with birth defects; however, use during late pregnancy can produce neonatal hypoglycemia, hyponatremia, hypokalemia, and thrombocytopenia.121,128,129,258

ELECTROLYTES

Long-term infusion of magnesium sulfate during the second trimester occasionally causes bone abnormalities and dental enamel hypoplasia.259,260 However, infusions between 24 and 34 weeks of gestation in women with preterm labor generally do not cause adverse neonatal outcomes.261

(continued)
RENAL AND ELECTROLYTES

Use of magnesium sulfate as a tocolytic agent near term can cause dosage-related neonatal hypotonia, hyporeflexia, respiratory distress, hypocalcemia, and hypermagnesemia.\textsuperscript{259,262} Magnesium also is used for prophylaxis of seizures due to eclampsia.\textsuperscript{263} Serial serum magnesium concentrations should be used to guide therapy.

RESPIRATORY DRUGS

ANTIALLERGICS

Cromolyn sodium use during pregnancy is not known to have any teratogenic effects. A prescription event monitoring study of nedocromil in the United Kingdom identified 79 exposed pregnancies.\textsuperscript{264} Routine follow-up showed no abnormalities among the 33 infants with first-trimester exposure.

ANTIASTHMATICS

\(\beta\)-Adrenergic Agonists

Aerosol inhalation of metaproterenol, albuterol, isoetharine, or terbutaline is considered safe during pregnancy. Less is known about the oral use of these products, but experience suggests no adverse outcome. In one series, salmeterol was taken during the first trimester during 64 pregnancies and only during the second and third trimesters in 3 pregnancies.\textsuperscript{265} The 67 pregnancies resulted in 50 live births, 6 spontaneous abortions, 2 ectopic pregnancies, 4 elective terminations, and 5 unknown outcomes. The small number of fetuses exposed to salmeterol during this study precludes any definitive conclusions about its safety. Albuterol, terbutaline, and ritodrine have been given in large IV or PO doses for their tocolytic effects in the third trimester without permanent harm to the fetus, although hypertension, hypoglycemia, hypokalemia, and hypocalcemia have been reported.\textsuperscript{262} Ephedrine use during labor can alter neonatal sleep patterns.\textsuperscript{170}

Inhaled Corticosteroids

This route is preferred during pregnancy to minimize fetal exposure. (See Corticosteroids.)

Theophylline

Maternal theophylline pharmacokinetics can change during the second and third trimesters, with decreased theophylline protein binding, decreased nonrenal clearance, and increased renal clearance.\textsuperscript{266} \(V_d\) and half-life increase during the third trimester.\textsuperscript{266} Pregnant women should have serum theophylline concentrations monitored frequently for dosage adjustments. No adverse fetal effects from long-term theophylline use during pregnancy are known.\textsuperscript{267} Theophylline toxicity occurred in three newborns whose mothers received theophylline or aminophylline in late pregnancy and during labor. Symptoms included jitteriness, vomiting, and tachycardia, all of which resolved.\textsuperscript{40}

ANTIHISTAMINES

The relative risk of malformations for first-generation antihistamines (eg, chlorpheniramine, doxylamine) is not statistically significant. Despite reports implicating meclizine and Bendectin (doxylamine and pyridoxine with or without dicyclomine) as teratogens, large-scale studies and two meta-analyses have shown no association between antihistamines and fetal malformation.\textsuperscript{3,40,268,269} Limited data from studies of astemizole, hydroxyzine, and cetirizine have not shown a significant teratogenic risk.\textsuperscript{269,270} Dimenhydrinate might have an oxytocic

(continued)
RESPIRATORY DRUGS

Effect on the term uterus, causing shortened labor. The same concerns as with oxytocin (hyperstimulation and the possibility of uterine rupture) apply.189 (See Antiemetics.)

COUGH AND COLD

Use of sympathomimetics (eg, pseudoephedrine) for treatment of nasal congestion can cause increased fetal activity and fetal tachycardia. Systemically administered sympathomimetics should be avoided in patients with hypertension or eclampsia or situations in which there is poor fetal cardiac reserve.40,271

MISCELLANEOUS DRUGS

ANESTHETICS, LOCAL

Maternal exposure to local anesthetics during the first 4 months of pregnancy does not cause fetal malformations.169,170,272 However, local anesthetics have resulted in fetal and neonatal CNS and myocardial depression and fetal hyperthermia after maternal use during labor. Fetal bradycardia can occur after paracervical blocks. Epidural administration of bupivacaine or lidocaine might be the safest method for obstetric analgesia. Epidural lidocaine was shown to decrease neonatal neurobehavioral performance, but the effect was short lived and of minimal clinical consequence.169 High-concentration (0.75%) bupivacaine is not recommended for use in obstetrics because of the profound neonatal myocardial depression and prolonged difficult resuscitation that follow accidental intravascular injection.169,272 Lower concentrations are popular because of the high quality of analgesia with minimal degree of motor block.169 Mepivacaine should not be used in obstetric anesthesia because the neonate cannot metabolize it.169,170

ERGOT ALKALOIDS

The use of ergonovine or methylergonovine before delivery carries the same risk of uterine stimulation as oxytocin.13,262 (See Hormonal Drugs, Oxytocin.)

PODOPHYLLIN

Podophyllotoxins are antimitotic agents, and their use during pregnancy is contraindicated. Maternal oral use during the 5th through 9th weeks of gestation might be associated with congenital malformation. High doses used topically at the 34th week resulted in severe maternal toxicity and a stillborn fetus. There are several cases in which topical podophyllin use during pregnancy produced no adverse outcome.171,273–276

RETINOIDS

Etretinate

Etretinate is a known teratogen in humans. Fetal abnormalities include facial dysmoria; syndactylies; absence of terminal phalanges; neural tube closure defects; malformations of the hip, ankle, and forearm; low-set ears; high palate; decreased cranial volume; and alterations of the cervical vertebrae and skull.277 Etretinate accumulates in adipose tissue with repeated administration and has been detected in the serum of some patients up to 3 yr after discontinuing therapy. The importance of this, relative to the risk of teratogenicity, is unknown. An effective form of contraception must be used for at least 1 month before etretinate therapy, during therapy, and for an indefinite time after...
*MISCELLANEOUS DRUGS*

therapy, some say for at least 2 yr. Unpublished data from the manufacturer indicate suspected teratogenicity in 8 of 95 fetuses reportedly exposed 1–24 months after drug discontinuation.279

**Isotretinoin**

Use during pregnancy causes defects of the CNS, heart, external ear, and thymus. Other reported malformations include cleft palate, microphthalmia, micrognathia, facial dysmorphia, and limb reduction defects. Infants might exhibit hearing and visual impairments and mental retardation. One report estimates the relative risk for major birth defects from isotretinoin to be 25.6. Data suggest that the risk is substantial even when exposure to isotretinoin is brief or dosage is low. The risk for spontaneous abortion is also increased. In an analysis of cases reported to the manufacturer, 28% of exposed fetuses were malformed. Exposures before day 14 of gestation resulted in the same rates of malformation and spontaneous/missed abortion as exposures between days 14 and 83. An effective form of contraception must be used for at least 1 month after discontinuation of isotretinoin and before conceiving.

**Tretinoin**

Topical application of tretinoin results in absorption of vitamin A with equivalent activity less than that of a prenatal vitamin. Therefore, teratogenicity of this retinoid, although theoretically possible, is unlikely when used topically as directed. Studies of first trimester use of topical retinoids and a review of cases of malformations typical of retinoids do not implicate topical retinoids in malformations. Of note, two cases of malformations similar to those of isotretinoin have been reported after first-trimester tretinoin exposure.

**Vitamin A**

Retinol or retinyl esters (but not beta-carotene) in toxic doses are teratogenic in experimental animals, producing defects in almost all organ systems. Human data on vitamin A teratogenicity consist of only a few anecdotal reports and epidemiologic studies. Epidemiologic studies show that doses of vitamin A contained in prenatal vitamin supplements (ie, ≤10,000 IU/day) do not increase the risk of fetal malformation. However, because of methodologic flaws and incomplete data, the degree of risk with higher daily dosages of vitamin A is unclear. Defects observed when mothers ingested ≥25,000 IU/day of vitamin A during pregnancy include craniofacial, CNS, cardiac, urinary, vertebral, and other skeletal malformations.

**VACCINES**

See Immunization, page 996, for information regarding vaccination during pregnancy.

**VAGINAL SPERMICIDES**

Use of vaginal spermicides was associated with major congenital anomalies in two retrospective analyses, but there were many flaws in the studies, casting doubt on the results. Subsequent studies, including a meta-analysis of nine studies, did not implicate these agents as teratogens.

**VITAMIN D**

Supplementation with vitamin D in pregnant women with poor dietary intake and little exposure to light is not associated with adverse outcomes. Excessive use has been associated with an idiopathic hypercalcemic syndrome including cardiovascular malformation, abnormal bone mineralization, elfin facies, mental retardation, hypercalcemia, and nephrocalcinosis. Definitive conclusions await further investigation.

(continued)
MISCELLANEOUS DRUGS

DRUGS FOR NONMEDICAL USE

Alcohol

Animal studies show that alcohol and acetaldehyde exposure in utero results in morphologic changes in the structure and protein and endocrine content of the CNS. As many as 5% of human congenital anomalies might be due to maternal alcohol consumption, and it might be responsible for 10% of all cases of mental retardation. (Alcohol is the most frequent recognizable cause of mental retardation.) A 2-fold increase in spontaneous abortion was noted among women who drank one to two drinks daily for the first 2 months of pregnancy; the rate was higher in those who drank more than two drinks daily. Moderate drinking (1–13 fl oz of absolute alcohol per week) results in an increase in some of the features of the "fetal alcohol syndrome" (FAS), including IUGR. Chronic heavy alcohol consumption can cause full-blown FAS in up to 50% of children exposed in utero. Binge drinking has been associated with behavioral defects. Features of FAS include IUGR, microcephaly, postnatal growth deficiency, developmental delay, mental retardation, and craniofacial anomalies. Joint, limb, cardiac, ocular, brain, urogenital defects, and eustachian tube dysfunction also can occur. Neonates can have withdrawal symptoms similar to those of adults. Alcohol-related birth defects can occur at rates higher than those of full-blown FAS, and CNS dysfunction can occur with lower alcohol exposure than that required to produce the FAS. One investigator proposes that the threshold dosage for alcohol teratogenesis is 1 fl oz of absolute alcohol per day. Alcohol consumption during pregnancy should be avoided because the minimum dosage that can produce adverse fetal effects has not been established.

Amphetamines

Data on the effect of prenatal amphetamines, prescribed and abused, are conflicting so no consistent pattern of abnormalities has emerged. Most studies found no increase in severe congenital malformations, but others reported biliary atresia, congenital heart disease, and eye and CNS defects. Reports of decreased birth weight and length, head circumference, and IUGR might reflect poor maternal nutrition, but use of other drugs and alcohol might have confounded those findings. Investigations with term neonates exposed antenatally to methamphetamine, with or without cocaine, document an increased prevalence of prematurity, IUGR, altered behavioral patterns (abnormal sleep patterns, poor feeding, tremors, and hypertonia), and the presence of cerebral injury as detected by ultrasonography. (See also Cocaine.)

Caffeine

Caffeine is not suspected of causing fetal malformations. Data suggest that caffeine use before and during pregnancy increases the rate of spontaneous abortion. A small reduction in birth weight can occur when caffeine consumption during pregnancy exceeds 300 mg/day, although cigarette smoking might contribute. There are conflicting data about whether 400–600 mg/day of caffeine increases the risk of miscarriage, stillbirth, or prematurity. One study showed that rates of infant central and obstructive apnea positively correlated with increasing maternal caffeine consumption. In one study of long-term outcome, no adverse effects were observed on the IQ of children exposed in utero to caffeine.

Cocaine

Maternal cocaine use during pregnancy can cause adverse fetal and maternal outcomes. Although most data relate to use near term, preliminary findings suggest some risks from exposure early in pregnancy. Poor health care and nutrition, a high infection rate, and frequent concomitant drug (eg, narcotics, nicotine, or marijuana) and alcohol use complicate risk evaluation. A meta-analysis (continued)
DRUGS FOR NONMEDICAL USE

found no significant differences in adverse fetal effects between cocaine users and polydrug users.338 Maternal cocaine use results in an increase in the frequency of spontaneous abortion, placental hemorrhage, abruptio placentae, and stillbirth. There also is an increase in frequency of prematurity and premature rupture of the membranes. A few cases of precipitate delivery have been reported, as have cases of precipitate rupture of ectopic pregnancy. Fetal genitourinary malformations and intrauterine death have been reported when mothers used cocaine.338–343 Cocaine most likely disrupts morphogenesis by vasoconstriction of uterine and fetal circulation. Interruption of calcium metabolism might contribute to hydronephrosis.340,342,344,345 Infants can experience a withdrawal syndrome consisting of jitteriness, tremor, hyperreflexia, hypertonia, irritability, high-pitched crying, frantic sucking, poor feeding, tachypnea, abnormal sleep patterns, vomiting, or loose stools. Neurologic signs of toxicity, including seizures, also have been reported. Exposed infants have an increased frequency of abnormal pneumograms, respiratory distress, or other cardiorespiratory abnormalities. Some investigators feel these abnormalities can predispose infants to sudden infant death syndrome (SIDS). Retrospective population-based studies found that the risk of SIDS in cocaine-exposed infants was higher than that of infants whose mothers did not abuse any drugs; however, several confounding independent risks for SIDS were not controlled.346,347 Several reports of neonatal or fetal cerebral infarction or intracranial hemorrhage exist.338,339,344,348–376 Long-term behavioral abnormalities have been reported after in utero exposure to cocaine.377

Heroin

Poor nutrition and health care, lack of prenatal care, high infection rates, and frequent concomitant nicotine, nonmedical drug, and alcohol use complicate risk assessment. No specific pattern of fetal malformation has been noted. Some studies have associated maternal heroin use with a decrease in birth weight and length, reduced head circumference, small-for-gestational-age infants, low Apgar scores, meconium staining, neonatal respiratory distress, jaundice, and increased neonatal mortality. A narcotic withdrawal syndrome occurs frequently, usually within the first 24–48 hr after birth, although it can be delayed for as long as 6 days. The symptoms, which can persist for as long as 20 days, include irritability; feeding and sleeping problems; hyperactivity; and excessive sneezing, yawning, vomiting, mucous secretion, sweating, and face scratching. Other withdrawal symptoms such as increased muscle tone, vague autonomic nervous system symptoms, tremulousness, high-pitched crying, frantic and uncoordinated sucking, and seizures can occur in neonates born to narcotic-addicted women and nonaddicted women who use narcotics near term. The frequency of withdrawal is directly related to the daily dosage and duration of maternal heroin use. The results of studies evaluating development show conflicting results ranging from no effect to problems with behavior and perceptual and organizational abilities.40,378 One retrospective, population-based study found that the relative risk of SIDS was 15.5 in infants of mothers who abused illicit opiates during all or part of pregnancy.346

Marijuana

Marijuana smoke contains carbon monoxide and can constrict uterine and placental vessels, resulting in fetal hypoxia and decreased nutrient supply (growth retardation).374,378 Marijuana use during pregnancy did not result in any specific pattern of malformation, but exposures were confounded by nicotine and other nonmedical drug and alcohol use.374,376,378–381

Phencyclidine

Long-term phencyclidine use during pregnancy, especially near term, can produce neonatal withdrawal symptoms of hypertonicity, occasional darting eye movements, lethargy, and coarse flappy...
tremors after stimulation. These infants have a marked increase in lability of behavioral states and poor consolability.\textsuperscript{382,383} Long-term behavioral abnormalities have been reported in some studies but not in others.\textsuperscript{374,382,384,385}

**Tobacco**

Cigarette smoke contains numerous toxins, including carbon monoxide and nicotine, that are probably responsible for reported fetal hypoxia because of decreased oxygen exchange and placental vasoconstriction.\textsuperscript{386,387} Smoking during pregnancy increases rates of IUGR or low birth weight (5\% decrease/pack smoked daily), prematurity, spontaneous abortion, neonatal and postnatal deaths, abruptio placentae, premature rupture of membranes, and placenta previa.\textsuperscript{2,226,387–391} Women exposed to second-hand smoke also can be at risk for delivering a growth-retarded or preterm infant.\textsuperscript{388} If a woman stops smoking by the 20th week of pregnancy, the risk of a low birth weight infant is similar to that of the general population.\textsuperscript{386} Cigarette smoking alters the placental arterial linings and accelerates placental senescence, which might explain the placental complications of smoking.\textsuperscript{387,390–392} One study involving 17,152 infants (15.7\% of the mothers were smokers) found an increased frequency of minor malformations in infants of mothers aged \geq 35 yr who smoked.\textsuperscript{393} Some data suggest infants of smokers have increased nervous system excitation, hypertonicity, and altered breathing patterns with an increased rate of central apnea.\textsuperscript{336,394} There also might be long-term mental and physical effects.\textsuperscript{395} Some studies suggest an increased risk of childhood acute lymphocytic and lymphoblastic leukemias and lymphoma in those whose fathers and/or mothers smoked before or during pregnancy.\textsuperscript{396,397} Tobacco chewing during pregnancy also greatly increases the rate of stillbirth, IUGR, and prematurity.\textsuperscript{388}

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Drugs and Breastfeeding

Philip O. Anderson

With the increasing recognition of the benefits of breastfeeding, the clinician often must weigh the benefits against the risks of drug therapy in lactating women. The physicochemical, pharmacokinetic, and clinical factors involved with drug use in nursing women have been described. These factors are summarized as follows.

I PHYSICOCHEMICAL FACTORS

Small water-soluble nonelectrolytes pass into breastmilk by simple diffusion through pores in the mammary epithelial membrane that separates plasma from milk. Equilibration between the two fluids is rapid, and milk concentrations of drugs approximate plasma concentrations. For larger molecules, only the lipid-soluble, nonionized forms pass through the membrane by crossing the cell wall and diffusing across the interior of the cell to reach the milk. Because the pH of milk is generally lower than that of plasma, milk can act as an “ion trap” for basic drugs. At equilibrium, these compounds can be concentrated in milk relative to plasma. Conversely, acidic drugs are inhibited from entering milk. The pKa of weak electrolytes is an important determinant of their equilibrium concentration in milk.

Protein binding also is an important determinant because plasma proteins bind drugs much more avidly than do milk proteins. Highly protein-bound drugs do not pass into milk in high concentrations. Lipid solubility favors passage of some drugs into milk because the fat component of milk can concentrate lipid-soluble drugs. However, because milk contains only 3 to 5% fat, its capacity for concentrating drugs is limited. Active or facilitated transport of drugs into breastmilk might occur in humans, but it is rare.

I PHARMACOKINETIC FACTORS

Because the breast is periodically emptied by the nursing infant and refilled with newly formed milk, equilibrium between plasma and milk is rarely reached. Therefore, the rate of drug passage from plasma into milk is important in determining the concentration of a drug in milk. Factors favoring rapid passage into milk are high lipid solubility and low molecular weight.

Passage of drugs between plasma and milk occurs in both directions. When the concentration of nonionized free drug is higher in milk than in plasma, net transfer of drug from milk to plasma occurs. Thus, the maneuver of pumping and discarding milk does not appreciably hasten the elimination of most drugs from milk and does not have a marked effect on overall clearance of the drug from the mother’s body.

I METHODS OF EXPRESSING THE EXTENT OF PASSAGE

The ratio of concentrations of a drug in milk and plasma (the milk/plasma, or M/P, ratio) often has been used as a measure of a drug’s passage into breastmilk. However, the M/P ratio has shortcomings that make it meaningless as a measure of
drug safety during nursing. There is no standard method of calculating the value, and the value is not constant, as often calculated, but changes with the time after the dose and the number of doses given. It also does not take into account the potential toxicity of the drug.

The percentage of the maternal dosage that is excreted into milk also is used to express the extent of passage. This value alone is not predictive of safety in a nursing infant but can be used to calculate the actual dosage received by the infant. Usually a weight-adjusted (ie, mg/kg) infant dosage <10% of the mother’s is considered safe; of 205 drugs studied, 87% of drugs fell into this category. The likelihood of an adverse effect in the infant increases markedly in those few drugs (about 3%) that have a dosage in milk that is >25% of the maternal weight-adjusted dosage.2

Drug clearance can be a useful factor for identifying drugs that can accumulate in infants and thereby have a pharmacologic effect.3 Drugs with an adult total body clearance of ≥0.3 L/hr/kg and that have no active metabolites are unlikely to have pharmacologic effects in a nursing infant because they are rapidly eliminated from the mother and infant.

All of the above methods fall short of providing a complete assessment of the safety of a drug during breastfeeding in a specific mother–infant pair. Several additional factors must be considered.

## CLINICAL CONSIDERATIONS

Factors that should be considered when determining the advisability of using a particular drug in a nursing mother are the potential acute toxicity of the drug, dosage and duration of therapy, age of the infant (<2 months are the most susceptible), quantity of milk consumed, experience with the drug in infants, oral absorption of the drug by the infant, potential long-term effects, and possible interference with lactation.1,4

A stepwise approach to using medications in breastfeeding women can be followed to minimize infant exposure to medications in milk.1 Starting from the strategies that are least disruptive to nursing and progressing to those that are most disruptive, the prescriber can consider the following steps: withhold the drug; delay drug therapy temporarily; choose drugs that pass poorly into milk; use alternative routes of administration (eg, topical, inhalation); avoid nursing at times of peak milk levels; administer the drug before the infant’s longest sleep period; withhold breastfeeding temporarily; and, infrequently, discontinue breastfeeding.

Another consideration is non–dose-related adverse effects such as allergic reactions and some hemolytic anemias; however, these reactions are relatively uncommon. GI intolerance caused by antimicrobial agents in breastmilk can occur whether or not the drugs are absorbed by the infant. Antimicrobial agents are among the most commonly used maternal medications during nursing and, although serious side effects are rare, diarrhea might occur in up to 12% of infants.5 Disruption of the infant’s GI flora is uncommon but occasionally leads to thrush and rarely to pseudomembranous colitis.6 Severe diarrhea or blood in the infant’s stool during maternal antimicrobial use is an indication to stop nursing and seek medical attention.

Although the above considerations are important, follow-up of mothers who took medications while breastfeeding their infants has shown that serious side ef-
Effects are uncommon. Nursing seldom needs to be completely discontinued because of concern of acute toxicity from maternal drug therapy.

The following table contains information on the use of specific drugs during nursing. The risks are assessed and alternatives are presented based on the principles discussed. Information in the table is from reference 1 unless noted otherwise.

### DRUGS AND BREASTFEEDING

#### ANALGESICS AND ANTI-INFLAMMATORY DRUGS

#### ANTIMIGRAINE DRUGS

**Ergotamine**
When given daily for 6 days postpartum, ergotamine did not affect lactation or infant weight in one study; however, the excretion of ergotamine into milk during lactation has not been studied. Avoid its use during lactation because older ergot preparations have produced toxicity in infants.

**Sumatriptan**
Minimally excreted in milk, sumatriptan has poor oral absorption by the infant. It poses little risk during breastfeeding.

#### NONSTEROIDAL ANTI-INFLAMMATORY DRUGS

**Acetaminophen**
The amount of acetaminophen excreted into milk is small. Acetaminophen is a good analgesic choice during nursing.

**Nonsteroidal Anti-inflammatory Drugs**
Amounts of most NSAIDs in milk are low because they are weak acids that are extensively plasma protein bound. However, short-acting agents are preferred, particularly in the case of breastfed neonates. Some agents have active metabolites (eg, sulindac) or glucuronide metabolites (eg, salicylate, fenoprofen, and ketoprofen) that can add to infant intake. Because of the increased likelihood of accumulation, avoid long-acting agents such as diflunisal, naproxen, piroxicam, and sulindac in mothers of neonates, although amounts of piroxicam in milk are low. Naproxen caused prolonged bleeding time, thrombocytopenia, and acute anemia in one 7-day-old infant, and possibly drowsiness and vomiting in others. The more toxic NSAIDs such as mefenamic acid and indomethacin also should be avoided, although recent studies on indomethacin indicate that it might not be contraindicated.

**Ketorolac** is contraindicated during nursing. **Diclofenac** was not detected in milk after a single dose of 50 mg IM, or 100 mg/day for 1 week, and the amount of tolmetin in one woman’s milk was low. **Ibuprofen** and **flurbiprofen** have the best documentation of safety during breastfeeding; the dose of ibuprofen that an infant receives in milk is <0.001% of the mother’s dosage, and flurbiprofen concentrations are low to undetectable after dosages up to 50 mg tid.

**Salicylates**
Salicylate enters milk in a low concentration relative to that in plasma, although its glucuronide metabolite increases the overall infant dosage. Doses >1 g yield markedly higher salicylate concentrations in milk and can result in high infant serum concentrations. One case of thrombocytopenic purpura from aspirin in breastmilk (confirmed by rechallenge) was reported in a 5-month-old infant. The risk of Reye’s syndrome caused by salicylate in milk is unknown. If aspirin is taken occasionally, avoid breastfeeding for 1–2 hr after a dose to minimize antiplatelet effects. NSAIDs such as ibuprofen are preferred to aspirin for long-term therapy.

(continued)
ANALGESICS AND ANTI-INFLAMMATORY DRUGS

OTHER ANTI-INFLAMMATORY DRUGS (See also Antimalarials.)

Gold

During maternal administration of aurothioglucose and gold sodium thiomalate, gold was detected in the blood and urine of some nursing infants. The weight-adjusted infant dosage might be greater than the maternal dosage, but the amount of gold that infants absorb orally is not known. Sufficient amounts are absorbed, however, to potentially cause adverse effects. Gold therapy is a reason to very carefully monitor the breastfed infant and might be a reason for withholding breastfeeding.1,12

Penicillamine

Penicillamine was used during 3 months of breastfeeding in two women who nursed 3 infants without harm, but it is not recommended.13,14

OPIOIDS

Neonates are particularly susceptible to narcotics in breastmilk.4 Postpartum maternal opioids (oral codeine or propoxyphene with or without prior IM meperidine) might be a causative factor in episodes of apnea, bradycardia, and cyanosis during the first week of life. Avoid maternal narcotics when the breastfed neonate has experienced such an episode. Although single analgesic doses of most narcotics are excreted into milk in small amounts, infant drowsiness caused by repeated administration of postpartum oral narcotics in milk is more prevalent than commonly thought—about 20% in one study.5 Drowsiness is dose related and can be severe with the maximum dosage. Limiting oral dosage to 1 tablet (eg, codeine 30 mg, hydrocodone 5 mg, or oxycodone 5 mg) q 4 hr is advisable; analgesia can be supplemented with additional acetaminophen or ibuprofen.

Meperidine

Meperidine is particularly likely to interfere with infant nursing behavior when given during labor.15,16 Furthermore, repeated postpartum meperidine doses, including patient-controlled analgesia, cause diminished alertness and orientation in breastfed neonates compared with equivalent doses of morphine.1,17 Meperidine should be avoided during labor and nursing, although a single small dose for anesthesia or conscious sedation usually does not cause problems in older breastfed infants.18

Morphine

Morphine 10–15 mg in single parenteral doses produces only low concentrations in milk, but repeated doses can result in drug accumulation in infant serum to near therapeutic concentrations. Morphine glucuronides in milk contribute an additional 50 to 100% to the infant. Epidural administration and patient-controlled analgesia cause fewer effects in an infant than IV administration and are preferred.4,17

Fentanyl and Sufentanil

IV or epidural fentanyl, alfentanil, and sufentanil produce low milk levels.19,20 In addition, these drugs have poor oral bioavailability, so they are good choices for maternal analgesia during nursing.

Narcotic Partial Agonists

IV narcotic agonist/antagonists given during labor can interfere with establishment of lactation.15,16 Despite breastfeeding and relatively high infant serum drug levels, mild withdrawal occurred in the neonate of a mother taking buprenorphine during pregnancy and postpartum for heroin addiction.21 Oral buprenorphine for narcotic abstinence appears to have little impact on the (continued)
ANALGESICS AND ANTI-INFLAMMATORY DRUGS

Breastfed infant. Butorphanol and nalbuphine concentrations in milk are low, and oral bioavailability in the infant should be low.Only about 0.1% of the maternal dose of tramadol is found in milk according to the manufacturer.

ANTIMICROBIAL DRUGS

AMINOGLYCOSIDES

Systemic effects of amikacin, gentamicin, streptomycin, tobramycin, and other aminoglycosides are unlikely in infants because of the small amounts in milk and poor oral absorption; however, observe infants for disruptions of the GI flora such as diarrhea and thrush.

ANTIFUNGAL DRUGS

Amphotericin B and nystatin are virtually unabsorbed orally, and the latter is frequently used orally for treating thrush in infants; therefore, both are safe for use in nursing mothers, including topical application to the nipples. Likewise, clotrimazole has poor oral bioavailability and has been used orally in infants with thrush, sometimes successfully after nystatin has failed. Miconazole has efficacy and safety similar to clotrimazole. These two imidazoles are preferred for topical or vaginal application during nursing. Fluconazole amounts in milk are much less than the dosage prescribed for infants and can be used for recalcitrant Candida infections given to the mother and infant simultaneously. Ketoconazole concentrations in milk are low, but it is best avoided in nursing mothers orally or topically to the nipples because of its oral absorption, occasional hepatotoxicity, and the availability of safer alternatives. Other imidazole antifungals have not been studied. Gentian violet is potentially toxic (toxic to mucous membranes, potential tattooing of the skin, and carcinogenic and mutagenic in rodents) and is best avoided topically on the nipples or in the infant’s mouth.

ANTIMYCOBACTERIAL DRUGS

Clofazimine

Clofazimine is excreted into milk, reportedly coloring it bright pink. Infants receive about 15 to 30% of the maternal mg/kg dose. Breastfed infants can develop the typical skin discoloration. The skin color returned to normal 5 months after the end of maternal therapy in one infant.

Antituberculars

Antituberculars pass into milk in small quantities. Use caution in nursing mothers because many of these drugs can cause hepatic damage. However, inadequate maternal therapy poses a much greater risk to the infant than the drugs in milk. The mother may take single daily doses of many of these drugs at bedtime and substitute a bottle for a nighttime feeding to minimize infant exposure. Isoniazid is excreted into milk in amounts that are less than those given to treat an infant. Pyrazinamide concentrations in milk in one woman were low and would give the baby less than a therapeutic dosage. Rifampin has not been well studied, but amounts in milk are small. Cycloserine is excreted in small amounts, and no adverse reactions have been reported in infants. Ethambutol has not been adequately studied.

Sulfones

Newborns and G-6-PD–deficient infants are particularly susceptible to dapsone hemolysis. Older infants might tolerate the amounts of sulfones excreted into milk.

(continued)
ANTIMICROBIAL DRUGS

ANTIPARASITIC DRUGS

Anthelmintics

Mebendazole was undetectable in milk in one woman and is poorly absorbed orally; therefore, it is unlikely to cause adverse effects in a breastfed infant. In contrast to an earlier report, it does not inhibit lactation. Only small amounts of praziquantel and ivermectin reach the infant and these drugs seem safe during nursing.  

Antimalarials

Undertake breastfeeding cautiously during long-term daily therapy with chloroquine or hydroxychloroquine because the importance of the small amounts of drug and metabolites in milk is unclear and accumulation can occur. Weekly prophylactic doses are probably safe because the amount of drug in milk is less than the infant prophylactic dose. Small amounts of quinine in milk are unlikely to harm the infant, although allergic reactions can occur. Pyrimethamine appears to be safe and might be excreted into milk in quantities sufficient to treat or protect infants <6 months of age against malaria; however, breastfeeding is not a reliable method of drug administration. Mefloquine appears in milk in small amounts after a single dose but has not been studied after repeated weekly administration for malaria prophylaxis.

Lindane

Lindane was excreted into milk at up to 30 times the typical background concentration (from environmental pollution) after maternal topical application of a 0.3% emulsion daily for 3 days. Milk concentrations remained elevated over background concentrations for at least 7 days. Although they have not been studied, alternative drugs (eg, permethrin, and pyrethrins) are preferred for nursing mothers because of their low toxicity.

ANTIVIRAL DRUGS

Acyclovir

Acyclovir has not been well studied, but a breastfed infant would receive about 1% of the mother’s weight-adjusted oral dosage. The low dosage in milk and its poor oral bioavailability indicate that it may be well tolerated by the nursing infant, even with large IV doses. Topical acyclovir applied to small areas of the mother’s body away from the breast should pose no risk to the infant.

Amantadine

Amantadine is a dopamine agonist that decreases serum prolactin and theoretically can decrease lactation, so it is best avoided during nursing. Rimantadine might not have the same effect.

Antiretrovirals

Lamivudine in breastmilk adds negligibly to the neonatal treatment dose. Nevirapine and zidovudine levels are also low but might offer some protection against breastmilk transmission of HIV-1.

β-LACTAMS

Cephalosporins

Cephalosporins appear in trace amounts in milk and can lead to disruption of the GI flora or, rarely, allergic sensitization. Breastfeeding is safe with first- and second-generation agents. The risk might be greater with the third-generation cephalosporins and similar agents (eg, aztreonam) that are more active against GI flora. Observe infants for diarrhea, thrush, and rashes.

(continued)
ANTIMICROBIAL DRUGS

Penicillins
Penicillins appear in trace amounts in milk that can occasionally lead to allergic sensitization, allergic reactions in previously sensitized infants, or disruption of the GI flora, especially with the broader-spectrum agents. Unless the infant is allergic to penicillin, breastfeeding is safe. Observe infants for diarrhea, rashes, and thrush.

MACROLIDES
Erythromycin, clarithromycin, and azithromycin are excreted into the milk in amounts much smaller than a typical infant dosage and are usually safe.1,40,41

QUINOLONES
Ciprofloxacin, fleroxacin, nalidixic acid, ofloxcin, and pefloxacin have been detected in milk.42,43 Ciprofloxacin seems to have caused pseudomembranous colitis in an infant via breastmilk,6 and nalidixic acid caused hemolytic anemia in a breastfed neonate.44 Most fluoroquinolones are best avoided during nursing. Norfloxacin was undetectable in milk after a 200 mg dose and might be acceptable for maternal UTI treatment because of its low milk excretion and poor oral bioavailability.

SULFONAMIDES
Some sulfonamides can cause hemolysis in G-6-PD–deficient infants; theoretically, sulfonamides increase the risk of kernicterus in neonates. Sulfamethoxazole, with or without trimethoprim, and sulfisoxazole can be used by mothers of healthy, full-term infants >2 months old.

TETRACYCLINES
Tooth staining from a tetracycline in breastmilk has not been reported. Milk calcium apparently inhibits absorption of the small amounts of tetracycline in milk. Infant absorption and serum concentrations have not been reported with other tetracyclines, but infants would only receive a few milligrams per day of demeclocycline, doxycycline, or minocycline with usual maternal dosages. Minocycline has caused black milk.45 Although other drugs are preferred for most infections, tetracyclines can be used for a short time (7–14 days); avoid prolonged or repeat courses during nursing.

MISCELLANEOUS ANTIMICROBIALS
Chloramphenicol
Breastfeeding is contraindicated during maternal chloramphenicol treatment. Milk concentrations are not sufficient to induce “gray baby” syndrome but theoretically might be enough to cause the rare, idiosyncratic aplastic anemia. Adverse reactions in infants, including refusal of the breast, falling asleep during feeding, and vomiting after feeding, have occurred.

Clindamycin
Clindamycin is excreted variably in small amounts into milk. It is not certain what effects these amounts have on infants’ GI flora (eg, pseudomembranous colitis), but a single case of bloody stools in an infant with normal stool flora was reported during maternal clindamycin use. Clindamycin is best avoided, if possible, but a few days of therapy with close monitoring of the infant is acceptable. Vaginal clindamycin presents less infant risk than oral or IV use.

Furazolidone
Furazolidone is poorly absorbed orally and can be used to treat maternal giardiasis if the infant is >1 month old.

(continued)
ANTIMICROBIAL DRUGS

Methenamine
The hippurate and methenamine salts of methenamine pass into milk in small quantities and seem safe to use.

Metronidazole
Metronidazole and its hydroxy metabolite are found in the serum of nursing infants in concentrations that are 10 to 20% of maternal serum concentrations. Anecdotal cases of “spitting up” (possibly from a bad taste), diarrhea, and isolation of Candida species in breastfed infants have been reported, but most infants do not have immediate reactions. Because of the carcinogenicity in animals, possible mutagenicity, and the relatively high infant serum concentrations achieved, metronidazole probably should be avoided in nursing mothers.\(^1\)\(^4\) When essential to treat trichomoniasis, metronidazole may be given as a single 2 g dose, and an alternative feeding method used for the next 24 hr. After longer courses for anaerobic infections, nursing can resume 12–24 hr after the final dose.

Nitrofurantoin
Nitrofurantoin is excreted into milk in pharmacologically unimportant amounts but avoid it with infants <1 month old and those with G-6-PD deficiency because of the risk of hemolysis.

Trimethoprim
Trimethoprim is excreted into milk in amounts that are not harmful.

Vancomycin
Because it is excreted into milk in only small amounts\(^4\)\(^7\) and is not orally absorbed, vancomycin is safe during nursing.

ANTINEOPLASTICS AND IMMUNOSUPPRESSANTS

Few reports exist, but breastfeeding is generally considered to be contraindicated in women receiving antineoplastics because of the potential for immunosuppression and carcinogenicity.

ALKYLATING AGENTS

Busulfan
Busulfan in a dosage of 4 mg/day for 5 weeks was taken by one woman while breastfeeding, with no apparent adverse effects on her infant’s leukocytes or hemoglobin. This case is not conclusive and breastfeeding is not recommended.

Cisplatin
Platinum was not detected in the milk of one patient at any time after an IV infusion of 100 mg/m\(^2\) of cisplatin. In another patient, milk platinum was 0.9 mg/L 19.5 hr after her third daily dose of 20 mg/m\(^2\). Because the platinum might be in a reactive form, nursing is not recommended during cisplatin therapy.

Cyclophosphamide
Cyclophosphamide is detectable in milk and has caused bone marrow depression in infants of women who nursed while receiving the drug. Breastfeeding is contraindicated during cyclophosphamide therapy.

ANTIMETABOLITES

Methotrexate
Low amounts of methotrexate were found in milk in one patient; however, this case is not conclusive. Low weekly doses for arthritis probably pose only a slight risk to the infant.

(continued)
ANTINEOPLASTICS AND IMMUNOSUPPRESSANTS

CYTOKINES

Interferons
An IV dose of 30 million units of interferon alfa-N3 resulted in only a slight increase over physiologic milk levels in one woman.48

DNA INTERCALATING DRUGS

Doxorubicin
Doxorubicin and its primary active metabolite, doxorubicinol, appear in milk, with their highest milk concentrations occurring 24 hr after a dose.

Mitoxantrone
Measurable levels of mitoxantrone occurred in milk for at least 28 days after 6 mg/kg was given daily for 3 days.49

MITOTIC INHIBITORS

Etoposide
Etoposide is undetectable in milk 24 hr after a dose.49

MISCELLANEOUS ANTINEOPLASTICS

Hydroxyurea
Only small amounts of hydroxyurea are found in milk, but breastfeeding is not advised.

IMMUNOSUPPRESSANTS

Three infants reportedly were breastfed safely during maternal azathioprine use (75–100 mg/day) after renal transplantation. Low concentrations of the azathioprine metabolite, mercaptopurine, were found in milk. Breastfeeding can be undertaken with close infant monitoring for infection or other signs of immunosuppression during azathioprine therapy, although there is concern over potential carcinogenicity.1,12 Maternal cyclosporine therapy results in the infant receiving ≤2% of the mother’s mg/kg dosage.50,51 At least 9 infants have been breastfed safely for 4–12 months during maternal therapy with cyclosporine, prednisone, and azathioprine. Infant serum cyclosporine levels were undetectable (<30 µg/L), renal function was unaffected, and the infants grew normally.50,51 Tacrolimus colostrum concentrations are about 50% of maternal serum concentrations.52 The implications of these low concentrations for the infant are not known.

CARDIOVASCULAR DRUGS

ANTIARRHYTHMIC DRUGS

Some antiarrhythmics reach near-therapeutic serum concentrations in breastfed infants. Amiodarone is excreted in amounts that might pose a hazard to the infant and it should not be used during nursing.1,53 Data on disopyramide indicate that infants can receive relatively large amounts of the drug and its active metabolite, with serum concentrations near the therapeutic range. Disopyramide can be used cauiously while breastfeeding older infants when other alternatives are unacceptable. Observe the infant for anticholinergic symptoms, and monitor infant serum concentrations if there is a concern. The anticholinergic activity of disopyramide might suppress lactation. (See Anticholinergics.) Sparse data from one patient indicate that tocainide should be used with caution during nursing. Because of its low oral bioavailability, maternal bretylium is unlikely to harm nursing infants; 400 mg q 8 hr was taken orally by one mother while nursing, with

(continued)
CARDIOVASCULAR DRUGS

no apparent effects on her infant. Infants receive trivial doses of digoxin via breastmilk. Amounts of flecainide in milk are small and unlikely to affect the infant. Lidocaine concentrations in milk during continuous IV infusion and epidural administration and in high doses as a local anesthetic are low and poorly absorbed by the infant, so it poses no hazard to the infant.54–56 Amounts of mexiletine in milk are too low to be detected in the serum of breastfed infants. Procainamide and its active metabolite, N-acetylprocainamide, are found in milk in fairly low concentrations; procainamide may be used with caution in nursing mothers. Propafenone milk concentrations are very low, but no clinical experience has been reported.57 Quinidine excretion seems inconsequential.

ANTIHYPERTENSIVE DRUGS

Certain antihypertensives are less desirable than others during nursing. Breastfed infants have serum clonidine concentrations approaching those of the mother.1,58 Clonidine and guanfacine also can decrease prolactin secretion. These drugs must be used with caution during breastfeeding and avoided if possible. Avoid reserpine because it can cause nasal stuffiness and increased tracheobronchial secretions in the infant. The angiotensin-converting enzyme (ACE) inhibitors, benazepril, captopril, and enalapril, are found in small amounts and no adverse effects have occurred in breastfed infants.1,59 In addition, milk ACE activity was in the normal range after a dose of enalapril. These ACE inhibitors are good choices during lactation; others have not been studied. Limited data indicate that low-dose, short-term use of hydralazine (ie, a few days postpartum) is safe. There is limited information on oral minoxidil in milk, but amounts are small. However, use minoxidil with caution, particularly when therapy involves large dosages and long-term use. Several studies indicate that methylldopa is excreted in unimportant amounts.

β-ADRENERGIC BLOCKING DRUGS

The excretion of β-blockers into breastmilk has been studied extensively. The infant’s dosage differs greatly among the different compounds, allowing a range of choices. The most water-soluble drugs reach the infant in the greatest amounts because of low serum protein binding. Water-soluble agents also have the longest half-lives, are renally eliminated, and therefore are more likely to accumulate in infants. Maternal therapy with atenolol and acebutolol have resulted in adverse effects (eg, bradycardia, hypotension, tachypnea, and cyanosis) in breastfed infants. These two drugs, as well as betaxolol, nadolol, sotalol, and timolol, should be avoided in mothers of newborn infants or when high dosage is required. Oxprenolol and mepindolol excretions are intermediate and should be avoided in neonates. Propranolol, metoprolol, and labetalol are excreted in low enough quantities to allow nursing even in the neonatal period.

CALCIUM-CHANNEL BLOCKING DRUGS

Case reports indicate that only small amounts of diltiazem, nifedipine, nimodipine, and nitrendipine are excreted into milk.1,60 Several case reports indicate that the amounts of verapamil and norverapamil in milk and infant serum are low. Verapamil appears to be safe during nursing.

CENTRAL NERVOUS SYSTEM DRUGS

ANTICONVULSANTS

Breastfed infants can achieve serum anticonvulsant concentrations that produce pharmacologic effects. Mild drowsiness, irritability, and feeding difficulties are common in the infants of mothers taking sedating anticonvulsants, especially during the early neonatal period.1,61 Breastfeeding can (continued)
mitigate withdrawal symptoms in infants whose mothers took sedating anticonvulsants during preg-
nancy, and withdrawal symptoms have been observed after abrupt weaning. Serum concentration
monitoring in breastfed infants might be indicated, particularly in infants who are excessively drowsy,
feed poorly, or gain weight inadequately. Long-term effects of exposure are not well studied. Infants
of mothers taking anticonvulsants might have more difficulty nursing and breastfed for a shorter du-
ration, possibly because of negative or equivocal safety advice given by health professionals.62–64
No data are available for some of the newer anticonvulsants such as felbamate, gabapentin,
levetiracetam, oxcarbazepine, tiagabine, and topiramate. Breastfeeding is not recommended
during felbamate use.65–67

Carbamazepine
Carbamazepine and its major active metabolite are excreted into milk and can be detected in
nursing infants’ serum; concentrations are usually low but near the therapeutic range in some in-
fants. Two cases of hepatic dysfunction in breastfed neonates have been reported. Poor feeding
also has been reported. Carbamazepine can be used during lactation, but close observation of the
infant for jaundice and other signs of possible adverse idiosyncratic effects is advisable.67 Measure-
ment of infant serum concentration might be indicated if symptoms occur.

Clonazepam
Serum concentrations of clonazepam were low in two nursing infants, and no effects were noted.1
In another infant, breastfeeding increased serum concentrations over those present at birth.68
Clonazepam has been detected in the serum of a breastfed neonate whose mother was receiving
the drug before and after delivery but was undetectable in 4 others.69 Observation of the infant for
drowsiness and monitoring of the infant’s serum concentration might be indicated.

Ethosuximide
Breastfed infants can attain ethosuximide serum concentrations near the therapeutic range, and
some infants might become drowsy or fussy. Breastfeed with caution and keep the mother’s
serum concentrations as low as possible while remaining in the therapeutic range. Infant serum
drug concentration monitoring is indicated.

Lamotrigine
Lamotrigine concentrations in infants breastfed during maternal lamotrigine therapy have ranged
from 22 to 85% of the maternal serum concentration, but no adverse effects have been reported
with these relatively high levels.67,70,71 Infants can be allowed to nurse, but close monitoring for
side effects such as rash (which can be life-threatening), drowsiness, or poor sucking is essential.
Obtain an infant serum concentration if adverse effects are suspected and discontinue breastfeed-
ing if rash occurs.

Phenobarbital
The effect of phenobarbital is unpredictable: drowsiness leading to feeding difficulties can occur;
breastfeeding can prevent withdrawal symptoms in infants whose mothers took phenobarbital dur-
ing pregnancy; and withdrawal symptoms have been observed after abrupt weaning. Phenobarbi-
tal can be used in low to moderate dosages but monitor infant behavior, weight gain, and, if there
is concern, serum concentrations. Sometimes breastfeeding must be discontinued because of ex-
cessive drowsiness and poor weight gain.

Phenytoin
Only small amounts of phenytoin are excreted into milk. Rarely, infants might experience idiosyn-
cratic reactions such as cyanosis and methemoglobinemia, but infants generally tolerate phenytoin
in milk well.
CENTRAL NERVOUS SYSTEM DRUGS

**Primidone**
Primidone and its metabolites (phenylethylmalonamide, phenobarbital, and parahydroxyphenobarbital) appear in milk in large amounts. Considerations are the same as those for phenobarbital. (See Phenobarbital.)

**Valproic Acid**
Milk concentrations of valproate are low, and usually no effects occur in infants. One case of probable infant thrombocytopenic purpura from valproate in milk has been reported.72 Observe infants for rare idiosyncratic effects such as thrombocytopenia and hepatotoxicity.67

**Vigabatrin**
Limited data from two mothers indicate that a breastfed infant would receive <4% of the mother’s mg/kg dose of vigabatrin.73

**Zonisamide**
Peak milk concentrations were 9–10 mg/L with a maternal dose of 300 mg/day in one mother. No data are available on effects in breastfed infants.74

ANTIDEPRESSANTS

**Heterocyclic Antidepressants**
Most of these drugs have not been well studied during lactation. Some investigators recommend against the use of antidepressants because of theoretical (but undemonstrated) long-term effects on infants’ neurologic development; others consider tricyclic antidepressants to be acceptable. Follow-up for 1–3 yr in a small group of breastfed infants indicates no adverse effects on growth and development.75 Another study found that breastfed infants of mothers taking dothiepin had cognitive development equal to controls at 3 yr of age.76 Sedating TCAs and those with active metabolites (eg, amitriptyline, doxepin, and imipramine) might be less desirable than other TCAs. Respiratory depression was reported in one breastfed infant whose mother was taking doxepin 25 mg tid, but an infant whose mother was taking 150 mg at night had no problems. Another report found poor sucking and swallowing, muscle hypotonia, and vomiting in a 9-day-old whose mother was taking doxepin 35 mg/day.77 Maternal dosages of amitriptyline up to 150 mg/day, clomipramine 150 mg/day, desipramine 300 mg/day, imipramine 200 mg/day, or nortriptyline 125 mg/day have not caused observable effects in the infants studied. In several infants, nortriptyline serum concentrations were undetectable with maternal nortriptyline dosages of up to 125 mg/day or amitriptyline 175 mg/day.69,78,79 One nortriptyline metabolite has been detected in low levels in the serum of breastfed infants without adverse consequences.78,80 Nortriptyline (and probably the other secondary amine, desipramine) is the TCA of choice during breastfeeding. Doxepin should be avoided. Giving the drug as a single dose at bedtime and skipping nighttime feeding(s) can further minimize infant exposure. A dose of 250 mg/day of amoxapine or 100–150 mg/day of maprotiline produces low drug concentrations in milk, but effects of these drugs on infants have not been well studied.

**Selective Serotonin Reuptake Inhibitors**
Although the average daily dosages of fluoxetine and norfluoxetine in milk are about 7% of the mother’s weight-adjusted dosages, some mothers excrete as much as 12% of a dosage and the drugs’ half-lives are very long.51 One case of colic (increased crying, decreased sleep, watery stools, and vomiting) and unexplained high serum concentrations were reported in a breastfed 6-week-old infant. The infant improved after switching to formula and colic reappeared with rechallenge. Other case reports include seizure-like activity, irritability, hyperglycemia and
CENTRAL NERVOUS SYSTEM DRUGS

Fluoxetine in breastmilk had no effect on neurologic development in 4 infants, but a larger retrospective study found that fluoxetine can reduce the growth rate of infants who are exposed via breastmilk from birth. Fluoxetine should be avoided during breastfeeding if possible, although older infants might be less susceptible to fluoxetine’s effects than newborns. Monitor infants carefully for behavioral symptoms and adequate weight gain. Citalopram reaches the infant in dosages of about 5% of the mother’s mg/kg dosage. The manufacturer states that drowsiness and weight loss in breastfed infants have occurred, and uneasy sleep occurred in the infant of a mother taking citalopram. Citalopram is not a good choice while breastfeeding a newborn. Infants receive a dose <1% of the maternal fluvoxamine dose. Several infants grew and developed normally with maternal fluvoxamine use. With paroxetine, infants receive about 1.5% of the maternal dosage. Of 23 infants studied, only 1 had detectable serum concentrations of paroxetine. No adverse behavioral or growth effects have been observed in studies, but one case of infant agitation and feeding difficulties has been reported. Sertraline dosage to the breastfed infant is <2% of the maternal dosage; concentrations in infant serum are usually low to undetectable, platelet serotonin is unaffected, and no adverse effects on growth have been seen in controlled follow-up. One case of infant agitation and one of somnolence and developmental difficulties have been reported spontaneously to Australian authorities. Sertraline and paroxetine are considered the SSRIs of choice during breastfeeding, especially with a neonate.

Monoamine Oxidase Inhibitors

There are no data on the amounts of older nonselective MAOIs excreted into milk. Because of their potential for toxicity and lactation inhibition, avoid MAOIs during nursing. With moclobemide, a reversible MAO-A inhibitor not available in the United States, infants receive a dose <1% of the mother’s dose and no side effects have been reported in a small number of infants studied.

Other Antidepressants

Bupropion and its metabolites were undetectable in one 14-month-old infant whose mother was taking 300 mg/day and nursing twice daily. Nefazodone and trazodone dosages in the infant are <1% of the mother’s mg/kg dosage, but only a few cases have been reported. One case of drowsiness, lethargy, poor feeding, and inability to maintain normal body temperature was reported in a small preterm breastfed infant whose mother was taking nefazodone 300 mg/day. Infants receive venlafaxine doses of up to 9.2% of the mother’s mg/kg dosage and the active metabolite is detectable in the infant’s serum. Although adverse effects were not seen in 3 breastfed infants, caution should be used with venlafaxine until more experience is gained.

ANTIPSYCHOTIC DRUGS

Data on the use of antipsychotics during lactation are sparse. Phenothiazines and thio-
cause no problems for nursing infants unless dosages are at the high end of the range or combinations of drugs are used.\textsuperscript{104,105} Breastfeeding during clozapine use in 4 infants resulted in sedation in 1 and agranulocytosis in another, which resolved with discontinuation; nursing is not recommended during clozapine use.\textsuperscript{100,107} Exposure of 2 infants to olanzapine in breastmilk for a few days each caused no untoward events, but more experience is needed.\textsuperscript{108} One mother taking risperidone excreted about 4\% of her mg/kg dosage into breastmilk; no infant side effects were noted.\textsuperscript{109}

ANXIOLYTICS, SEDATIVES, AND HYPNOTICS

Many sedatives and hypnotics pass into breastmilk in measurable and potentially important amounts. Minimize sedative and hypnotic intake during lactation.

Anesthetics, General

Compared with epidural anesthesia, general anesthesia used during cesarean delivery can decrease the frequency and duration of breastfeeding.\textsuperscript{19} Excretion of most inhalation anesthetics in breastmilk has not been well studied. Blood levels of anesthetic gases such as desflurane, enfurane, halothane, isoflurane, nitrous oxide, and sevoflurane drop rapidly after termination of anesthesia, are predicted to pass poorly into milk, and are probably poorly absorbed by the infant.\textsuperscript{19,20} Etomidate milk levels drop rapidly after a dose and should pose little risk to the infant.\textsuperscript{110} Amounts of propofol in milk are small and do not have good oral bioavailability in the infant. Typical IV doses of methohexital or thiopental for induction of anesthesia produce low concentrations in milk that do not cause effects in the infant.\textsuperscript{1,18,110} Current opinion suggests that breastfeeding can be resumed as soon as the mother has recovered sufficiently from general anesthesia to nurse.\textsuperscript{19,20}

Barbiturates

These drugs can stimulate metabolism of endogenous compounds in the infant when small amounts pass into milk. Short-acting agents are preferable to long-acting agents because smaller amounts are excreted into milk. Large single doses might have more potential for causing infant drowsiness than multiple small doses. (See also Anesthetics, General; Anticonvulsants.)

Benzodiazepines

Long-acting benzodiazepines and those with active metabolites (eg, diazepam) can accumulate and cause adverse effects in infants, especially with repeated doses, and in neonates because of their immature excretory mechanisms. Bromazepam taken by the mother might have contributed to the death of her 4-week-old breastfed infant with a 5-day history of apneic episodes.\textsuperscript{111} A single dose of diazepam for short dental, surgical, or diagnostic procedures is not likely to cause sedation in infants past the neonatal period.\textsuperscript{19} Milk alprazolam concentrations are low,\textsuperscript{112} but infant drowsiness and withdrawal symptoms have been reported with alprazolam use during nursing.\textsuperscript{1,5} When oral therapy is essential, the short-acting agents, oxazepam or lorazepam, are preferred; temazepam also might be acceptable.\textsuperscript{103,113} Midazolam concentrations in milk are low and unlikely to affect the infant after a single dose or short course of therapy.\textsuperscript{19,114}

Chloral Hydrate

Chloral hydrate and its active metabolite, trichloroethanol, appear in milk in dosages that approximate an infant sedative dosage and are detectable for up to 24 hr after a single dose. Using another hypnotic is advisable during nursing.
GASTROINTESTINAL DRUGS

**Zaleplon**

The dose in milk is very small and the drug disappears from breastmilk rapidly.\(^{115}\)

**Zolpidem**

Zolpidem milk concentrations are low for 3 hr after a dose and undetectable thereafter.\(^{116}\)

LITHIUM

Lithium in milk can adversely affect the infant when its elimination is impaired, as in dehydration or in neonates or premature infants. Neonates also can have transplacentally acquired serum lithium levels. The long-term effects of lithium on infants are not known; many investigators consider lithium therapy a contraindication to breastfeeding, but others do not. Lithium may be used cautiously in mothers who are carefully selected for their ability to monitor their full-term infants. Discontinue breastfeeding immediately if the infant appears restless or looks ill. Measurement of serum lithium concentrations in the infant can help rule out lithium toxicity.\(^1,67,80\)

PARKINSONISM DRUGS

**Dopamine Agonists**

Some ergot alkaloids have dopaminergic activity that can suppress prolactin release and lactation. Bromocriptine was used therapeutically for this purpose but has lost this indication in the United States because of potentially serious maternal toxicity (ie, stroke, death).

**Levodopa**

Levodopa decreases serum prolactin in non-nursing women with hyperprolactinemia and galactorrhea in a dose-dependent fashion and inhibits lactation in animals at high dosages.\(^1\) One mother taking sustained-release levodopa/carbidopa 200 mg/50 mg qid successfully breastfed her infant whose development was normal at age 2 yr.\(^{117}\)

GASTROINTESTINAL DRUGS

**ACID-PEPTIC THERAPY**

**Antacids**

Although aluminum, calcium, and magnesium antacids are partially absorbed, they are unlikely to appreciably increase concentrations of these ions in milk and are safe to use.

**Histamine H\(_2\)-Blockers**

Cimetidine is concentrated in milk because of ion trapping and possibly active secretion;\(^{118}\) ranitidine doses in milk are lower. Famotidine and nizatidine have the lowest concentrations in milk and are preferred during nursing.

**Proton Pump Inhibitors**

Omeprazole and lansoprazole have not been adequately studied. In one mother, omeprazole milk levels were low and her newborn infant was breastfed without harm.\(^{119}\)

**Sucralfate**

Because sucralfate is virtually nonabsorbable, it might be preferable to H\(_2\)-receptor antagonists.

GASTROINTESTINAL MOTILITY

**Antidiarrheals**

Nonabsorbable products such as kaolin-pectin are preferred in nursing mothers. The loperamide prodrug loperamide oxide results in only small amounts of loperamide in breastmilk.

(continued)
GASTROINTESTINAL DRUGS

Diphenoxylate excretion into milk has not been studied. One or two small doses of loperamide or diphenoxylate daily should pose little risk to the nursing infant. Avoid bismuth subsalicylate because salicylate is absorbable.

Cathartics and Laxatives

Some anthraquinone derivatives, such as aloe and cascara, and other stimulant cathartics (eg, phenolphthalein) should be avoided during nursing because of a laxative effect in breastfed infants. Laxatives that are nonabsorbable or poorly absorbed, such as bulk-forming (eg, psyllium), osmotic (eg, magnesium or phosphate salts), or stool-soothing (eg, docusate) types, are preferred during lactation. Senna in moderate dosages is acceptable if other measures fail. Bisacodyl is virtually unabsorbed from the GI tract and should be safe.

Gastrokinetic Agents

Metoclopramide elevates serum prolactin via central dopaminergic antagonism and results in increased milk production and a more rapid transition from colostrum to mature milk. It can be used therapeutically in mothers who are producing insufficient quantities of milk, such as the mothers of premature or sick infants or adoptive mothers. Although infant dosages of metoclopramide from milk are low, the infant’s serum prolactin concentrations can become elevated. Metoclopramide can induce depression, so caution is warranted. Limiting the duration of metoclopramide therapy to 14 days is essential, and it should not be used in mothers with a history of depression. Domperidone (not available in the U.S.) also has been used to increase milk supply and results in lower milk drug levels than metoclopramide.120,121

MISCELLANEOUS GASTROINTESTINAL DRUGS

Mesalamine Derivatives

Small amounts of sulfasalazine and sulfapyridine have been found in milk and infants’ sera after oral sulfasalazine use. The small amount of sulfapyridine released should cause no bilirubin displacement. Olsalazine is not detectable in milk, but its metabolite, N-acetyl-5-ASA, is found in small amounts.122 Small amounts of mesalamine and larger amounts of its metabolite are found in milk after oral administration.123 Diarrhea has been reported in infants of mothers using mesalamine derivatives, but a controlled study found the frequency of diarrhea to be no greater than that in infants of untreated mothers.124 Sulfasalazine and mesalamine and its derivatives may be used during nursing.

Ursodiol

Ursodiol was undetectable in the milk of 1 lactating mother, and her nursing infant developed normally during therapy.125 Maternal ursodiol therapy decreased the bile acid concentration in colostrum and was found in trivial amounts in breastmilk in 16 mothers with intrahepatic cholestasis of pregnancy.126 Their infants showed no adverse effects.

HEMATOLOGIC DRUGS

COAGULANTS AND ANTICOAGULANTS

Coumarins

Amounts of warfarin in milk are of no clinical consequence with a maternal dosage of ≤12 mg/day because of extensive protein binding. Higher dosages have not been studied. Other coumarin derivatives (eg,acenocoumarol, dicumarol, and phenprocoumon) also appear to be safe.127
HEMATOLOGIC DRUGS

Heparins
Although minimal documentation exists, it is unlikely that heparin or low-molecular-weight heparins (e.g., enoxaparin, dalteparin) pass into milk or are absorbed orally by the infant; anticoagulant activity was undetectable in 1 breastfed infant whose mother received 20–40 mg/day of enoxaparin. Hirudin was not detectable in milk.

Indandiones
Anisindione and phenindione are contraindicated because infant hemorrhage has occurred.

HORMONES AND SYNTHETIC SUBSTITUTES

ADRENAL HORMONES

Corticosteroids
Prednisone and prednisolone excretions into milk are minimal even with large oral doses. The infant dosage can be reduced even further by using prednisolone rather than prednisone and avoiding nursing for 3–4 hr after a dose. Three infants have been breastfed during long-term maternal use of methylprednisolone 6–8 mg/day with apparent safety. Large IV doses of corticosteroids or use of long-acting agents such as dexamethasone have not been studied, and caution is warranted. Depot injections, inhaled corticosteroids (e.g., beclomethasone, fluticasone), or topical corticosteroids should present little or no risk to the infant because of low maternal serum concentrations. However, topical application to the nipple has caused adverse effects in the infant because of direct ingestion.

ANTIDIABETIC DRUGS

Insulin
Diabetic mothers using insulin may nurse their infants. However, it has been found empirically that the mother might need to reduce her insulin dosage to 55–75% of the prepregnancy dosage. Close monitoring is required postpartum because the return to prepregnancy insulin dosage has been variably reported to take 1–6 weeks.

Sulfonylureas
Tolbutamide is excreted in milk in small amounts that should cause no harm. The manufacturer reports that chlorpropamide concentrations in milk are low, but no published clinical data are available on this or other sulfonylureas.

CONTRACEPTIVES

Estrogen–Progestin Combinations
Although present in milk in small amounts, estrogens and progestins are readily metabolized by nursing infants. Rare case reports of breast enlargement in infants have been attributed to estrogen-containing oral contraceptives. These effects occur primarily with products containing >50 μg of estrogen. These high-estrogen contraceptives also markedly suppress lactation, especially when administered immediately postpartum. When currently available low-dose estrogen–progestin combination contraceptives are begun ≥6 weeks postpartum, a dramatic immediate effect on milk supply is usually not seen, but long-term negative effects on milk yield lead to early feeding supplementation and discontinuation of breastfeeding and decreased infant growth. An 8-year follow-up of breastfed infants of mothers taking contraceptives containing ethinyl estradiol 50 μg found no adverse effects on the infants’ development or behavior. Progestin-only contraceptives are preferred during lactation.

(continued)
HORMONES AND SYNTHETIC SUBSTITUTES

Progestin Only

No immediate effects have been reported with progestin-only contraceptives such as levonorgestrel implants, depot medroxyprogesterone acetate, or oral norethindrone or norgestrel. Progestin-only contraceptives generally have no effect on, or enhance, milk supply and might extend the duration of lactation. Although infant growth might undergo a slight, transient depression after insertion of levonorgestrel implants, large multicenter studies have found no effect of progestin-only contraceptives on growth and development of infants and children up to puberty. Early (ie, immediately postpartum) initiation of these agents is controversial. Because physiologic postpartum progesterone withdrawal is a primary stimulus for lactation, it appears best to wait for at least 3 days postpartum before starting a progestin-only contraceptive. One small study found no adverse effects on lactation or infant growth when depot medroxyprogesterone was given immediately postpartum, but anecdotal reports of lactation suppression with immediate postpartum administration exist. Progestin-only contraceptives started 6 weeks postpartum are the preferred hormonal contraceptives during lactation. (See also Progesterone.)

FEMALE SEX HORMONES

Progesterone

Contraceptive use via implants (investigationally) or intrauterine devices transfers little progesterone to the breastfed infant, and any drug in milk is minimally absorbed by the infant.1,140,141 Milk progesterone concentrations have not been measured after higher doses used to treat premenstrual syndrome.

THYROID AND ANTITHYROID DRUGS

Iodides

Inorganic iodide is contraindicated during breastfeeding because of possible thyroid suppression and rash. Topical and vaginal povidone–iodine in nursing mothers results in elevated milk iodine concentrations and occasional thyroid suppression in nursing infants. Avoid povidone–iodine preparations while nursing and minimize their use during delivery.

Thioamides

Propylthiouracil is the antithyroid drug of choice during lactation; little passes into milk and infant thyroid suppression does not occur. Dosages as high as 750 mg/day have been given to nursing mothers with no adverse effects in their infants. Methimazole 20 mg/day or carbimazole (a methimazole prodrug) 15 mg/day also can be used, but these drugs pass into milk in greater quantities and have longer half-lives than propylthiouracil. Infants of mothers who took 20 mg/day of methimazole while nursing had no decrease in intellectual or physical development at age 1 yr. A potential for idiosyncratic reactions (eg, agranulocytosis) and hypothyroidism exists, and measurement of the infant’s serum thyroxine and TSH concentrations at 2–4-week intervals might be prudent during maternal antithyroid drug use.

Thyroid Hormones

Normal lactation requires thyroid hormones. Levothyroxine (T₄) passes into milk poorly, although liothyronine (T₃) might pass in more physiologically relevant amounts. Milk concentrations of thyroid hormones have not been measured after exogenous administration, but a physiologic replacement dosage of levothyroxine to a breastfeeding mother is not expected to result in excessive thyroid administration to the infant. Replacement therapy with liothyronine or supraphysiologic maternal levothyroxine dosage might transfer larger amounts of liothyronine to the infant.

(continued)
HORMONES AND SYNTHETIC SUBSTITUTES

Protirelin

Protirelin (thyrotropin-releasing hormone [TRH]) causes an increase in prolactin secretion and can enhance milk yield.\(^{145}\)

MISCELLANEOUS HORMONAL AGENTS

Ergot Alkaloids

Ergonovine can lower postpartum serum prolactin concentrations, but methylergonovine apparently does not. Methylergonovine is not found in milk in important quantities. Short-term, low-dose regimens of these agents immediately postpartum pose no hazard to the infant, but methylergonovine is preferred because it does not inhibit lactation. Courses of these drugs given several days postpartum can expose the infant to greater risk of ergot side effects because of the larger amount of milk consumed at this age.

Calcitriol

Calcitriol requirements in hypoparathyroid women decrease during lactation. Failure to substantially decrease (by up to two-thirds) the calcitriol dosage results in maternal hypercalcemia.\(^ {146,147}\)

Desmopressin

Desmopressin is excreted in negligible amounts into milk and is poorly absorbed orally by the infant, so it appears safe to use.

Human Growth Hormone

Somatropin can increase milk production in mothers with an insufficient milk supply.\(^{148,149}\)

RENAL AND ELECTROLYTES

DIURETICS

Large dosages of short-acting thiazide-type diuretics (eg, hydrochlorothiazide), usual dosages of loop diuretics (eg, furosemide), or long-acting thiazide-type diuretics (eg, chlorthalidone and bendroflumethiazide) can suppress lactation and should be avoided. Long-acting agents also can accumulate in infants’ serum. Low dosages of short-acting thiazide-type diuretics should pose no problems to the infant or suppress lactation. Acetazolamide appears in milk in small amounts that are unlikely to harm the infant. The amounts of spironolactone and its metabolites in milk are inconsequential.

ELECTROLYTES

Bisphosphonates

Pamidronate was used successfully in one patient to treat bone loss associated with reflex sympathetic dystrophy. The drug was undetectable in breastmilk.\(^ {150}\)

Fluoride

Fluoride supplementation is not recommended during the first 6 months after birth; from 6 months to 3 yr of age, fluoride supplementation of the breastfed infant is recommended only if the mother’s water supply contains <0.3 ppm fluoride.\(^ {154}\)

Magnesium Sulfate

When given IV, magnesium sulfate increases milk magnesium concentrations only slightly. Oral absorption of magnesium is poor, so maternal magnesium therapy is not a contraindication to breastfeeding.

(continued)
RENAL AND ELECTROLYTES

ANTIGOUT AGENTS

**Allopurinol**

This drug and its active metabolite, oxypurinol, are excreted into milk in nearly therapeutic amounts, and oxypurinol is detectable in the nursing infant’s serum in near-therapeutic levels. Although one infant breastfed without harm during maternal allopurinol therapy, observe infants for side effects, especially hypersensitivity reactions. If possible, give allopurinol to the mother in a single dose after the last nursing of the day.

**Colchicine**

Several infants have been breast-fed safely during long-term, low-dose administration of colchicine to the mother for familial Mediterranean fever. The amount excreted in milk indicates that toxicity might occur with higher dosages. Colchicine decreases milk production and alters milk composition in animals when infused into the udder. Use it with great caution and in low dosages when breastfeeding, especially with a neonate.

RESPIRATORY DRUGS

ANTIASTHMATICS

**Anticholinergics**

Excretion of anticholinergics into milk has not been studied. Theoretical hazards of the orally absorbable compounds include anticholinergic effects such as drying of secretions, temperature elevations, and CNS disturbances in the infant. Anticholinergics might inhibit lactation by inhibiting growth hormone and oxytocin secretion. Observe infants carefully for anticholinergic symptoms and signs of decreased lactation (eg, insatiety, poor weight gain) when anticholinergics are given to the mother. It is unlikely that inhaled ipratropium affects the infant or milk production.

**Terbutaline**

Oral administration results in low milk terbutaline concentrations, causes no symptoms in breastfed infants, and is not expected to decrease milk supply. Other β2-receptor agonists (eg, albuterol) appear safe to use orally, but inhaler products should transfer less drug to the infant and are preferred.

**Theophylline**

Maternal theophylline use occasionally can cause irritability and fretful sleep in infants. Newborn infants are most likely to be affected because of their slow elimination and low serum protein binding of theophylline. There is no need to avoid theophylline products; however, keep maternal serum concentrations in the lower part of the therapeutic range and measure infant serum concentrations if side effects occur. The related drug dyphylline is excreted into milk in greater amounts and is best avoided.

**ANTIHISTAMINES**

There are few studies on antihistamine use during lactation. One study found drowsiness or irritability in 12% of breastfed infants whose mothers took antihistamines. Older sedating (and more anticholinergic) antihistamines are more problematic because they can affect the infant and might suppress lactation. Nonsedating antihistamines are preferred agents for long-term therapy. However, single bedtime doses of a sedating antihistamine after the last feeding of the day might be adequate and minimize the amount the infant receives. Avoid sedating antihistamines in high dosages, in SR formulations, or in combinations with sympathomimetic agents.

(continued)
RESPIRATORY DRUGS

Cetirizine
Cetirizine has not been studied and is not a preferred agent because of its sedative and anticholinergic effects.

Cyproheptadine
Cyproheptadine lowers maternal serum prolactin and should be avoided during lactation.

Fexofenadine
Based on terfenadine experience, fexofenadine is likely to be well tolerated by breastfed infants.5,156

Loratadine
Loratadine is excreted into milk in seemingly unimportant amounts.

Triprolidine
Only small amounts of triprolidine are found in breastmilk.

COUGH AND COLD

α-Adrenergic sympathomimetics decrease milk flow in animals by central inhibition of secretion and release of oxytocin and by peripheral vasoconstriction, which limits the access of oxytocin to myoepithelial cells in the mammary glands. Norepinephrine also might decrease prolactin release. Although these effects are not well documented in humans, lactation inhibition seems to occur with oral decongestant (e.g., pseudoephedrine) use; therefore, sympathomimetic nasal sprays (e.g., oxymetazoline) are recommended over oral decongestant products. Pseudoephedrine also can cause irritability in some infants.5

MISCELLANEOUS DRUGS

CHOLINERGIC DRUGS

Six infants of mothers treated with neostigmine for myasthenia gravis were reportedly breastfed successfully. Neostigmine was not found in milk, but 1 infant appeared to have abdominal cramps after each breastfeeding. Pyridostigmine has been used safely during breastfeeding in 3 patients with myasthenia gravis.

Baclofen
Only small amounts of baclofen appear in milk, and it may be used in nursing mothers with caution.

Bupivacaine
Bupivacaine appears in milk in small amounts when administered to the mother by intrapleural or epidural routes but has no effect on the infant.55 Epidural analgesia with bupivacaine postcesarean section improved breastfeeding performance in one study.157 (See also Lidocaine in Antiarrhythmics.)

Dantrolene
Several dantrolene doses totaling 720 mg IV over 2 days to a postpartum mother yielded peak milk levels of 12 mg/L. Dantrolene half-life in milk was 9.2 hr.158

Pyridoxine
In high doses (200–600 mg/day), pyridoxine has been used therapeutically to suppress lactation, although it is often not effective. With usual dosages found in foods and low-dose vitamin supplements, pyridoxine has no effect on prolactin or lactation.

(continued)
MISCELLANEOUS DRUGS

Radiopharmaceuticals
Exposure of the infant to excessive amounts of radioactivity is usually the primary concern raised by administration of radiopharmaceuticals to nursing mothers, rather than any pharmacologic toxicity of the agent. Some, but not all, radiopharmaceuticals require discontinuation of breastfeeding, at least temporarily, after administration to a nursing mother. Radioactive iodine compounds are the most dangerous and might require complete cessation of breastfeeding. The period needed for milk radioactivity to decline (by means of radioactive decay and maternal excretion) to a safe exposure level depends on several factors: dosage, biological half-life, radionuclide half-life, and “contamination” with other isotopes. The age of the infant, potential for oral absorption of the radionuclide from the infant’s GI tract, and threshold level that is considered safe are also important factors. Measurement of milk radioactivity can aid in determining when breastfeeding can resume. Consult specialty sources for more detailed information.159,160

Retinoids
Acitretin passes into breastmilk in a quantity sufficient to merit avoidance of nursing while taking it. Although there is no information on use during lactation, the manufacturers of oral isotretinoin and topical tretinoin state that they are not compatible with nursing. Based on the systemic bioavailability of tretinoin applied topically to a small area such as the face, it is unlikely that harmful amounts reach the infant via breastmilk. Avoid contact of the infant’s skin with treated areas of the mother’s skin.

Vaccines
Breastfeeding is not a contraindication to the use of any vaccine (live or inactivated) in the nursing mother.161

DIAGNOSTIC AGENTS

Iodinated Contrast Media
Iopanoic acid contains free iodide that can be detected in milk. (See Iodides.) Diatrizoate, iodamide, iohexol, metrizoate, and metrizamide are detectable in milk after IV administration. Although no adverse effects have been reported in infants, breastfeeding probably should be withheld for a period after administration of most iodinated contrast media, the period depending on its rate of elimination. A few hours is probably adequate after an IV pyelogram. Large amounts of iodine are excreted into milk for weeks after lymphangiography with ethiodized oil, and nursing should be discontinued after this procedure.

Fluorescein
Fluorescein is detectable in milk after IV or topical administration. After IV administration, it had a milk half-life of 62 hr in one mother. The drug might present a risk to neonates who are undergoing phototherapy. Temporarily withholding nursing after fluorescein use (especially IV) seems appropriate in this situation.

Gadolinium
Gadodiamide and gadopentetate, used in magnetic resonance imaging, are detectable in milk but have poor oral absorption and are rapidly excreted renally. Suspension of breastfeeding is not necessary after use of these agents.162

(continued)
**Alcohol**

Alcohol equilibrates rapidly between blood and milk, resulting in milk concentrations equivalent to simultaneous blood concentrations. Peak maternal serum alcohol levels occur later (1 hr after the drink) in nursing mothers than in non-nursing women.\(^{163}\) Alteration in milk odor parallels milk alcohol levels.\(^{164,165}\) Potential effects on infants depend on the pattern of use. Drunkenness (deep, un arousable sleep with snoring, deep respiration, no reaction to pain, inability to suck, excessive perspiration, and a feeble pulse) was reported after maternal binge drinking. Pseudo-Cushing syndrome was reported in the infant of a chronic alcoholic mother. One prospective study suggests that as little as 1 drink daily can cause slight impairment of the infant’s motor development; the impairment increases in a dose-dependent fashion.\(^{166}\) Infants suck more but consume less milk after maternal alcohol ingestion.\(^{164}\) Alcohol also affects lactation; it inhibits the milk ejection reflex in a dose-dependent fashion, with single doses >2 g/kg completely blocking suckling-induced oxytocin release. Animal studies show that alcohol consumption results in a reduced suckling-induced prolactin release and reduced milk yield. An unknown substance in beer increases maternal serum prolactin; this effect also occurs with nonalcoholic beer.\(^{1,167}\) Use alcohol in moderation during lactation and withhold nursing temporarily after alcohol consumption, with the duration dependent on the amount consumed—at least 2 hr per drink is suggested.\(^{168}\)

**Amphetamine**

In a mother taking amphetamine 20 mg/day therapeutically, amphetamine concentrations in milk were less than those in serum and no adverse effects on the infant were noted. However, there is likely to be substantial intersubject variation in excretion, and concentrations in milk have not been measured during high-dose abuse of amphetamines. Anecdotally, infants breastfed by amphetamine abusers seem to experience drug-induced behavioral abnormalities such as agitation and crying. Amphetamine also inhibits prolactin release and, in high dosages, can interfere with lactation.

**Caffeine**

Anecdotal reports of infant jitteriness and difficulty sleeping have been reported with very high maternal intake of caffeine, but infant serum caffeine concentrations were not measured. Systematic studies have indicated that caffeine and its metabolites are excreted into milk in relatively small amounts with usual maternal intake and infants are usually not affected, even with high maternal intake.\(^{1,169,170}\) Effects are more likely in premature and newborn infants because of their greatly diminished ability to metabolize caffeine.

**Cocaine**

Although not well studied in humans, the chemical nature of cocaine and results from animal studies indicate that it probably appears in milk in amounts that affect the infant. Cocaine was detectable in milk for 24–36 hr after use. In addition, serum cholinesterase, which is needed to metabolize the drug, is low in newborns. Cocaine and its toxic metabolite can be detected in milk and can cause adverse effects (vomiting, diarrhea, irritability, and dilated pupils) in breastfed infants. Convulsions occurred in an infant whose mother used topical cocaine to treat sore nipples. Breastfeeding is not recommended when the mother is a chronic cocaine user, and even occasional use of cocaine is discouraged during breastfeeding. Withhold breastfeeding for at least 24 hr after occasional cocaine use.\(^{1,171}\)

**Heroin**

Abuse can result in high enough concentrations in milk to cause addiction or alleviate withdrawal symptoms in infants; however, breastfeeding is not a reliable method of preventing withdrawal.
DRUGS FOR NONMEDICAL USE

Most authorities consider breastfeeding safe during methadone maintenance in doses up to 80 mg/day.\(^1\)\(^2\)

**Marijuana**

Marijuana excretion into milk is not well studied, but dronabinol (tetrahydrocannabinol) can reach high concentrations in milk and be detected in the infant, particularly with heavy maternal use. Short-term effects in infants have not been reported, but a decrement in motor development at age 1 yr in the infants of marijuana-smoking mothers was reported in one study. Marijuana lowers serum prolactin slightly in nonlactating women and oxytocin release in rodents. One survey indicated that women who smoke marijuana breastfed for a shorter duration than nonsmokers and that the effect appears to be dose related.\(^1\)\(^2\) Avoid breastfeeding in heavy marijuana users and during therapeutic dronabinol use. Withhold breastfeeding for several hours after occasional marijuana use and use caution to avoid exposing the infant to marijuana smoke.

**Phencyclidine**

Phencyclidine is concentrated in milk and remains detectable in milk for weeks after heavy use. Avoid breastfeeding after phencyclidine use; a sufficient duration of abstinence has not been defined.

**Tobacco**

Nicotine and its metabolite, cotinine, are excreted into breastmilk in amounts proportional to the number of cigarettes smoked by the mother.\(^1\)\(^7\)\(^4\) The milk of smokers contains higher concentrations of cadmium than the milk of nonsmokers; other toxins from smoke have not been measured. Smokers also produce lower milk volumes, have lower milkfat content, use formula supplements more often, and wean their infants from breastfeeding earlier than nonsmokers, in part because nicotine lowers maternal basal prolactin concentrations.\(^1\)\(^7\)\(^5\),\(^6\) Infants of smoking mothers have increased infantile colic, large postnursing decreases in respiratory rate and oxygen saturation, and more respiratory infections.\(^1\)\(^7\)\(^4\) However, among infants of smokers, breastfeeding reduces the risk of respiratory illness by half that of formula-fed infants.\(^1\)\(^7\) In nonsmokers, breastfeeding reduces the risk of sudden infant death syndrome compared with formula feeding, but smoking negates this advantage.\(^1\)\(^7\)\(^8\) Advise nursing mothers to (1) stop or decrease smoking to the greatest degree possible, (2) not breastfeed right after smoking, and (3) not smoke in the same room with the infant.\(^1\)\(^7\) The use of nicotine chewing gum, topical patches, or nasal spray has not been studied during lactation. Although they are not recommended by the manufacturer during nursing, these products are likely to be less hazardous to the nursing infant than maternal smoking.

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DRUG USE IN SPECIAL POPULATIONS


DRUGS AND BREASTFEEDING

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Pediatric drug therapy presents a challenge to the practitioner in many respects. The pediatric population is comprised of a range of patient weights and organ maturity. Often there are no pediatric-specific data in the literature from which to derive appropriate dosage regimens. At times, medications must be used for which data are extrapolated on the basis of limited pharmacokinetic knowledge about the pediatric population. It must be remembered that children should not be treated as “little adults” when designing dosage regimens. Dosage administration nomograms derived from adult data should not be used in the pediatric population. Pharmacodynamic responses for the majority of medications used in children are even less well known. Children often react much differently from adults to certain medications. Examples are the use of stimulants such as methylphenidate to control hyperactivity common with attention deficit disorders and paradoxical hyperactivity, which can be observed in children taking phenobarbital. With therapeutically monitored medications, the standard adult therapeutic range is typically used because age-specific, concentration-effect information is scarce. Because of protein binding differences, infants might respond to lower total drug concentrations than those used in adults for certain medications (eg, phenytoin, theophylline).

One of the problems facing the clinician and caregiver of small children is the administration of medications. Dosage forms are usually designed with the adult population in mind, and the dosage cannot easily be individualized in small patients. This is especially true for most sustained-release products. Most young children cannot swallow tablets and capsules; thus, liquid preparations are generally preferred in this age group. For many drugs, liquid forms are not commercially available and must be extemporaneously compounded. Stability of these preparations is often unknown or of limited duration. Even when appropriate dosage forms suitable for young children are available, palatability, resistance to taking medications, and compliance issues can hinder optimal therapy.

## PHARMACOKINETICS

### ABSORPTION

At birth, gastric pH is neutral but falls to values of 1–3 in the first day of life. Subsequently, gastric pH returns toward neutrality because gastric acid secretion is low in the first several weeks to months. Adult values are usually achieved after the age of 2 yr.\(^1\,^2\) Medications that require gastric acidity for absorption can have poor bioavailability in this age group, rendering them ineffective or requiring much higher doses than normal for therapeutic serum concentrations to be reached. Examples of medications in this group are phenytoin, ketoconazole, and itraconazole.\(^1\,^3\) Alternative agents might have to be used if adequate serum levels cannot be documented when these drugs are administered orally. Certain medications that are acid labile actually might have increased bioavailability in infants, and these are antibiotics such as penicillin G and ampicillin.\(^4\)
Gastric emptying time can be delayed in infants, especially premature infants.\textsuperscript{1,3,5} Peak drug concentrations can occur much later in infants than in older children and adults. Other factors that can influence overall bioavailability of a particular medication in infants are the relatively high frequencies of gastroesophageal reflux, which can cause the dose to be spit up or vomited, and acute gastroenteritis (diarrhea), which can considerably shorten intestinal transit time. The oral route must be used with caution in these instances, especially in critically ill patients.

Other routes of administration can pose difficulties in the pediatric population. Overall muscle mass is decreased, and intramuscular administration might not be practical and certainly is not appreciated by most children. Most adults still remember their first injections in the doctor’s office when they were children. Also, the dose of drug to be administered might require multiple injections.

Rectal administration may be used in situations where the oral route is not practical or available; however, absorption might be incomplete and/or erratic. Topical administration of medications can lead to undesired systemic absorption, especially in infants in whom the skin thickness is less and the total skin surface area is proportionally greater than in adults.\textsuperscript{1,2,4}

DISTRIBUTION
Rapid changes in body composition can dramatically alter the $V_d$ for many medications during the first several months of life. Newborns have a higher percentage of total body water and extracellular fluid than older children and adults.\textsuperscript{1,3,6} Hydrophilic drugs such as the aminoglycosides have a much larger $V_d$ in newborns; this gradually decreases over the first year of life to approach adult values.

Total body fat in newborns (especially premature infants) is much lower than in older children and adults.\textsuperscript{6} Medications that are lipophilic might have a lower weight-adjusted $V_d$ in the very young.

Protein binding is an important determinant of the $V_d$ for drugs that are bound by albumin and other plasma proteins. In the neonatal period, the binding affinity of albumin is decreased compared with that in older children and adults (because of the persistence of fetal albumin).\textsuperscript{1–3} Highly protein-bound drugs such as phenytoin have higher free fractions in neonates, and there might be an increased pharmacodynamic response at lower concentrations of total drug. The $V_d$ of these drugs is inversely related to the degree of protein binding.

In addition, the clinician must be aware of the potential for highly protein-bound substances to displace bilirubin from binding sites on albumin, particularly in the newborn.\textsuperscript{1–3,7} The blood–brain barrier in newborns is more permeable than in older patients, and free bilirubin can readily cross into the CNS and cause kernicterus.

Tissue binding for many medications is unknown but can differ dramatically from that in adults. One example is digoxin, which binds to erythrocytes in pediatric patients to a much greater extent than in adult patients.\textsuperscript{2,4} Digoxin has a much larger $V_d$ in pediatric patients, and recommended loading doses in this age group are much larger on a mg/kg basis than in adult patients. In general, drug distribution volumes are larger in neonates and gradually approach adult values (in L/kg) by the first year of life.
METABOLISM

Metabolic processes show dramatic changes in the first weeks to months of life. At birth, most hepatic enzymes are immature and drug metabolizing capacity is greatly reduced. Phase I reactions (ie, oxidation) are controlled largely by the mixed-function oxidase system, of which the cytochrome P450 enzymes are the major determinant. These enzymes are largely undeveloped in newborns, especially premature infants, but maturation can take place quickly in the first weeks to months of life. Phase II reactions (ie, conjugation) include glucuronidation, sulfation, and acetylation. These reactions also are immature at birth, and drug toxicity has resulted (eg, with chloramphenicol) because of the absence of knowledge about reduced dosage requirements in newborns.1–3,6

The liver size relative to body weight in newborns is much larger than that in adults.1 Rapid weight gain, with subsequent increases in liver size and metabolic capacity, might require many dosage adjustments to prevent newborns from growing out of their dosages for many medications. When full metabolic capacity is reached in the pediatric patient, the hepatic clearance can greatly exceed that observed in adult patients on a weight-adjusted basis. Pediatric dosages of many medications on a mg/kg basis are often much greater than adult dosages. Figure 2–2 illustrates the change in clearance with age for theophylline.6 Most medications have similar curves but can be shifted to the left or have different relative peaks compared with adult values. A decrease in hepatic clearance relative to body weight typically begins after a child weighs approximately 30 kg.8 Thereafter, the increase in total body weight in proportion to liver size becomes greater. Thus, in adolescence, drug dosages typically begin to approach adult values. Drug toxicity can be observed in the adolescent patient if drug dosages on a mg/kg basis (designed for younger patients) are used.

RENAL ELIMINATION

The kidneys are the major route of drug elimination for many drugs. The kidneys are functionally immature at birth with regard to glomerular filtration and tubular secretion. Glomerular filtration at birth adjusted for body surface area is only

![Figure 2–2. Maturation of theophylline metabolism.](image-url)
30 to 40% of values in older infants and healthy young adults. Premature infants often have even lower values during the first few weeks of life. Dosages of many medications (eg, aminoglycosides, vancomycin) that are eliminated largely by glomerular filtration must be decreased on the basis of the relative immaturity of the kidneys at birth. Maturation of glomerular filtration occurs over the first several weeks to months of life. The dosages of most medications are similar to those in older children by age 4–6 months. Although the frequency of renal disease in children is much lower than in the adult population, factors that can alter renal function, such as shock, nonsteroidal anti-inflammatory drugs, or hypoxia, must be considered when evaluating dosage regimens. Serum creatinine, the usual marker for renal function, is usually lower in young children than in adults because of children’s lower muscle mass. Thus, a serum creatinine that indicates normal renal function in an adult might indicate renal impairment in a young child.

Tubular secretion also is diminished in the newborn. Drugs that have a component of tubular secretion (eg, penicillin) are typically administered at reduced dosages in the newborn. Maturation of tubular secretion occurs somewhat more slowly than glomerular filtration, but approaches adult values by age 8–12 months.

EVALUATING DRUG DATA IN CHILDREN

With the numerous maturational changes observed in children from birth through adolescence, results of pediatric drug studies must be used with caution in children whose ages differ from those in the study. Dosages extrapolated only on a weight basis have the potential to underdose or overdose other age groups, depending on the population studied. Body surface area might correlate better than body weight with total body water and extracellular water and can be useful in certain instances in calculating dosage regimens. With the exception of cancer chemotherapeutic agents, information on drug dosage is more widely available in mg/kg than by body surface area. Medications with narrow therapeutic ranges should have serum concentrations measured to aid in individualizing drug therapy, especially in critically ill children or those with known decreased renal or hepatic function.

Pharmacodynamic changes are poorly studied in the pediatric population, and responses to specific drug concentrations might be much different from those in the adult population. Diseases of childhood often differ from those in adults. Medications tolerated by adult patients might be inappropriate for the pediatric population (eg, aspirin for fever).

Caution must be used in the interpretation of drug levels because there might be much greater fluctuation in serum concentrations because of shorter drug half-lives in children than in adults. Further, the total volume of blood needed for drug level monitoring in small children can limit monitoring.

Detailed information on specific drugs can be found in the Pediatric Dosage sections of the individual drug monographs.

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Geriatric drug therapy is an important area of therapeutics and research, because of the growing elderly population, their disproportionately high use of medications, and their increased risk of drug misadventures. Although they represent approximately 12% of the U.S. population, the elderly consume more than 30% of all medications. Trends include increasing numbers of the extreme elderly (over age 80) and elderly with functional disabilities. It is estimated that the number of elderly who are dependent in their activities of daily living will triple from 1985 to 2060. Ethical considerations, such as a patient’s right to exercise decisions regarding treatment, are particularly relevant to the elderly population. As the number of elderly increases and health care resources diminish, cost–benefit considerations will become increasingly important.

The elderly are the most physiologically heterogeneous category of the adult population. The rate of normal aging varies considerably, and comparing data from persons of chronologically similar age can be misleading; health status is probably as important as age. Optimization of drug therapy in the elderly requires an understanding of how aging and concomitant pathology affect the pharmacokinetics and pharmacodynamics of drugs, the need to assess elderly patients individually, and elderly patients’ expectations of therapy.

Compliance issues leading to misuse and medication errors can be important in the elderly. The cost of medications, physical difficulty in opening medication containers, swallowing large tablets, reading the prescription label, and the presence of depression or cognitive impairment can contribute to compliance problems.

Adverse drug reactions are more common in the elderly, although the correlation with age alone is debatable. Increased medication use, especially medications with greater potential for toxicity, and chronic pathology with intermittent acute exacerbations are thought to contribute to the higher frequency and severity of adverse drug reactions. Most reactions in the elderly are dose related rather than idiosyncratic as a result of changes in pharmacokinetics and/or pharmacodynamics. Given the wide physiologic variability in the elderly population, the contribution of pharmacokinetic and pharmacodynamic changes can vary considerably. Additionally, the elderly are more sensitive to specific adverse reactions. For example, they have an increased sensitivity to anticholinergic side effects, especially central effects such as disorientation and memory impairment. These effects can be additive because many drugs commonly taken by the elderly are centrally active. Varying degrees of cognitive impairment or even delirium can be induced by drugs in several classes including benzodiazepines, centrally acting antihypertensive agents, and antidepressants. The onset can be insidious and mistakenly attributed solely to the aging process.
PHARMACOKINETICS

ABSORPTION

With aging there is some decrease in gastric secretions, acidity, gastric emptying, peristalsis, absorptive surface area, and splanchnic blood flow, although the effect on gastric pH may not be as pronounced as once believed. Taken together, the changes predict an altered extent or rate of absorption of orally administered drugs, yet most formal studies show no difference in oral bioavailability. Some factors might counterbalance each other (eg, acidity and gastric emptying; decreased absorptive surface and longer transit time). Some drugs (eg, digoxin) have shown a clinically unimportant slowed rate of absorption with equivalent quantities absorbed. Drugs with high extraction ratios may have increased bioavailability in the elderly compared with young patients, because of a decreased first-pass effect secondary to reduced hepatic blood flow. Decreased first-pass metabolism in the elderly has been shown for labetalol, propranolol, lidocaine, and verapamil.

22 It is known that the elderly have drier skin with lower lipid content, which is expected to be less permeable to hydrophilic compounds. Although neither conclusively nor well studied, percutaneous drug absorption appears to decrease with age.

DISTRIBUTION

Body weight generally decreases, but more important, body composition changes with age. Total body water and lean body mass decrease, while body fat increases in proportion to total body weight. The percentage of body weight contributed by fat changes from 18% and 33% in young men and women, respectively, to 36% and 45% in their elderly counterparts. These factors can alter the Vd of drugs in the elderly, although other aspects of drug disposition (binding, metabolism, elimination) can be additive or negate the effect. The Vd changes are most marked for highly lipophilic and hydrophilic drugs, and elderly patients are particularly susceptible to overdosage from drugs whose doses should be based on ideal body weight or lean body weight. Theoretically, highly lipidsoluble drugs (eg, long-acting benzodiazepines, lidocaine) may have an increased Vd and a prolonged effect if drug clearance remains constant. Conversely, watersoluble drugs (eg, gentamicin) may have a decreased Vd, and at least transiently increased serum levels, leading to possible toxicity if initial doses are not conservative. Although cardiac output does not appear to decrease with age, some chronic diseases affecting the elderly do contribute to a decrease in cardiac output and regional blood flow. There is some evidence that blood is preferentially shunted away from the liver and kidneys to the brain, heart, and muscles. These changes could explain the slowed elimination of some drugs and the heightened sensitivity to others.

PROTEIN BINDING

The proportion of albumin among total plasma proteins decreases with frailty, catabolic disease states, and immobility seen in many elderly, but it is no longer believed that serum albumin decreases with age alone. Serum albumin determinations should be performed to aid monitoring and dosage adjustment of drugs.
that are highly protein bound in the chronically immobile or ill elderly. A decrease in serum albumin can increase the percentage of free drug available for pharmacologic effect and elimination. Changes in albumin binding are more important with highly bound (greater than 90%) acidic drugs such as salicylates, phenytoin, and warfarin. Conversely, basic drugs, including lidocaine, propranolol, and meperidine, have affinity for α1-acid-glycoprotein, which may increase with age, especially when associated with conditions such as inflammatory diseases and malignancies. Protein binding theoretically may be increased and result in less free drug available, although the clinical relevance of this is unclear. With both types of binding, the net effect on clearance varies, depending on metabolism and elimination. Although not always available, free drug concentration measurements are often desirable in the elderly. There is also some evidence that the elderly may have a greater potential for protein displacement drug interactions.

METABOLISM
Liver size and hepatic blood flow decrease with age and especially with disease. Studies show hepatic blood flow decreases by 35%, and liver volume by 44% and 28% in elderly women and men, respectively, when compared to younger counterparts. Such a decrease in hepatic blood flow can limit the first-pass effect of drugs with high extraction ratios and markedly reduce their systemic clearance. Studies on phase I drug metabolism (ie, oxidation) do not consistently show a correlation with age, although most show that the elderly, especially men, have prolonged elimination. Differences may be explained by environmental factors such as smoking habits and genetics. Phase II metabolism (ie, conjugation) does not appear to be influenced as much by age, although there has been less study in this area. The effect of aging on drug acetylation is inconsistent and the importance unclear. There does not appear to be any age difference in the degree of inhibition or induction of cytochrome P450 isozymes. Monitoring and management of interactions with drugs such as cimetidine should be handled in the same manner as in younger patients. The changes described in liver size and metabolic function help to explain why certain drugs may have prolonged elimination; however, the variability of data cautions against generalizing about the effect of age alone. The initial dosage of metabolized drugs should be conservative and subsequent dosage adjustments based on careful monitoring of therapeutic and toxic parameters.

RENA L ELIMINATION
The effect of aging on the renal elimination of drugs is probably the most completely understood and important aspect affecting geriatric drug therapy. Glomerular filtration, tubular secretion, and renal blood flow all decrease with age. Creatinine clearance decreases approximately 1% per year after age 40, the effect is variable, and volume depletion, CHF, and renal disease can further decrease organ function. Because creatinine production also decreases with age, serum creatinine may be normal despite a substantial decrease in renal function. It is therefore recommended that Clc be measured or estimated using a method that
incorporates age and weight. The dosage of renally excreted drugs with low therapeutic indices should be conservative initially, with subsequent dosage titrated by close clinical and serum drug level monitoring, if applicable.

PHARMACODYNAMICS

Heightened drug effects that cannot be explained by altered pharmacokinetic variables alone have been hypothesized to be caused by changes in compensatory homeostasis, drug receptor sensitivity, or complications of chronic diseases that occur in the elderly. There is a gradual decrease in homeostatic reserve with aging. Postural control and orthostatic circulatory response are examples of compensatory mechanisms that are slowed in aging. Adequate postural blood pressure control relies on several factors, including central coordination, muscle tone, and proprioception, all of which can be blunted in the elderly. As a result, side effects that are minimal or absent in a young patient with normal compensatory response can be marked in the elderly. The administration of long-acting anxiolytics, hypnotics, or antipsychotics can further alter these mechanisms and lead to an increased risk of falls in the elderly. Similarly, symptomatic postural hypotension can result from the administration of a variety of antihypertensive agents (especially calcium-channel blockers and ACE inhibitors) and other drugs (eg, antipsychotics, antidepressants) that affect vasomotor tone. Physiologic mechanisms such as vasoconstriction and tachycardia cannot fully compensate for postural hypotension in the elderly. Temperature regulation and intestinal motility are other homeostatic mechanisms that change with aging and can explain heightened effects of certain drugs.

The number and characteristics of drug receptors can change with aging and produce altered, often heightened, drug response. Research has shown age-related decreases in several autonomic receptors. There is some evidence of increased sensitivity to oral anticoagulants and digoxin, apart from the alterations in pharmacokinetics, which might contribute to the higher frequency of adverse reactions to these two agents in the elderly.

Preliminary data indicate a possible increase in brain sensitivity to certain drugs with aging. It is unknown whether this effect is caused by changes in blood-brain permeability or tissue receptor sensitivity. More research into drug pharmacodynamics in the elderly is needed, especially the interrelationship with pharmacokinetic alterations. The presence and impact of multiple concurrent pathologies and their treatments cannot be overemphasized in their contribution to the various drug effects seen in the elderly.

OTHER FACTORS

Cigarette smoking can cause clinically important induction of the metabolism of some drugs to a similar degree in both the elderly and the young. This, and the fact that many published studies do not indicate smoking history, could explain some interpatient variability of pharmacokinetic data.
Nutritional intake is sometimes diminished in the elderly and can lead to nutritional and vitamin deficiencies. Nutritional status of the elderly can impact the outcome of drug therapy, and, conversely, drug therapy can affect nutritional status.\textsuperscript{19,36,37}

\section*{Evaluating Drug Data for the Elderly}

Because of age-related changes that may impact the outcome of drug therapy as outlined in this chapter, the results of drug studies using young subjects cannot always be extrapolated accurately to the elderly. Studies on diseases and drugs in the elderly do not always include sufficient numbers of elderly, especially extremely aged subjects, to draw appropriate conclusions.\textsuperscript{38} Studies that include the elderly do not always separate results by decade of age and health status, two criteria that are helpful in assessing applicability of data in this heterogeneous population. Many studies also do not mention data on nutrition, alcohol, and smoking, which might explain some variability of results.\textsuperscript{19} Although single-dose studies in healthy volunteers can be useful, long-term studies in afflicted elderly patients often yield data more applicable to therapeutics. Drugs are often not studied over a wide dosage range, so a minimal effective dosage in the elderly cannot be determined.\textsuperscript{39}

When reviewing studies that include the elderly, one should consider the following potential problems: numbers of subjects must be sufficient to allow for high attrition rates and the typically wide variation in this population; study lengths must be sufficient for a chronic disease; concomitant diseases and medications must be acknowledged and their impact assessed; and “normal” values can be different from those of a younger population.\textsuperscript{40–42}

\section*{Conclusion}

The effects of aging as related to drug therapy illustrate the challenges in caring for the elderly. Clinical practice guidelines that have been developed for conditions commonly afflicting the elderly, such as those published by the Agency for Health Care Research and Quality, can be a helpful guide.\textsuperscript{43} Conservative dosage, especially initially, with close clinical monitoring for dose-dependent effects is critical and should be emphasized by all health care practitioners caring for the elderly. For detailed information on specific drugs in the elderly, refer to the Geriatric Dosage section of the individual drug monographs.

\section*{References}

Renal Disease
Gary R. Matzke

DOSAGE REGIMEN OPTIMIZATION FOR PATIENTS WITH RENAL INSUFFICIENCY

Eleven million Americans have early renal insufficiency, defined as a Cr ≥1.5 mg/dL or CrCl < 70 mL/min; approximately 1 million have concentrations > 2 mg/dL. The number of individuals with end-stage renal disease (ESRD) has been increasing at a rate of about 7–9% annually during the past decade.

Reduced renal function can be associated with drug effects, age, or chronic disease states. Medical problems can contribute to the development of a patient’s initial renal injury, enhance the rate of their progressive decline in renal function, or develop as sequelae of chronic renal disease. Hypertension, diabetes mellitus, infection, bone disease, neurological dysfunction, GI disturbances, and bleeding abnormalities are but a few of the medical conditions frequently encountered in renal failure patients. These patients are often given medications early in the course of their disease in an attempt to slow the rate of decline in renal function and prevent cardiovascular complications. Surveys of dialysis patients have found that they average more than eight scheduled prescription drugs and two or more “prn” drugs. Thus, patients with early renal insufficiency and those with ESRD are at increased risk for adverse reactions because of the number of drugs received, concurrent medical problems, and impaired drug excretion.

Renal insufficiency in any patient requires that the clinician understand the aspects of drug disposition that are altered and the appropriate methods to individualize drug therapy. Complications of drug administration can be minimized by the application of pharmacokinetic and pharmacodynamic principles. The advent of specific and sensitive methods for measuring drug concentrations in biological fluids has resulted in a voluminous literature on drug disposition in renal disease and evaluations of the effects of dialysis.

This section provides a conceptual discussion of how renal disease alters drug disposition with selected literature examples. It also describes an approach for determining the individual dosage adjustment necessary to achieve the optimal therapeutic effect with minimal toxicity for a patient with a given degree of renal function. The subsequent section, Dialysis of Drugs, presents the concepts of drug removal by hemodialysis, peritoneal dialysis, and continuous renal replacement therapies that are now frequently used in critically ill patients. Data on the amount of drug removed by dialysis are tabulated and dosage modification schemes for a number of drugs during dialysis are presented.

FOUR BASIC QUESTIONS

A practical approach to drug therapy in patients with renal insufficiency can be arrived at if one considers the following questions:
1. What is the patient’s renal function status?
2. What is the degree of alteration in the pharmacokinetics or pharmacodynamics of the patient’s drug(s) in the presence of renal insufficiency?
3. What approaches to dosage modification are useful for a specific drug?
4. What is the impact of dialysis on drug disposition, and is dosage modification or supplementation necessary?

**QUANTIFYING RENAL FUNCTION**

Several common laboratory tests provide an assessment of a patient’s renal function: blood urea nitrogen (BUN), serum creatinine (Cr), the ratio of BUN to Cr, and creatinine clearance (Clcr). The BUN concentration can change because of many factors in addition to changes in renal function. Urea is filtered and reabsorbed by the nephron, and its renal excretion is a function of urine flow. Diuretic use, dehydration, and bleeding can increase the BUN concentration without a decline in renal function. These conditions usually result in an increased BUN/Cr ratio to values above the normal range of 10–15. Creatinine production and elimination in adults are usually constant at approximately 20 mg/kg/day under steady-state conditions. Creatinine is filtered predominantly by the glomerulus with little renal tubular secretion (about 10%) in those with normal renal function. However, secretion becomes an important excretory pathway for patients with a Clcr <50 mL/min. In these individuals, accurate measurement of Clcr can be obtained by giving cimetidine before initiating the urine collection because cimetidine inhibits the tubular secretion of creatinine.

Because the nonrenal factors that can affect BUN do not alter serum or urine creatinine concentrations, Cr and Clcr serve as better markers of changing renal function. The relationship between Cr and Clcr is a hyperbolic one, as is shown in Figure 2–3. Small increases in Cr represent a larger absolute decrease in renal function in subjects with normal renal function than do similar increases in Cr in individuals with moderate to severe renal insufficiency. For example, doubling the Cr is associated with a halving in Clcr (ie, as Cr changes from 1 to 2 mg/dL, the Clcr declines from 120 to 60 mL/min, whereas an increase from 2 to 4 mg/dL represents a decrease in Clcr of 60 to 30 mL/min).

Although Cr is easy to determine, requiring collection of only a single blood sample, measurement of Clcr is more difficult. The standard method consists of a continuous 24-hr urine collection for urine creatinine with a single blood sample for Cr at approximately the middle of the urine collection period. The most difficult problem from a practical standpoint is obtaining a complete urine collection. Almost invariably, the urine collection is incomplete and consequently the Clcr is underestimated. However, the accuracy of the Clcr and urine collection can be assessed by determining the daily creatinine excretion rate—the amount of creatinine (in mg/day) excreted in the urine during the 24-hr collection period. This can be compared with the expected amount of creatinine to be excreted, which is approximately 20–25 mg/day/kg ideal body weight in males and 15–20 mg/day/kg in females who are age 18–50 yr. Urinary creatinine excretion declines in males and females who are >50 yr. For example, in a 70-kg man, the expected creatinine production and excretion in the urine is approximately 1.4 g/day. If his total creatinine excretion is less than this value, it is likely he did not collect all his urine and...
the calculated Cl\text{cr} is an underestimate. The use of this approach is valid only under steady-state conditions when creatinine production and excretion are equivalent. If it is impractical to measure a patient’s Cl\text{cr}, it can be estimated from equations based on the patient’s age, height, and weight. The most frequently used equation for adults with stable renal function was derived by Cockcroft and Gault. Equations are given for men, women, and children in Appendix 2, Anthropometrics. These equations assume steady-state serum creatinine values and do not provide valid Cl\text{cr} estimates in patients with fluctuating renal function or those receiving dialysis of any type. The advantages and disadvantages of the several methods for Cl\text{cr} estimation in patients with changing renal function have been reviewed recently.

\section*{PHARMACOKINETIC/PHARMACODYNAMIC ALTERATIONS OF DRUGS IN RENAL FAILURE}

Decreased renal function can alter the absorption, distribution, protein binding, metabolism, or excretion of drugs. Furthermore, the pharmacodynamic effects of a drug can be different in patients with renal insufficiency because of biochemical or pathophysiologic changes associated with renal disease. The bioavailability of drugs can be altered in symptomatic (uremic) patients because of GI disturbances such as nausea, vomiting, and diarrhea, increased gastric pH because of the ingestion of histamine H\textsubscript{2}-receptor antagonists, or increased salivary urea concentration as a result of markedly increased BUN (>100–120 mg/dL). This can decrease the...
absorption of ferrous sulfate and other drugs that are best absorbed from an acidic environment. In addition, patients who routinely take aluminum or calcium antacids might have reduced bioavailability of some drugs because of complexation in the GI tract. Propoxyphene, dihydrocodeine, and some β-blockers might have increased bioavailability because of reduced first-pass metabolism.24–26

The plasma protein binding of some drugs is altered in patients with severe renal insufficiency. This might be secondary to hypoalbuminemia; accumulation of acidic byproducts of uremia resulting in competitive displacement of drugs from binding sites; or changes in the structure of albumin resulting in a decreased number of effective binding sites.3,22,27 Most weak organic acid drugs, such as cefazolin, phenytoin, salicylate, valproic acid, and warfarin, exhibit decreased plasma protein binding (increased free fraction). Weak organic basic drugs might have decreased or unchanged binding. The protein binding of carbamazepine, dapsone, diazepam, and morphine is decreased, whereas the binding of propranolol, quinidine, verapamil, and trimethoprim is unchanged. Propranolol and lidocaine are bound primarily to α1-acid glycoprotein from which little displacement occurs in renal disease or hypoalbuminemia. If the protein binding of a drug is decreased, the patient can experience an increased pharmacodynamic effect, an increased \( V_d \), and increased or unchanged total body clearance depending on whether it is a high or low extraction ratio drug.

Phenytoin has altered protein binding and disposition in ESRD patients that results in important differences in dosage.28 The percentage of unbound phenytoin in plasma is normally 10% but increases to 20 to 35% in ESRD patients. This results in an increase in the \( V_d \) from 0.65 L/kg in those with normal kidney function to 1–1.8 L/kg in ESRD patients. Further, the terminal half-life is decreased from 11–16 hr to 6–10 hr and the apparent plasma clearance increases from 28–41 mL/hr/kg to 64–225 mL/hr/kg in ESRD patients compared with those with normal renal function. These changes in the pharmacokinetics of phenytoin result in a change in its therapeutic concentration range. In patients with normal kidney function, the usual therapeutic plasma concentration range for total phenytoin (unbound plus bound) is 10–20 mg/L; in those with ESRD, the range is approximately 4–8 mg/L. Both ranges of total drug represent the same concentration of unbound drug, 1–2 mg/L.

The \( V_d \) of drugs can be increased, decreased, or unchanged in renal failure patients.3,22,27 An increase in \( V_d \) could be due to decreased protein binding, fluid overload secondary to reduced renal excretion, or increased tissue binding. A decrease in \( V_d \) could be due to decreased tissue binding or increased protein binding. Examples of drugs with increased \( V_d \) are cefazolin, furosemide, gentamicin, naproxen, phenytoin, and vancomycin. Digoxin exhibits a decreased \( V_d \) in renal impairment, whereas minoxidil and procainamide are drugs whose \( V_d \) does not change markedly in ESRD.

Drugs are eliminated from the body by two primary pathways: renal and nonrenal elimination (predominantly hepatic metabolism).39 The degree of reduction in renal clearance depends on the percentage of drug excreted unchanged by the kidney. The influence of renal function is very important for aminoglycosides, cephalosporins, penicillins, vancomycin, acyclovir, lithium, and ranitidine, all of which are extensively (>80%) eliminated unchanged renally. For many of these drugs, linear correlations have been established between the drug’s plasma and
renal clearance and Clcr. These correlations can be used as guides to project the drug dosage requirement for those with a given degree of renal insufficiency. For example, the linear correlation between gentamicin plasma clearance and Clcr demonstrates that the clearance of gentamicin can change from 120 mL/min with normal renal function to as little as 2 mL/min in ESRD. The half-life of gentamicin in ESRD is markedly prolonged (range 40–60 hr) compared with the 1–2 hr values of patients with normal renal function. This relationship then can be used to determine the desired maintenance dosage of gentamicin in patients with different degrees of renal insufficiency, as outlined later in this chapter.

Drug metabolism typically involves enzymatic conversion of drugs to more water-soluble compounds. These metabolites are formed through the processes of oxidation, reduction, synthesis (eg, conjugation), or hydrolysis. Once formed, these metabolites often are excreted predominantly by the kidney. Most metabolites are inactive or have minimal pharmacologic activity. However, some active metabolites might accumulate, especially in ESRD patients, and lead to exaggerated pharmacodynamic responses that warrant dosage reduction. Active metabolites that are excreted by the kidney include oxypurinol from allopurinol, which is an active inhibitor of xanthine oxidase; desacetylcefotaxime from cefotaxime, which is microbiologically active; normeperidine from meperidine, which can cause seizures; and N-acetylprocainamide from procainamide, which has its own unique antiarrhythmic properties.

Renal insufficiency also can lead to alterations in drug metabolism. Animal experiments indicate that the activity of many drug metabolic pathways is reduced in the presence of renal insufficiency by up to 70%. The decrement in enzyme activity is larger in the animals with the most severe renal dysfunction. The nonrenal clearance of several drugs is decreased in ESRD. For example, the antiviral agent acyclovir and the antihypertensive agent captopril demonstrate 50% decreases in nonrenal clearance in patients with ESRD. As a consequence, the elimination half-life for both drugs is increased 6-fold in the presence of renal failure. This shows that predictions of the disposition of drugs in renal failure based on general principles and nomograms are subject to considerable error if one assumes that nonrenal clearance is unaffected by renal disease.

The pharmacodynamics of a drug also can be altered in ESRD and result in the pharmacologic effects being different from those one would expect in patients with normal renal function. One well-defined example is that of nifedipine, where marked differences in Emax (maximal change in diastolic blood pressure) were observed. The average Emax values were 12 and 29% in healthy controls and ESRD patients, respectively. Thus, at the same plasma concentration of unbound nifedipine, a greater blood pressure reduction occurs in patients with renal insufficiency.

### Dosage Adjustment Approaches that are Useful and Practical for Specific Drugs

The general approaches for dosage adjustments of drugs in renal insufficiency are to (1) decrease the dose and maintain the usual dosage interval, (2) lengthen the dosage interval and maintain the usual dose, or (3) modify the dose and interval. The primary goal of these approaches is to provide average steady-state plasma
concentrations or AUCs in renal insufficiency similar to those in normal kidney function. The choice of approach depends on the type of drug and the desirability, from a therapeutic or toxic standpoint, of having small or large peak-to-trough fluctuations.\textsuperscript{38,39} Other considerations are that the dosage regimen adjustment should be practical and the reduced dose or prolonged dosage interval should be relatively easy to implement.\textsuperscript{14,27}

When presented with a patient with renal insufficiency for whom drug dosage regimen decisions must be made, the most practical and efficient approach is to first consult published tables or guidelines that provide a quick reference source for drug dosage in renal failure.\textsuperscript{14} Additional sources are the appendices of \textit{Handbook of Drug Therapy in Liver and Kidney Disease}\textsuperscript{29} and \textit{“Use of Drugs in Renal Failure”} in \textit{Diseases of the Kidney},\textsuperscript{40} which describe specific pharmacokinetic alterations of drugs in kidney disease, recommended dosage regimens, and tabulate the effects of dialysis. The reader also is advised to refer to specific drug monographs in this book and in \textit{AHFS Drug Information},\textsuperscript{41} which briefly describe the effect of renal failure on drug disposition and provide initial dosage recommendations. These sources allow the user to determine whether dosage adjustments are necessary and if there are any important toxicities or precautions in using a particular drug. These sources, however, provide only general guidelines.

For drugs requiring marked dosage adjustment in renal insufficiency or for which the achievement of specific therapeutic plasma concentrations is critical, the reader should consult the original publications, which provide specific data on individual drugs. Consulting the original publications or authoritative reviews will provide details regarding the relationship of renal function to drug elimination and can provide a dosage nomogram or specific dosage recommendations and precautions for the use of the drug in patients with various degrees of renal insufficiency.\textsuperscript{17}

If drug-specific data or guidelines are not available, one can use general dosage equations such as those developed by Rowland and Tozer.\textsuperscript{42} Only basic pharmacokinetic information about the drug is needed—primarily the fraction of the available dose that is normally excreted unchanged in the urine, $f_e$. The fraction of normal renal function (KF) in a given patient is determined as the ratio of the patient’s Cl\textsubscript{cr} to the accepted normal value of 120 mL/min. The patient’s Cl\textsubscript{cr} can be determined from Cr\textsubscript{s} values at steady state with the method of Cockcroft and Gault.\textsuperscript{21}

The following equation, which takes into consideration the renal clearance and extrarenal clearance of unbound drug, can be used to determine the dosage adjustment factor, Q:\textsuperscript{42}

$$Q = \frac{[(KF \times f_e) + (1 - f_e) \times ((140 - \text{age}) \times \text{weight in kg}^{0.7})]}{1660}$$

Q is analogous to the ratio of the unbound drug clearance of the patient to that observed in those with a Cl\textsubscript{cr} \geq 120 mL/min, or $\frac{Cl_e \text{ (failure)}}{Cl_e \text{ (normal)}}$. If a drug is minimally protein bound ($<25\%$) and not extensively metabolized ($f_e \geq 70\%$), this equation can be simplified to:\textsuperscript{5}

$$Q = 1 - [f_e (1 - KF)]$$
Once the value of $Q$ is obtained, the dosage regimen adjustment can be made with the following equations and scenarios:

$$D_{RI} = Q \times D_N$$

where $D_{RI}$ is the maintenance dose in the renally insufficient patient that is to be given at the normal dosage interval and $D_N$ is the normal dose for those with $\text{Cl}_{cr} \geq 120 \text{ mL/min}$. 

$$\tau_{RI} = \frac{\tau_N}{Q}$$

where $\tau_{RI}$ is the maintenance dosage interval for the renally insufficient patient at which the $D_N$ is the dose to be given and $\tau_N$ is the dosage interval for those with $\text{Cl}_{cr} \geq 120 \text{ mL/min}$. 

The final scenario incorporates a modification of $D_N$ and $\tau_N$. This scenario usually is used when the calculated $D_{RI}$ or $\tau_{RI}$ are impractical. In that situation, one chooses a clinically relevant $\tau_{RI}$ and calculates the $D_{RI}$ to be given at that time.

$$D_{RI} = \frac{[D_N \times Q \times "\tau_{RI}" ]}{\tau_N}$$

An example will clarify the use of this approach. An 80-kg, 45-yr-old man with a $\text{Cr}_r$ of 5.4 mg/dL requires treatment with ceftazidime for a pseudomonal infection. This drug is 70% excreted unchanged in the urine, and the usual dosage is 1 g q 8 hr IV. With the equation of Cockcroft and Gault, the patient’s $\text{Cl}_{cr}$, KF, and $Q$ are calculated as follows:

$$\text{Cl}_{cr} = \frac{(140 - 45) \times 80}{(5.4 \times 72)} = 20 \text{ mL/min}$$

$$\text{KF} = \frac{20 \text{ mL/min}}{120 \text{ mL/min}} = 0.17$$

$$Q = (0.17 \times 0.7) + (1 - 0.7)$$
$$= 0.120 + 0.3$$
$$= 0.43$$

If the maintenance dose for this patient was reduced and the dosage interval maintained every 8 hr, the $D_{RI}$ would be:

$$D_{RI} = 0.43 \times D_N$$
$$= 430 \text{ mg q 8 hr}$$

This regimen will result in reduced peak and increased trough concentrations relative to subjects with normal renal function receiving the standard dose. The average concentration would, however, be the same.
Alternatively, one might extend the dosage interval and maintain the standard dose size ($D_N$). This will produce the same peak and trough concentrations for the renal patient that one would expect in a patient with $Cl_{cr} \geq 120$ mL/min. Unfortunately, the use of nonstandard dosage intervals often has been associated with drug administration errors.

$$\tau_{RI} = \frac{\tau_N}{Q} = \frac{8}{0.43} = 18.6 \text{ hr}$$

In this example and many patient scenarios, the best dosage adjustment strategy might be to select a feasible prolonged dosage interval ("$\tau_{RI}$"), eg, 12 hr, and then calculate the $D_{RI}$.

$$D_{RI} = \frac{[D_N \times Q \times \"\tau_{RI}\"]}{\tau_N} = \frac{[1000 \text{ mg} \times 0.43 \times 12]}{8} = 650 \text{ mg}$$

This general approach provides a reasonable initial method for adjusting drug dosage regimens in patients with renal insufficiency until more specific guidelines can be consulted or serum concentrations are measured. This method is based on several assumptions: (1) bioavailability is unchanged in renal failure; (2) metabolites are not therapeutically active or toxic; (3) decreased renal function does not alter metabolism of the drug; (4) metabolism or renal excretion does not exhibit concentration-dependent pharmacokinetics; (5) renal function is constant with time; and (6) the renal clearance of the drug is directly proportional to the renal clearance of the compound used to measure renal function. If any of these assumptions is invalid, the accuracy of the projected dosage regimen will be reduced.

The time to reach steady state is longer for a patient with renal insufficiency than one with normal renal function. Consequently, it is common to initiate therapy for many drugs with a loading dose (ie, at least the $D_N$ and in some cases an even greater dose) to achieve the desired concentration in the expanded $V_d$ and/or shorten the time to reach a therapeutic plasma concentration. The amount of the loading dose depends on the particular drug being used and the desired therapeutic objectives.

It should be noted that any dosage regimen modification for renally insufficient patients might require plasma concentration determinations of the drug, if available, and close clinical observation for assessment of toxicity and verification of achievement of the desired therapeutic serum concentrations or effects.

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DIALYSIS OF DRUGS

DIALYSIS REMOVAL OF DRUGS AND DOSAGE SUPPLEMENTATION

In the United States, more than 250,000 patients with ESRD receive chronic maintenance hemodialysis or continuous ambulatory or cycling peritoneal dialysis. Further, 8000–10,000 patients each year receive a kidney transplant. Therapeutic advances in immunosuppressive drugs and dialysis techniques have increased survival, and a reasonable quality of life is now possible for many patients with ESRD. These patients are managed primarily with hemodialysis, but the use of continuous ambulatory and/or continuous cycling peritoneal dialysis is now common. Although the purpose of dialysis is to remove unwanted toxic waste products from the body, it also removes many pharmacotherapeutic agents. Thus, it is important to know to what extent a drug is removed by dialysis because this can affect the patient’s therapy. Supplemental doses or a revised dosage regimen might be required if the dialysis procedure markedly augments the patient’s clearance of the drug. Dialysis procedures, including hemoperfusion, also have been used in drug overdose situations as a means of enhancing drug removal from the body. Therefore, it is important to consider how effective these procedures are and whether they offer any substantial advantage over conventional means of treating overdoses.

The objectives of this section are to review those drug and dialysis prescription factors that affect the efficiency of the removal of drugs by dialysis. The impact of hemodialysis on the disposition of a drug can be quantified by direct measurement of the clearance by the dialyzer. This provides the most accurate assessment of dialyzability and requires one to assay the dialysate and/or obtain multiple blood samples. Alternatively, one can assess the clearance or half-life during dialysis just by collecting 2–3 blood samples. These pharmacokinetic data can be used to estimate the fraction of the drug removed by dialysis from the patient during the dialysis procedure.

Peritoneal dialysis is a much less efficient means of removing waste products and drugs than hemodialysis. The additional clearance provided by this mode of dialysis is only of clinical significance for a few agents: aminoglycosides, some antifungals, phenobarbital, theophylline, and vancomycin. In contrast, the continuous renal replacement therapies, which are frequently used in critically ill patients, can dramatically alter disposition of many drugs used in the acute care setting. These therapies are now widely used and employ hemofilters that are made of the same materials as hemodialyzers. Thus, the dialyzer filter factors, which influence drug removal, are similar. However, the patient factors are not affected as dramatically because these patients usually have some residual renal function.
DETERMINANTS OF DRUG DIALYZABILITY

The Dialysis Prescription. The literature on drug dialyzability dates back to the early 1970s. Since then there have been several marked improvements in hemodialysis therapy. The blood and dialysate flow rates, two of the primary determinants of the removal of uremic waste products and drug, have increased from 200–300 and ~500 mL/min in the 1970s and much of the 1980s to 400–600 and ~700–1000 mL/min in the 1990s. As a result, the clearance of many drugs has been markedly increased. Further, in the 1990s, a major shift in the composition of dialyzer membranes occurred. Before the 1990s, >80% of ESRD patients were dialyzed with conventional dialyzers: cellulose, cuprophane, or slightly modified cellulose filters. By 2000, >75% of ESRD patients received hemodialysis with semisynthetic or synthetic dialyzers: polysulfone, polymethylmethacrylate, polyamide, or cellulose triacetate. The clearance of waste products, BUN, creatinine, and all drugs evaluated to date are much higher than reported in the literature. In fact, some drugs such as vancomycin, which was not dialyzable with conventional dialyzers, is now highly dialyzable with these new filters. For other agents such as the aminoglycosides and cephalosporins, increases in clearance of 200 to 300% are common.

In light of these dramatic improvements in hemodialysis, earlier data might no longer be applicable. Thus, one must accurately document what dialyzer is used, what the blood and dialysate flow rates are, and the duration of the dialysis procedure. With this information, one can begin the literature search for data on dialyzer clearance of the drugs of concern. If only older, conventional data are available, it will be necessary to extrapolate to the current situation, usually resulting in an increase in the projected impact of hemodialysis.

Pharmacokinetic Factors. Certain drug properties can help predict drug dialyzability. Drugs with low molecular weights, usually <500 d, cross conventional dialysis membranes readily. Large-molecular-weight drugs, such as vancomycin (MW 1449) and amphotericin B (MW 924), cross those membranes poorly and are not effectively removed by conventional hemodialysis. The use of semisynthetic and synthetic dialyzers, however, allows for removal of large-molecular-weight compounds such as vancomycin and has been associated with marked increases in the clearance of low-molecular-weight agents.

Drugs with high water solubility are removed more easily to the aqueous dialysate than more lipid-soluble compounds. In addition, the latter usually have a larger V_d than more water-soluble drugs. Drugs such as digoxin or tricyclic antidepressants, which have V_d >5–7 L/kg, are usually minimally removed by dialysis because the majority of drug is contained in tissue compartments rather than in the blood, and the drug in these tissues is not as readily accessible for removal. A large V_d and slow transport between tissue and blood also can limit the use of hemodialysis and hemoperfusion. Although one can rapidly clear the blood compartment of a drug (shown by a dramatic decrease in plasma concentrations once the procedure has ended), plasma drug concentrations can increase (rebound) as a result of re-equilibration of drug from tissue stores.

Plasma protein binding of a drug also determines how effectively it can be dialyzed. Drugs with a high degree of protein binding, such as propranolol (90–94%) and warfarin (99%), are poorly removed by dialysis because the
drug–protein complex is too large to cross all dialysis membranes. This is not a limitation of hemoperfusion because the drug is removed from plasma proteins as the complex passes through the high surface area absorbent material.4

**LITERATURE EVALUATION PITFALLS**

Several problems are encountered when attempting to assess the literature on removal of drugs by dialysis.2,26 First, for some drugs, only anecdotal reports are available. This is primarily true in the overdose setting in which the effect of dialysis on drug removal often is determined primarily by clinical response.3,5 For example, a comatose patient awakens during or shortly after dialysis and it is assumed that dialysis removed the drug, accounting for the improved clinical status. Second, the amount of drug ingested and/or the amount of drug recovered in the dialysate often is unknown. Third, the type of dialysis system employed is frequently not specified—this is extremely important when comparing the patient’s dialysis prescription with published data. Advances in dialysis technology make much of the data evaluating conventional methods inapplicable. Fourth, there often is a lack of patient data, such as weight, hematocrit, and renal and liver function. Fifth, the method used to calculate drug clearance often is unspecified. For example, was clearance determined from the amount of drug recovered in the dialysate or from differences in arterial and venous plasma concentrations across the dialyzer? The proper method for clearance calculations in hemodialysis has been described.2,6,27

A common error is misinterpretation of plasma drug concentrations obtained before and after dialysis. Declining plasma concentrations during dialysis often are believed to be the result of the dialysis procedure. However, a declining concentration might be due to drug elimination by metabolism or renal excretion, and the contribution of dialysis to this decline might be very small. The situation in which drug concentrations are relatively unchanged during dialysis usually means that little or no drug is being removed by dialysis. However, the drug might continue to be absorbed from the GI tract as dialysis is being carried out, as in the delayed and prolonged absorption observed in drug overdose.3 Another problem is interpreting drug removal rate by dialysis. If 200 mg of a drug was removed in the first hour of dialysis, one might assume that 5 hr of dialysis would remove 5 times as much (ie, 1000 mg). This is incorrect because drug removal by dialysis occurs by a first-order process; as the amount of drug in the body declines, so does the amount removed per hour. Thus, the total amount removed is less than that calculated from the initial estimates.

For many drugs, there is a lack of correlation between plasma drug concentration and clinical response. Some drugs have been found to have active or toxic metabolites that correlate well with the toxic effects of the drug.3 In attempting to collect information on dialysis removal of drugs, attention must be given to metabolites as well. In the overdose and critically ill patients, the pharmacokinetic disposition of a drug can be altered.9 In making predictions of drug dialyzability, pharmacokinetic data are usually derived from healthy subjects receiving therapeutic dosages. However, in critically ill patients, especially those who have ingested an overdose, there might be changes in drug metabolism, Vd, or protein binding. For example, large amounts of drug in the body might saturate plasma protein binding, which in turn might alter drug distribution and metabolism.
ESTIMATION OF THE IMPACT OF DIALYSIS

The clearance of a drug by dialysis can be compared with the clearance of the drug by the body because clearance terms are additive:

\[ Cl_{TD} = Cl_T + Cl_D \]

where

- \( Cl_{TD} \) = total body clearance of drug during dialysis
- \( Cl_T \) = patient’s residual total body clearance of drug
- \( Cl_D \) = dialysis clearance of drug

If dialysis clearance adds substantially to total body clearance, thereby forming a much larger total clearance, then the drug will be eliminated that much faster. For example, if the dialysis clearance of a drug is 50 mL/min and the body clearance is 50 mL/min, then the drug would be eliminated from the body twice as fast during the dialysis period. To relate clearance to drug half-life (\( t_{1/2} \)), the following equations are useful:

- \( t_{1/2} \) (off dialysis) \[ t_{1/2} \text{ (off dialysis)} = \frac{0.693 \times V_d}{Cl_T} \]

- \( t_{1/2} \) (on dialysis) \[ t_{1/2} \text{ (on dialysis)} = \frac{0.693 \times V_d}{Cl_T + Cl_D} \]

The more the dialysis clearance adds to the patient’s residual total body clearance, the shorter the drug half-life will be on dialysis (assuming \( V_d \) remains constant). Another extension of this allows one to calculate the fraction of the drug in the body that is lost during a dialysis period.

\[ \text{Fraction Lost} = 1 - e^{-\left( Cl_T + Cl_D \right) \left( \tau_d / V_d \right)} \]

where \( \tau_d \) is the duration of the dialysis period.

This calculation represents the fraction of drug in the body that is lost during a dialysis period by all routes of elimination (ie, dialysis, metabolism, and renal excretion). It is necessary to acquire from literature sources (keeping in mind the limitations discussed previously) values for \( V_d \), \( Cl_T \), and \( Cl_D \). If renal or liver function is diminished, this must be taken into consideration because it will result in a lower \( Cl_T \). In addition, changes in \( V_d \) in certain disease states (eg, the decreased \( V_d \) of digoxin in renal failure) also must be taken into account.

Clearance data are not always available in the literature. Many reports, especially those published in the 1970s and 1980s, contain only the half-lives of the drugs, on and off dialysis. The following equation can be used to estimate the fraction of drug removed by dialysis alone with half-life data.
The assumptions made when using this equation are that all drug elimination (including dialysis removal) occurs by first-order processes, and dialysis is initiated after the completion of the absorption and distribution phases. The primary limitations of this equation are that inaccurate values of half-lives result in incorrect estimates of drug removal by dialysis and the fact that drugs might be given during dialysis. Because drug concentrations in plasma are higher in the distribution phase, especially for intravenously administered drugs, more drug can be removed by dialysis than one would predict by using this method. As an example, up to 30% of a dose of vancomycin is removed if it is given during the last hour of dialysis.

It is important to note the duration of dialysis in relation to the estimate of \( t^{1/2} \). For example, if the \( t^{1/2} \) is reported as 24 hr, but the dialysis duration is only 4 hr, the half-life value is probably not accurate. Conversely, if the \( t^{1/2} \) is reported as 1 hr during a 4-hr dialysis period, the half-life value might be more reliable.

Two examples illustrate the use of pharmacokinetic data to calculate drug clearance during dialysis. Phenobarbital has a \( V_d \) of approximately 50 L, a total body clearance of 0.3 L/hr, and a conventional hemodialysis clearance of 4.2 L/hr. The half-life off dialysis is 115 hr and this will decrease to 8 hr during dialysis. Approximately 50% of the drug would be removed from the body during 8 hr of dialysis. As another example, digoxin has a \( V_d \) of about 300 L and total body clearance of 2.4 L/hr in an anephric patient. The hemodialysis clearance of digoxin is 1.2 L/hr. Therefore, in this patient, the half-lives of digoxin are 86 hr off dialysis and 58 hr on dialysis. Although this appears to be a substantial decrease in half-life, it means that the patient would have to be dialyzed continuously for 58 hr to remove one-half of the digoxin from the body. The fraction of drug lost during a routine hemodialysis period of 4 hr would be only 5%. Thus, a supplemental dose of digoxin after hemodialysis is not warranted.

USING THE TABLES

Tables 2–1 through 2–4 provide semiquantitative data on selected drugs. An additional authoritative source of information is Drug Prescribing in Renal Failure: Dosing Guidelines for Adults. Drugs are classified on the basis of the reported
range of drug removal by hemodialysis or derived by using pharmacokinetic parameters taken from the literature using the equations cited previously in the text. Drug removal is intentionally described in a semiquantitative fashion for a number of reasons. First, much of this information changes quite rapidly (eg, as new dialysis techniques are developed). Second, a given value for the amount of drug removed or the dialysis clearance determined in one study might differ from that found in another study because of differences in dialysate or blood flow during dialysis or the duration of the dialysis. The duration of dialysis has become shorter in the past 10 yr. Previously, most conventional hemodialysis runs were 4–6 hr, whereas typical dialysis procedures now lasts 2.5–3 hr. Third, the tables list comments for the clarification of certain points and selected references are provided for more specific information.

TABLE 2–1. READILY REMOVED BY DIALYSIS (50–100% WITHIN 1 HEMODIALYSIS SESSION)

<table>
<thead>
<tr>
<th>DRUG</th>
<th>TYPE OF DIALYZER</th>
<th>% REMOVED IN N HOURS</th>
<th>REFERENCES</th>
</tr>
</thead>
<tbody>
<tr>
<td>Acyclovir</td>
<td>CONV 1</td>
<td>51–60 in 4–5</td>
<td>32, 33</td>
</tr>
<tr>
<td>Amikacin</td>
<td>CONV 1</td>
<td>50 in 6</td>
<td>5</td>
</tr>
<tr>
<td>Amoxicillin</td>
<td>CONV 2</td>
<td>64 in 4</td>
<td>64</td>
</tr>
<tr>
<td>Cefadroxil</td>
<td>CONV 1</td>
<td>63 in 6–8</td>
<td>35</td>
</tr>
<tr>
<td>Cefazolin</td>
<td>SYN 1 and HE</td>
<td>40–60 in 3–4</td>
<td>36, 37</td>
</tr>
<tr>
<td>Cefmetazole</td>
<td>HE</td>
<td>52–67 in 3</td>
<td>38</td>
</tr>
<tr>
<td>Ceftazidime</td>
<td>CONV 2</td>
<td>40–50 in 4</td>
<td>22, 39, 40</td>
</tr>
<tr>
<td></td>
<td>HE</td>
<td>58 in 3</td>
<td></td>
</tr>
<tr>
<td></td>
<td>SYN2</td>
<td>60–75 in 3</td>
<td></td>
</tr>
<tr>
<td>Cefepime</td>
<td>HE</td>
<td>68 in 3</td>
<td>41, 42</td>
</tr>
<tr>
<td></td>
<td>SYN 1</td>
<td>72 in 3.5</td>
<td></td>
</tr>
<tr>
<td>Cefprozil</td>
<td>HE</td>
<td>55 in 3</td>
<td>43</td>
</tr>
<tr>
<td>Clavulanic acid</td>
<td>CONV 2</td>
<td>65 in 4</td>
<td>64</td>
</tr>
<tr>
<td>Flucytosine</td>
<td>CONV 1</td>
<td>69 in 4</td>
<td>44</td>
</tr>
<tr>
<td>Ganciclovir</td>
<td>CONV 2</td>
<td>60 in 4</td>
<td>45</td>
</tr>
<tr>
<td>Gentamicin</td>
<td>CONV 1</td>
<td>30–50 in 4</td>
<td>46–48</td>
</tr>
<tr>
<td></td>
<td>HE</td>
<td>40–45 in 3–4</td>
<td></td>
</tr>
<tr>
<td></td>
<td>SYN 1</td>
<td>50–60 in 3</td>
<td></td>
</tr>
<tr>
<td>Isepancin</td>
<td>HE</td>
<td>42–85 in 3</td>
<td>49</td>
</tr>
<tr>
<td>Imipenem/cilistatin</td>
<td>CONV 2</td>
<td>55–63 in 4</td>
<td>5</td>
</tr>
<tr>
<td>Lithium</td>
<td>CONV 2</td>
<td>61 in 6</td>
<td>50, 51</td>
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<tr>
<td>Methotrexate</td>
<td>SYN 1</td>
<td>60 in 4–6</td>
<td>52</td>
</tr>
<tr>
<td>Mezlocillin</td>
<td>CONV 1</td>
<td>62 in 6</td>
<td>53</td>
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</table>

(continued)
### TABLE 2–1. READILY REMOVED BY DIALYSIS (50–100% WITHIN 1 HEMODIALYSIS SESSION) (continued)

<table>
<thead>
<tr>
<th>DRUG</th>
<th>TYPE OF DIALYZER</th>
<th>% REMOVED IN N HOURS</th>
<th>REFERENCES</th>
</tr>
</thead>
<tbody>
<tr>
<td>Netilmicin</td>
<td>CONV 1</td>
<td>50 in 4</td>
<td>54, 55</td>
</tr>
<tr>
<td></td>
<td>SYN 3</td>
<td>56–62 in 4</td>
<td></td>
</tr>
<tr>
<td>Phenobarbital</td>
<td>CONV 1</td>
<td>36–45 in 6–8</td>
<td>56</td>
</tr>
<tr>
<td></td>
<td>SYN 1</td>
<td>51 in 4</td>
<td></td>
</tr>
<tr>
<td>Theophylline</td>
<td>CONV 1</td>
<td>59 in 4</td>
<td>57–61</td>
</tr>
<tr>
<td>Tobramycin</td>
<td>CONV 1</td>
<td>40–50 in 4</td>
<td>18, 45, 62</td>
</tr>
<tr>
<td></td>
<td>SYN 2</td>
<td>60–75 in 3</td>
<td></td>
</tr>
</tbody>
</table>

*CONV 1 = conventional with dialyzers that are no longer available; CONV 2 = conventional with currently available dialyzers; HE = high efficiency with cellulose acetate dialyzers CA170 and CA210; SYN 1 = synthetic; polysulfone; SYN 2 = synthetic; polysulfone, polymethylmethacrylate and cellulose triacetate; SYN 3 = synthetic; polyacrylonitrile.

### TABLE 2–2. MODERATELY DIALYZABLE (20–50% WITHIN 1 HEMODIALYSIS SESSION)

<table>
<thead>
<tr>
<th>DRUG</th>
<th>REFERENCES</th>
<th>DRUG</th>
<th>REFERENCES</th>
</tr>
</thead>
<tbody>
<tr>
<td>Acebutolol</td>
<td>65, 66</td>
<td>Cefradine</td>
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</tr>
<tr>
<td>Allopurinol</td>
<td>5</td>
<td>Cyclophosphamide</td>
<td>84, 85</td>
</tr>
<tr>
<td>Acetaminophen</td>
<td>5</td>
<td>Didanosine</td>
<td>86, 87</td>
</tr>
<tr>
<td>Ampicillin/sulbactam</td>
<td>67</td>
<td>Enalapril</td>
<td>88</td>
</tr>
<tr>
<td>Atenolol</td>
<td>68–70</td>
<td>Ethoxyzimide</td>
<td>89</td>
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<tr>
<td>Azathioprine</td>
<td>71</td>
<td>Fluconazole</td>
<td>90, 91</td>
</tr>
<tr>
<td>Aztreonam</td>
<td>72</td>
<td>Foscarnet</td>
<td>92, 93</td>
</tr>
<tr>
<td>Bretyllium</td>
<td>73</td>
<td>Lisinopril</td>
<td>88</td>
</tr>
<tr>
<td>Cefaclor</td>
<td>74</td>
<td>Lorazepam</td>
<td>94</td>
</tr>
<tr>
<td>Cefamandole</td>
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<td>Methylidopa</td>
<td>5</td>
</tr>
<tr>
<td>Cefotaxime</td>
<td>75</td>
<td>Metronidazole</td>
<td>95, 96</td>
</tr>
<tr>
<td>Cefoxitin</td>
<td>76</td>
<td>Minoxidil</td>
<td>97</td>
</tr>
<tr>
<td>Cefpodoxime</td>
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<td>Nadolol</td>
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<td>Ceftizoxime</td>
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<td>Ofloxacin</td>
<td>98–100</td>
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<td>Cefuroxime</td>
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<td>Penicillin G</td>
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<td>Cephalexin</td>
<td>83</td>
<td>Pentoxifyline</td>
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(continued)
### TABLE 2–2. MODERATELY DIALYZABLE (20–50% WITHIN 1 HEMODIALYSIS SESSION) (continued)

<table>
<thead>
<tr>
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<th>REFERENCES</th>
<th>DRUG</th>
<th>REFERENCES</th>
</tr>
</thead>
<tbody>
<tr>
<td>Piperacillin</td>
<td>103–105</td>
<td>Sulfamethoxazole</td>
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<tr>
<td>Primidone</td>
<td>106, 107</td>
<td>Ticarcillin</td>
<td>115</td>
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<tr>
<td>Procarbazine</td>
<td>108, 109</td>
<td>Tocainide</td>
<td>116</td>
</tr>
<tr>
<td>Pyrazinamide</td>
<td>110, 111</td>
<td>Trimethoprim</td>
<td>114</td>
</tr>
<tr>
<td>Sotalol</td>
<td>112, 113</td>
<td>Vancomycin</td>
<td>19–21</td>
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### TABLE 2–3. NOT SIGNIFICANTLY REMOVED BY HEMODIALYSIS

<table>
<thead>
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<th>DRUG</th>
<th>REFERENCES</th>
<th>DRUG</th>
<th>REFERENCES</th>
</tr>
</thead>
<tbody>
<tr>
<td>Amantidine</td>
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<td>Doxepin</td>
<td>131</td>
</tr>
<tr>
<td>Amiodarone</td>
<td>117</td>
<td>Doxycycline</td>
<td>5</td>
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<tr>
<td>Amphotericin B</td>
<td>5</td>
<td>Epoetin alfa</td>
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<tr>
<td>Astemizole</td>
<td>118</td>
<td>Erythromycin</td>
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</tr>
<tr>
<td>Bleomycin</td>
<td>119</td>
<td>Esmolol</td>
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<td>Captopril</td>
<td>120</td>
<td>Ethambutol</td>
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<td>Carbamazepine</td>
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<td>Ethchlorvynol</td>
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<td>Cefonicid</td>
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<td>Etodolac</td>
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<td>Cefotaxime</td>
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<td>Famotidine</td>
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<td>Cefoperazone</td>
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<td>Felodipine</td>
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<tr>
<td>Chloramphenicol</td>
<td>124, 125</td>
<td>Filgrastim</td>
<td>138</td>
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<tr>
<td>Chloroquine</td>
<td>126</td>
<td>Flecainide</td>
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<td>Cimetidine</td>
<td>127</td>
<td>Flurbiprofen</td>
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<td>Ciprofloxacin</td>
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<td>Glutethimide</td>
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<td>Glyburide</td>
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<td>Cyclosporine</td>
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<td>Itraconazole</td>
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<td>Disopyramide</td>
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<td>Ketoconazole</td>
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### TABLE 2–3. NOT SIGNIFICANTLY REMOVED BY HEMODIALYSIS (continued)

<table>
<thead>
<tr>
<th>DRUG</th>
<th>REFERENCES</th>
<th>DRUG</th>
<th>REFERENCES</th>
</tr>
</thead>
<tbody>
<tr>
<td>Lidocaine</td>
<td>5</td>
<td>Ranitidine</td>
<td>153</td>
</tr>
<tr>
<td>Lomefloxacin</td>
<td>147, 148</td>
<td>Rifampin</td>
<td>111</td>
</tr>
<tr>
<td>Meprobamate</td>
<td>5</td>
<td>Rimantadine</td>
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<tr>
<td>Methadone</td>
<td>5</td>
<td>Secobarbital</td>
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<td>Methylprednisolone</td>
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<td>Sulindac</td>
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<td>Metoprolol</td>
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<td>Tacrolimus</td>
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<td>Miconazole</td>
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<td>Tetracycline</td>
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<td>Naproxen</td>
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<td>Thiabendazole</td>
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<td>Timolol</td>
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<td>Oxacillin</td>
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<tr>
<td>Propoxyphene</td>
<td>5</td>
<td>Valproic acid</td>
<td>5</td>
</tr>
<tr>
<td>Propranolol</td>
<td>5</td>
<td>Verapamil</td>
<td>162, 163</td>
</tr>
<tr>
<td>Quinapril</td>
<td>152</td>
<td>Zidovudine</td>
<td>164</td>
</tr>
<tr>
<td>Quinidine</td>
<td>5</td>
<td></td>
<td></td>
</tr>
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</table>

### TABLE 2–4. NO DATA

<table>
<thead>
<tr>
<th>DRUG</th>
<th>DRUG</th>
</tr>
</thead>
<tbody>
<tr>
<td>Alprazolam</td>
<td>Clorazepate</td>
</tr>
<tr>
<td>Amitriptyline&lt;sup&gt;a&lt;/sup&gt;</td>
<td>Clozapine</td>
</tr>
<tr>
<td>Amlodipine</td>
<td>Codeine</td>
</tr>
<tr>
<td>Azithromycin&lt;sup&gt;a&lt;/sup&gt;</td>
<td>Colchicine</td>
</tr>
<tr>
<td>Baclofen</td>
<td>Cytarabine</td>
</tr>
<tr>
<td>Betaxolol</td>
<td>Dapsone</td>
</tr>
<tr>
<td>Bumetanide</td>
<td>Desipramine</td>
</tr>
<tr>
<td>Carbofelol</td>
<td>Diltiazem&lt;sup&gt;b&lt;/sup&gt;</td>
</tr>
<tr>
<td>Chlorpheniramine</td>
<td>Dipyridamole&lt;sup&gt;b&lt;/sup&gt;</td>
</tr>
<tr>
<td>Chlorpromazine</td>
<td>Dopamine</td>
</tr>
<tr>
<td>Chlorpropanide</td>
<td>Ethacrynic acid</td>
</tr>
<tr>
<td>Cisplatin</td>
<td>Felbamate</td>
</tr>
<tr>
<td>Clarithromycin</td>
<td>Fenoprofen&lt;sup&gt;b&lt;/sup&gt;</td>
</tr>
<tr>
<td>Clofazimine</td>
<td>Fluorouracil</td>
</tr>
</tbody>
</table>

(continued)
The use of multiple modes of continuous renal replacement therapy in critically ill individuals is increasing. The principles of drug removal for some of the continuous renal replacement therapies are different from those discussed in this section. Therefore, the reader is referred to selected references that provide an introduction to this topic and tabulations of literature data on drug removal.13,14,31,32

### TABLE 2-4. NO DATA (continued)

<table>
<thead>
<tr>
<th>DRUG</th>
<th>DRUG</th>
</tr>
</thead>
<tbody>
<tr>
<td>Flurazepam</td>
<td>Penicillamine</td>
</tr>
<tr>
<td>Glipizide</td>
<td>Pentamidine(^c)</td>
</tr>
<tr>
<td>Griseofulvin</td>
<td>Pentazocine</td>
</tr>
<tr>
<td>Guanethidine</td>
<td>Phenothiazines</td>
</tr>
<tr>
<td>Haloperidol(^b)</td>
<td>Phenytoin(^b)</td>
</tr>
<tr>
<td>Hydralazine</td>
<td>Piroxicam(^b)</td>
</tr>
<tr>
<td>Hydrochlorothiazide</td>
<td>Pravastatin</td>
</tr>
<tr>
<td>Indinavir(^b)</td>
<td>Prazepam</td>
</tr>
<tr>
<td>Indomethacin</td>
<td>Prazosin</td>
</tr>
<tr>
<td>Lamivudine</td>
<td>Prednisone</td>
</tr>
<tr>
<td>Levodopa</td>
<td>Primaquine</td>
</tr>
<tr>
<td>Lovastatin</td>
<td>Probencid</td>
</tr>
<tr>
<td>Melphalan</td>
<td>Ramipril</td>
</tr>
<tr>
<td>Meperidine</td>
<td>Rifabutin(^d)</td>
</tr>
<tr>
<td>Methaqualone</td>
<td>Ritonavir(^b)</td>
</tr>
<tr>
<td>Metholazone</td>
<td>Saquinavir(^b)</td>
</tr>
<tr>
<td>Midazolam(^b)</td>
<td>Sargramostin</td>
</tr>
<tr>
<td>Mirinone</td>
<td>Simvastatin</td>
</tr>
<tr>
<td>Minocycline</td>
<td>Spirocolactone</td>
</tr>
<tr>
<td>Misoprostol</td>
<td>Stavudine</td>
</tr>
<tr>
<td>Morphine</td>
<td>Sulindac</td>
</tr>
<tr>
<td>Muromonab-CD3</td>
<td>Sulindac</td>
</tr>
<tr>
<td>Nelfinavir(^b)</td>
<td>Tamoxifen</td>
</tr>
<tr>
<td>Nevirapine</td>
<td>Tolmetin(^b)</td>
</tr>
<tr>
<td>Nitrofurantoin</td>
<td>Triamterene</td>
</tr>
<tr>
<td>Norfloxacin</td>
<td>Vinblastine(^c)</td>
</tr>
<tr>
<td>Nortriptyline(^a)</td>
<td>Vincristine(^c)</td>
</tr>
<tr>
<td>Ondansetron</td>
<td>Warfarin(^d)</td>
</tr>
<tr>
<td>Paromomycin</td>
<td>Zalcitabine</td>
</tr>
</tbody>
</table>

\(^a\)Removal unlikely due to large \(V_d\) and high degree of protein binding.
\(^b\)Removal unlikely due to extensive protein binding.
\(^c\)Removal unlikely due to large \(V_d\).
REFERENCES


123. Reitberg DP et al. Pharmacokinetics of cefoperazone (2.0 g) and sulbactam (1.0 g) coadministered to subjects with normal renal function, patients with decreased renal function, and patients with end-stage renal disease on hemodialysis. Antimicrob Agents Chemother 1988;32:503–9.
GENERAL RECOMMENDATIONS ON IMMUNIZATION*

Recommendations for immunizing infants, children, and adults are based on characteristics of immunobiologics, principles of active and passive immunization, and judgments by public health officials and specialists in clinical and preventive medicine. Benefits and risks are associated with the use of all immunobiologics; no vaccine is completely safe or completely effective. Recommendations for immunization practices balance scientific evidence of benefits, costs, and risks to achieve optimal levels of protection against infectious diseases. The recommendations in this chapter describe this balance and attempt to minimize the risks by providing information regarding dose, route, and spacing of immunobiologics, and delineating situations that warrant precautions or contraindicate the use of these immunobiologics. These recommendations are for use only in the United States because vaccines and epidemiologic circumstances often differ in other countries. Individual circumstances may warrant deviations from these recommendations. Tables 3–1 and 3–2 list the immunobiologics available in the United States.

IMMUNOBIOLOGICS

The specific nature and content of immunobiologics can differ. When immunobiologics against the same infectious agent are produced by different manufacturers, active and inert ingredients in the various products are not always the same. Practitioners are urged to become familiar with the constituents of the various products.

Suspending Fluids. These may be sterile water, saline, or complex fluids containing protein or other constituents derived from the medium or biologic system in which the vaccine is produced (eg, serum proteins, egg antigens, and cell culture–derived antigens).

Preservatives, Stabilizers, Antibiotics. These components of vaccines, antitoxins, and globulins are used to inhibit or prevent bacterial growth in viral cultures.

*Excerpted from reference 1.
### Table 3–1. Licensed Vaccines and Toxoids Available in the United States, by Type and Recommended Routes of Administration

<table>
<thead>
<tr>
<th>Vaccine Type</th>
<th>Route</th>
</tr>
</thead>
<tbody>
<tr>
<td>Adenovirus&lt;sup&gt;a&lt;/sup&gt;</td>
<td>Live virus</td>
</tr>
<tr>
<td>Anthrax</td>
<td>Inactivated bacteria</td>
</tr>
<tr>
<td>Bacillus of Calmette and Guérin (BCG)</td>
<td>Live bacteria</td>
</tr>
<tr>
<td>Cholera</td>
<td>Inactivated bacteria</td>
</tr>
<tr>
<td>Diphtheria-tetanus-acellular pertussis (DTaP)</td>
<td>Toxoids and inactivated bacterial components</td>
</tr>
<tr>
<td>Diphtheria-tetanus-pertussis (DTP)</td>
<td>Toxoids and inactivated whole bacteria</td>
</tr>
<tr>
<td>DTP–Haemophilus influenzae type b conjugate (DTP-Hib)</td>
<td>Toxoids, inactivated whole bacteria, and bacterial polysaccharide conjugated to protein</td>
</tr>
<tr>
<td>Haemophilus influenzae type b conjugate (Hib)&lt;sup&gt;d&lt;/sup&gt;</td>
<td>Bacterial polysaccharide conjugated to protein</td>
</tr>
<tr>
<td>Hepatitis A</td>
<td>Inactivated virus</td>
</tr>
<tr>
<td>Hepatitis B</td>
<td>Inactive viral antigen</td>
</tr>
<tr>
<td>Influenza</td>
<td>Inactivated virus or viral components</td>
</tr>
<tr>
<td>Lyme disease</td>
<td>Bacterial lipoprotein</td>
</tr>
<tr>
<td>Japanese encephalitis</td>
<td>Inactivated virus</td>
</tr>
<tr>
<td>Measles</td>
<td>Live virus</td>
</tr>
<tr>
<td>Measles-mumps-rubella (MMR)</td>
<td>Live virus</td>
</tr>
<tr>
<td>Meningococcal</td>
<td>Bacterial polysaccharides of serotypes A/C/Y/W-135</td>
</tr>
<tr>
<td>Mumps</td>
<td>Live virus</td>
</tr>
<tr>
<td>Pertussis</td>
<td>Inactivated whole bacteria</td>
</tr>
<tr>
<td>Plague</td>
<td>Inactivated bacteria</td>
</tr>
<tr>
<td>Pneumococcal conjugate</td>
<td>Bacterial polysaccharides of 7 pneumococcal types</td>
</tr>
<tr>
<td>Pneumococcal polyvalent</td>
<td>Bacterial polysaccharides of 23 pneumococcal types or subcutaneous</td>
</tr>
<tr>
<td>Poliovirus vaccine, inactivated (IPV)</td>
<td>Inactivated viruses of all 3 serotypes</td>
</tr>
</tbody>
</table>

(continued)
TABLE 3–1. LICENSED VACCINES AND TOXOIDS AVAILABLE IN THE UNITED STATES, BY TYPE AND RECOMMENDED ROUTES OF ADMINISTRATION (continued)

<table>
<thead>
<tr>
<th>VACCINE TYPE</th>
<th>ROUTE</th>
</tr>
</thead>
<tbody>
<tr>
<td>Poliovirus vaccine, oral (OPV)</td>
<td>Live viruses of all 3 serotypes</td>
</tr>
<tr>
<td>Rabies</td>
<td>Inactivated virus</td>
</tr>
<tr>
<td>Rubella</td>
<td>Live virus</td>
</tr>
<tr>
<td>Tetanus</td>
<td>Inactivated toxin (toxoid)</td>
</tr>
<tr>
<td>Tetanus-diphtheria (Td or DT)</td>
<td>Inactivated toxins (toxoids)</td>
</tr>
<tr>
<td>Typhoid (parenteral)</td>
<td>Inactivated bacteria</td>
</tr>
<tr>
<td>(Ty21a oral)</td>
<td>Live bacteria</td>
</tr>
<tr>
<td>(Vi CPS)</td>
<td>Bacterial polysaccharide</td>
</tr>
<tr>
<td>Varicella</td>
<td>Live virus</td>
</tr>
<tr>
<td>Yellow fever</td>
<td>Live virus</td>
</tr>
</tbody>
</table>

*Modified from reference 1.
*Available only to the U.S. Armed Forces.
*The intradermal dose is lower than the subcutaneous dose.
*The recommended schedule for infants depends on the vaccine manufacturer; consult the package insert and ACIP recommendations for specific products.
*The intradermal dose of rabies vaccine, human diploid cell (HDCV), is lower than the intramuscular dose and is used only for pre-exposure vaccination. Rabies vaccine, absorbed (RVA) should not be used intradermally.
*Preparations with adjuvants should be administered intramuscularly.
*Td = tetanus and diphtheria toxoids for use among persons ≥7 years of age. Td contains the same amount of tetanus toxoid as DTP or DT, but contains a smaller dose of diphtheria toxoid. DT = tetanus and diphtheria toxoids for use among children <7 years of age.
*Booster doses may be administered intradermally unless vaccine that is acetone killed and dried is used.
<table>
<thead>
<tr>
<th>IMMUNOBIOLOGIC</th>
<th>TYPE</th>
<th>INDICATION(S)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Botulinum antitoxin</td>
<td>Specific equine antibodies</td>
<td>Treatment of botulism</td>
</tr>
<tr>
<td>Cytomegalovirus immune globulin (CMV-IGIV)</td>
<td>Specific human antibodies</td>
<td>Prophylaxis for bone marrow and kidney transplant recipients</td>
</tr>
<tr>
<td>Diphtheria antitoxin</td>
<td>Specific equine antibodies</td>
<td>Treatment of respiratory diphtheria</td>
</tr>
<tr>
<td>Hepatitis B immune (HBIG)</td>
<td>Specific human antibodies</td>
<td>Hepatitis B postexposure prophylaxis</td>
</tr>
<tr>
<td>Immune globulin (IG)</td>
<td>Pooled human antibodies</td>
<td>Hepatitis A pre- and post-exposure prophylaxis; measles postexposure prophylaxis</td>
</tr>
<tr>
<td>Immune globulin, intravenous (IGIV)</td>
<td>Pooled human antibodies</td>
<td>Replacement therapy for antibody deficiency disorders; immune thrombocytopenic purpura (ITP); hypogammaglobulinemia in chronic lymphocytic leukemia; Kawasaki disease</td>
</tr>
<tr>
<td>Rabies immune globulinb</td>
<td>Specific human antibodies</td>
<td>Rabies postexposure management of persons not previously immunized with rabies vaccine</td>
</tr>
<tr>
<td>Tetanus immune globulin (TIG)</td>
<td>Specific human antibodies</td>
<td>Tetanus treatment; postexposure prophylaxis of persons not adequately immunized with tetanus toxoid</td>
</tr>
<tr>
<td>Vaccinia immune globulin (VIG)</td>
<td>Specific human antibodies</td>
<td>Treatment of eczema vaccinatum, vaccinia necrosum, and ocular vaccinia</td>
</tr>
<tr>
<td>Varicella-zoster immune globulin (VZIG)</td>
<td>Specific human antibodies</td>
<td>Postexposure prophylaxis of susceptible immunocompromised persons, certain susceptible pregnant women, and perinatally exposed newborn infants</td>
</tr>
</tbody>
</table>

*a*Immune globulin preparations and antitoxins are administered intramuscularly unless otherwise indicated.

*b*HRIG is administered around the wounds in addition to the intramuscular injection.
or the final product or to stabilize the antigens or antibodies. Allergic reactions can occur if the recipient is sensitive to one of these additives (eg, phenols, albumin, glycine, and neomycin).

**Adjuvants.** Many antigens evoke suboptimal immunologic responses. Efforts to enhance immunogenicity include mixing antigens with a variety of substances or adjuvants (eg, aluminum adjuvants such as aluminum phosphate or aluminum hydroxide).

**Storage and Handling of Immunobiologics.** Failure to adhere to recommended specifications for storage and handling of immunobiologics can make these products impotent. Recommendations included in a product’s package insert should be followed closely to ensure maximum potency of vaccines. Vaccines should be stored at recommended temperatures immediately upon receipt. Certain vaccines, such as oral polio vaccine (OPV) and yellow fever vaccine, are very sensitive to increased temperature. Other vaccines are sensitive to freezing, including diphtheria and tetanus toxoids and pertussis vaccine, adsorbed (DTP); diphtheria and tetanus toxoids and acellular pertussis vaccine, adsorbed (DTaP); diphtheria and tetanus toxoids for pediatric use (DT); tetanus and diphtheria toxoids for adult use (Td); inactivated poliovirus vaccine (IPV); *Haemophilus influenzae* type b conjugate vaccine (Hib); hepatitis B vaccine; pneumococcal vaccines; and influenza vaccine. Mishandled vaccine may not be easily distinguished from potent vaccine, and, when in doubt, contact the manufacturer.

## ADMINISTRATION OF VACCINES

**General Instructions.** Persons administering vaccines should take precautions to minimize risk for spreading disease and be adequately immunized against hepatitis B, measles, mumps, rubella, and influenza. Tetanus and diphtheria toxoids are recommended for all persons. Hands should be washed between patients. Gloves are not required when administering vaccinations, unless the persons who administer the vaccine will come in contact with potentially infectious body fluids or have open lesions on their hands. Syringes and needles used for injections must be sterile and preferably disposable to minimize the risk of contamination. A separate needle and syringe should be used for each injection. Different vaccines should not be mixed in the same syringe unless specifically licensed for such use. Disposable needles and syringes should be discarded in labeled, puncture-proof containers to prevent inadvertent reuse or needlestick injury.

Routes of administration are recommended for each immunobiologic (see Table 3–1). Injectable immunobiologics should be administered where there is little likelihood of local, neural, vascular, or tissue injury. In general, vaccines containing adjuvants should be injected into the muscle mass; when administered subcutaneously or intradermally, they can cause local irritation, induration, skin discoloration, inflammation, and granuloma formation. Once the needle is inserted into the injection site, the syringe plunger should be pulled back. If blood appears in the needle hub, the needle should be withdrawn and a new site selected before the vaccine is expelled. The process should be repeated until no blood appears.
Subcutaneous Injections. SC injections are usually administered into the thigh of infants and into the deltoid area of older children and adults. A ⅜- to ⅜-inch, 23- to 25-gauge needle should be inserted into the tissues below the dermal layer of the skin.

Intramuscular Injections. The preferred sites for IM injections are the anterolateral aspect of the upper thigh and the deltoid muscle of the upper arm. The buttock should not be used routinely for active vaccination of infants, children, or adults because of the potential for injury to the sciatic nerve. In addition, injection into the buttock has been associated with decreased immunogenicity of hepatitis B and rabies vaccines in adults. If the buttock is used for passive immunization when large volumes are to be injected or multiple doses are necessary (eg, large doses of immune globulin), the central region should be avoided; only the upper, outer quadrant should be used, and the needle should be directed anteriorly (ie, not inferiorly or perpendicular to the skin) to minimize the possibility of involvement with the sciatic nerve. For all IM injections, the needle should be long enough to reach the muscle mass and prevent vaccine from seeping into subcutaneous tissues, but not so long as to endanger the underlying neurovascular structures or bone.

Infants (<12 Months of Age). Among most infants, the anterolateral aspect of the thigh provides the largest muscle mass and is therefore the recommended site. However, the deltoid can also be used with the thigh, for example, when multiple vaccines must be administered on the same visit. In most cases, a ⅜- to 1-inch, 22- to 25-gauge needle is sufficient to penetrate muscle in the thigh of a 4-month-old infant. The free hand should bunch the muscle, and the needle should be directed inferiorly along the axis of the leg at an angle appropriate to reach the muscle while avoiding nearby neurovascular structures and bone.

Toddlers and Older Children. The deltoid may be used if the muscle mass is adequate. The needle size can range from 22 to 25 gauge and from ⅜ to 1¼ inches, based on the size of the muscle. As with infants, the anterolateral thigh may be used, but the needle should be longer—generally ranging from ⅜ to 1½ inches.

Adults. The deltoid is recommended for routine intramuscular vaccination among adults, particularly for hepatitis B vaccine. The suggested needle size is 1 to 1½ inches and 20- to 25-gauge.

Intradermal Injections. Intradermal injections are generally administered on the volar surface of the forearm, except for human diploid cell rabies vaccine (HDCV), for which reactions are less severe when the vaccine is administered in the deltoid area. With the bevel facing upward, a ⅜ to ⅜ inch, 25- or 27-gauge needle can be inserted into the epidermis at an angle parallel to the long axis of the forearm. The needle should be inserted so that the entire bevel penetrates the skin and the injected solution raises a small bleb. Because of the small amounts of antigen used in intradermal injections, care must be taken not to inject the vaccine subcutaneously, because it can result in a suboptimal immunologic response.

Multiple Vaccinations. If more than one vaccine preparation is administered or if live vaccine and an immune globulin preparation are administered simultaneously, it is preferable to administer each at a different anatomic site. It is also preferable
to avoid administering two IM injections in the same limb, especially if DTP is one of the products administered. However, if more than one injection must be administered in a single limb, the thigh is usually the preferred site because of the greater muscle mass; the injections should be sufficiently separated (ie, 1–2 inches apart) so that any local reactions are unlikely to overlap.

**Regurgitated Oral Vaccines.** Infants may sometimes fail to swallow oral preparations (eg, OPV) after administration. If a substantial amount of the vaccine is spit out, regurgitated, or vomited shortly after administration (ie, within 5–10 min), another dose can be administered at the same visit. If this repeat dose is not retained, neither dose should be counted, and the vaccine should be readministered at the next visit.

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**AGE AT WHICH IMMUNOBIOLOGICS ARE ADMINISTERED**

Recommendations for the age at which vaccines are administered are influenced by several factors: age-specific risks of disease, age-specific risks of complications, ability of persons of a given age to respond to the vaccine(s), and potential interference with the immune response by passively transferred maternal antibody. In general, vaccines are recommended for the youngest age group at risk for developing disease whose members are known to develop an adequate antibody response (see Tables 3–3, 3–4, and 3–5).

**SPACING OF IMMUNOBIOLOGICS**

**Interval Between Multiple Doses of Same Antigen.** Some products require administration of more than one dose for development of an adequate antibody response. In addition, some products require periodic reinforcement or booster doses to maintain protection (see Tables 3–3, 3–4, and 3–5). Longer-than-recommended intervals between doses do not reduce final antibody concentrations. Therefore, an interruption in the immunization schedule does not require reinstatement of the entire series of an immunobiologic or the addition of extra doses. However, administering doses of a vaccine or toxoid at less-than-recommended minimum intervals may decrease the antibody response and therefore should be avoided. Doses administered at less-than-recommended minimum intervals should not be counted as part of a primary series.

Some immunobiologics produce increased rates of local or systemic reactions in certain recipients when administered too frequently (eg, adult Td, pediatric DT, tetanus toxoid, and rabies vaccine). Such reactions are thought to result from the formation of antigen-antibody complexes.

**Simultaneous Administration.** Many of the commonly used vaccines can safely and effectively be administered simultaneously (ie, on the same day, not at the same anatomic site). Simultaneous administration is important in certain situations, including imminent exposure to several infectious diseases, preparation for foreign travel, and uncertainty that the person will return for further doses of vaccine.
<table>
<thead>
<tr>
<th>AGE</th>
<th>Birth</th>
<th>1 Mo</th>
<th>2 Mo</th>
<th>4 Mo</th>
<th>6 Mo</th>
<th>12 Mo</th>
<th>15 Mo</th>
<th>18 Mo</th>
<th>24 Mo</th>
<th>4–6 Yr</th>
<th>11–12 Yr</th>
<th>14–18 Yr</th>
</tr>
</thead>
<tbody>
<tr>
<td>VACCINE</td>
<td></td>
<td></td>
<td></td>
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<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Hepatitis B&lt;sup&gt;b&lt;/sup&gt;</td>
<td></td>
<td>Hep B–1</td>
<td></td>
<td></td>
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</tr>
<tr>
<td>Diphtheria and tetanus toxoids and pertussis&lt;sup&gt;c&lt;/sup&gt;</td>
<td>DTaP</td>
<td>DTaP</td>
<td>DTaP</td>
<td>DTaP</td>
<td>DTaP</td>
<td>DTaP</td>
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<td></td>
<td></td>
<td></td>
<td></td>
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</tr>
<tr>
<td>Haemophilus influenzae type b&lt;sup&gt;d&lt;/sup&gt;</td>
<td>IPV</td>
<td>IPV</td>
<td>IPV</td>
<td>IPV</td>
<td>IPV</td>
<td>IPV</td>
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<td></td>
<td></td>
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<td></td>
</tr>
<tr>
<td>Inactivated polio&lt;sup&gt;e&lt;/sup&gt;</td>
<td>PCV</td>
<td>PCV</td>
<td>PCV</td>
<td>PCV</td>
<td>PCV</td>
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</tbody>
</table>

(continued)
**TABLE 3–3. RECOMMENDED CHILDHOOD IMMUNIZATION SCHEDULE United States, January–December 2001 (continued)**

<table>
<thead>
<tr>
<th>AGE</th>
<th>Vaccines</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>VACCINE</strong></td>
<td>Birth</td>
</tr>
<tr>
<td>Measles-mumps-rubella&lt;sup&gt;a&lt;/sup&gt;</td>
<td>MMR</td>
</tr>
<tr>
<td>Varicella&lt;sup&gt;b&lt;/sup&gt;</td>
<td>Var</td>
</tr>
<tr>
<td>Hepatitis A&lt;sup&gt;c&lt;/sup&gt;</td>
<td></td>
</tr>
</tbody>
</table>

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- **Range of recommended ages for vaccination.**
- **Vaccines to be given if previously recommended doses were missed or given earlier than the recommended minimum age.**
- **Recommended in selected states and/or regions.**

<sup>a</sup>From reference 2. This schedule lists the recommended ages for routine administration of currently licensed childhood vaccines as of November 1, 2000, for children up to age 18 yr. Additional vaccines might be licensed and recommended during the year. Licensed combination vaccines may be used whenever any components of the combination are indicated and the vaccine’s other components are not contraindicated. Providers should consult the manufacturer’s package inserts for detailed recommendations.
TABLE 3–3. RECOMMENDED CHILDHOOD IMMUNIZATION SCHEDULE* UNITED STATES, JANUARY–DECEMBER 2001 (continued)

Infants born to hepatitis B surface antigen (HBsAg)–negative mothers should receive the first dose of hepatitis B vaccine (Hep B) by age 2 months. The second dose should be administered at least 1 month after the first dose. The third dose should be administered at least 4 months after the first dose and at least 2 months after the second dose, but not before age 6 months. Infants born to HBsAg-positive mothers should receive Hep B and 0.5 mL hepatitis B immune globulin (HBIG) within 12 hr of birth at separate sites. The second dose is recommended at age 1–2 months and the third dose at age 6 months. Infants born to mothers whose HBsAG status is unknown should receive Hep B within 12 hr of birth. Maternal blood should be drawn at delivery to determine the mother’s HBsAG status; if the HBsAG test is positive, the infant should receive HBIG as soon as possible (no later than age 1 week). All children and adolescents (through age 18 yr) who have not been immunized against hepatitis B should begin the series during any visit. Providers should make special efforts to immunize children who were born in or whose parents were born in areas of the world where hepatitis B virus infection is moderately or highly endemic.

The fourth dose of diphtheria and tetanus toxoids and acellular pertussis vaccine (DTaP) can be administered as early as age 12 months provided 6 months have elapsed since the third dose and the child is unlikely to return at age 15–18 months. Tetanus and diphtheria toxoids (Td) is recommended at age 11–12 yr if at least 5 yr have elapsed since the last dose of Td and pertussis vaccine (DTP), DTaP, or diphtheria and tetanus (DT) toxoids. Subsequent routine Td boosters are recommended every 10 yr.

Three Haemophilus influenzae type b (Hib) conjugate vaccines are licensed for use in infants. If Hib conjugate vaccine (PRP-OMP, Pedvax HIB or ComVax, Merck) is administered at ages 2 and 4 months, a dose at age 6 months is not required. Because clinical studies in infants have demonstrated that using some combination products can induce a lower immune response to the Hib vaccine component, DTaP/Hib combination products should not be used for primary immunization in infants at age 2, 4, or 6 months unless approved by the Food and Drug Administration for these ages.

An all-inactivated polio virus vaccine (IPV) schedule is recommended for routine childhood polio vaccination in the United States. All children should receive four doses of IPV at age 2 months, age 4 months, between 6–18 months, and 4–6 yr. Oral polio virus vaccine should be used only in selected circumstances.3

The heptavalent pneumococcal conjugate vaccine (PCV) is recommended for all children 2–23 months old. It is also recommended for certain children 24–59 months old.4

The second dose of measles–mumps–rubella (MMR) vaccine is recommended routinely at age 4–6 yr but can be administered during any visit provided at least 4 weeks have elapsed since receipt of the first dose and both doses are administered beginning at or after age 12 months. Those who previously did not receive the second dose should complete the schedule no later than the routine visit to a health care provider at age 11–12 yr.

Varicella vaccine (Var) is recommended at any visit on or after the first birthday for susceptible children (ie, those who lack a reliable history of chickenpox as judged by a health care provider and have not been immunized). Susceptible persons age ≥13 yr should receive two doses given at least 4 weeks apart.

Hepatitis A vaccine (Hep A) is recommended for use in selected states and/or regions and for certain high-risk groups. Information is available from local public health authorities.5

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* Infants born to hepatitis B surface antigen (HBsAg)–negative mothers should receive the first dose of hepatitis B vaccine (Hep B) by age 2 months. The second dose should be administered at least 1 month after the first dose. The third dose should be administered at least 4 months after the first dose and at least 2 months after the second dose, but not before age 6 months. Infants born to HBsAg-positive mothers should receive Hep B and 0.5 mL hepatitis B immune globulin (HBIG) within 12 hr of birth at separate sites. The second dose is recommended at age 1–2 months and the third dose at age 6 months. Infants born to mothers whose HBsAG status is unknown should receive Hep B within 12 hr of birth. Maternal blood should be drawn at delivery to determine the mother’s HBsAG status; if the HBsAG test is positive, the infant should receive HBIG as soon as possible (no later than age 1 week). All children and adolescents (through age 18 yr) who have not been immunized against hepatitis B should begin the series during any visit. Providers should make special efforts to immunize children who were born in or whose parents were born in areas of the world where hepatitis B virus infection is moderately or highly endemic.

The fourth dose of diphtheria and tetanus toxoids and acellular pertussis vaccine (DTaP) can be administered as early as age 12 months provided 6 months have elapsed since the third dose and the child is unlikely to return at age 15–18 months. Tetanus and diphtheria toxoids (Td) is recommended at age 11–12 yr if at least 5 yr have elapsed since the last dose of Td and pertussis vaccine (DTP), DTaP, or diphtheria and tetanus (DT) toxoids. Subsequent routine Td boosters are recommended every 10 yr.

Three Haemophilus influenzae type b (Hib) conjugate vaccines are licensed for use in infants. If Hib conjugate vaccine (PRP-OMP, Pedvax HIB or ComVax, Merck) is administered at ages 2 and 4 months, a dose at age 6 months is not required. Because clinical studies in infants have demonstrated that using some combination products can induce a lower immune response to the Hib vaccine component, DTaP/Hib combination products should not be used for primary immunization in infants at age 2, 4, or 6 months unless approved by the Food and Drug Administration for these ages.

An all-inactivated polio virus vaccine (IPV) schedule is recommended for routine childhood polio vaccination in the United States. All children should receive four doses of IPV at age 2 months, age 4 months, between 6–18 months, and 4–6 yr. Oral polio virus vaccine should be used only in selected circumstances.3

The heptavalent pneumococcal conjugate vaccine (PCV) is recommended for all children 2–23 months old. It is also recommended for certain children 24–59 months old.4

The second dose of measles–mumps–rubella (MMR) vaccine is recommended routinely at age 4–6 yr but can be administered during any visit provided at least 4 weeks have elapsed since receipt of the first dose and both doses are administered beginning at or after age 12 months. Those who previously did not receive the second dose should complete the schedule no later than the routine visit to a health care provider at age 11–12 yr.

Varicella vaccine (Var) is recommended at any visit on or after the first birthday for susceptible children (ie, those who lack a reliable history of chickenpox as judged by a health care provider and have not been immunized). Susceptible persons age ≥13 yr should receive two doses given at least 4 weeks apart.

Hepatitis A vaccine (Hep A) is recommended for use in selected states and/or regions and for certain high-risk groups. Information is available from local public health authorities.5
TABLE 3–4. RECOMMENDED ACCELERATED IMMUNIZATION SCHEDULE FOR INFANTS AND CHILDREN <7 YEARS OF AGE WHO START THE SERIES LATE OR WHO ARE >1 MONTH BEHIND IN THE IMMUNIZATION SCHEDULE (ie, children for whom compliance with scheduled return visits cannot be assured)

<table>
<thead>
<tr>
<th>TIMING</th>
<th>VACCINE(S)</th>
<th>COMMENTS</th>
</tr>
</thead>
<tbody>
<tr>
<td>First visit (≥4 mo of age)</td>
<td>DTP, IPV or OPV, Hib, Hepatitis B, MMR (should be given as soon as child is age 12–15 mo)</td>
<td>All vaccines should be administered simultaneously at the appropriate visit.</td>
</tr>
<tr>
<td>Second visit (1 mo after first visit)</td>
<td>DTP, Hib, Hepatitis B</td>
<td></td>
</tr>
<tr>
<td>Third visit (1 mo after second visit)</td>
<td>DTP, OPV, Hib, Hib, Hib</td>
<td></td>
</tr>
<tr>
<td>Fourth visit (6 weeks after third visit)</td>
<td>OPV</td>
<td></td>
</tr>
<tr>
<td>Fifth visit (≥6 mo after third visit)</td>
<td>DTaP or DTP, Hib, Hib</td>
<td>Preferably at or before school entry.</td>
</tr>
<tr>
<td>Additional visits (Age 4–6 yr)</td>
<td>DTaP or DTP, OPV, MMR</td>
<td></td>
</tr>
<tr>
<td>(Age 14–16 yr)</td>
<td>Td</td>
<td>Repeat every 10 yr throughout life.</td>
</tr>
</tbody>
</table>

DTP, diphtheria-tetanus-pertussis; DTaP, diphtheria-tetanus-acellular pertussis; Hib, Haemophilus influenzae type b conjugate; MMR, measles-mumps-rubella; OPV, poliovirus vaccine, live oral, trivalent; Td, tetanus and diphtheria toxoids (for use among persons ≥7 years of age).

If initiated in the first year of life, administer DTP doses 1, 2, and 3 and OPV doses 1, 2, and 3 according to this schedule; administer MMR when the child reaches 12–15 mo of age.

See individual ACIP recommendations for detailed information on specific vaccines.

Two DTP and Hib combination vaccines are available (DTP/HbOC [TETRAMUNE], and PRP-T [ActHIB, OmniHIB] which can be reconstituted with DTP vaccine produced by Connaught). DTaP preparations are currently recommended only for use as the fourth and/or fifth doses of the DTP series among children 15 mo–6 yr of age (before the seventh birthday). DTP and DTaP should not be used on or after the seventh birthday.

The Advisory Committee on Immunization Practices (ACIP) of the Centers for Disease Control and Prevention (CDC) recommends the use of enhanced inactivated poliomyelitis vaccine (IPV) injection for the first 2 doses of the series to minimize OPV-related paralysis.

The recommended schedule varies by vaccine manufacturer. For information specific to the vaccine being used, consult the package insert and ACIP recommendations. Children beginning the Hib vaccine series at age 2–6 mo should receive a primary series of three doses of HbOc, PRP-T, or a licensed DTP-Hib combination vaccine; or two doses of PRP-OMP. An additional booster dose of any licensed Hib conjugate vaccine should be administered at 12–15 mo of age and at least 2 mo after

(continued)
the previous dose. Children beginning the Hib vaccine series at 7–11 mo of age should receive a primary series of two doses of a vaccine containing HbOC, PRP-T, or PRP-OMP. An additional booster dose of any licensed Hib conjugate vaccine should be administered at 12–18 mo of age and at least 2 mo after the previous dose. Children beginning the Hib vaccine series at ages 12–14 mo should receive a primary series of one dose of a vaccine containing HbOC, PRP-T, or PRP-OMP. An additional booster dose of any licensed Hib conjugate vaccine should be administered 2 mo after the previous dose. Children beginning the Hib vaccine series at ages 15–59 mo should receive one dose of any licensed Hib vaccine. Hib vaccine should not be administered after the fifth birthday except for special circumstances as noted in the specific ACIP recommendations for the use of Hib vaccine.
TABLE 3–5. RECOMMENDED IMMUNIZATION SCHEDULE FOR PERSONS ≥7 YR OF AGE NOT VACCINATED AT THE RECOMMENDED TIME IN EARLY INFANCY

<table>
<thead>
<tr>
<th>TIMING</th>
<th>VACCINE(S)</th>
<th>COMMENTS</th>
</tr>
</thead>
<tbody>
<tr>
<td>First visit</td>
<td>Td, OPV, MMR, and Hepatitis B</td>
<td>Primary poliovirus vaccination is not routinely recommended for persons ≥18 yr of age.</td>
</tr>
<tr>
<td>Second visit (6–8 weeks after first visit)</td>
<td>Td, OPV, MMR, and Hepatitis B</td>
<td></td>
</tr>
<tr>
<td>Third visit (6 mo after second visit)</td>
<td>Td, OPV, Hepatitis B</td>
<td></td>
</tr>
<tr>
<td>Additional visits</td>
<td>Td</td>
<td>Repeat every 10 yr throughout life.</td>
</tr>
</tbody>
</table>

MMR, measles-mumps-rubella; OPV, poliovirus vaccine, live oral, trivalent; Td, tetanus and diphtheria toxoids (for use among persons ≥7 yr of age).

aSee individual ACIP recommendations for details.

bThe DTP and DTaP doses administered to children <7 yr of age who remain incompletely vaccinated at age ≥7 yr should be counted as prior exposure to tetanus and diphtheria toxoids (eg, a child who previously received two doses of DTP needs only one dose of Td to complete a primary series for tetanus and diphtheria).

cWhen polio vaccine is administered to previously unvaccinated persons ≥18 yr of age, inactivated poliovirus vaccine (IPV) is preferred. For the immunization schedule for IPV, see specific ACIP statement on the use of polio vaccine.

dPersons born before 1957 can generally be considered immune to measles and mumps and need not be vaccinated. Rubella (or MMR) vaccine can be administered to persons of any age, particularly to nonpregnant women of childbearing age.

eHepatitis B vaccine, recombinant. Selected high-risk groups for whom vaccination is recommended include persons with occupational risk, such as health care and public safety workers who have occupational exposure to blood, clients and staff of institutions for the developmentally disabled, hemodialysis patients, recipients of certain blood products (eg, clotting factor concentrates), household contacts and sex partners of hepatitis B virus carriers, injecting drug users, sexually active homosexual and bisexual men, certain sexually active heterosexual men and women, inmates of long-term correctional facilities, certain international travelers, and families of HBsAg-positive adoptees from countries where HBV infection is endemic. Because risk factors are often not identified directly among adolescents, universal hepatitis B vaccination of teenagers should be implemented in communities where injecting drug use, pregnancy among teenagers, and/or sexually transmitted diseases are common.

fThe ACIP recommends a second dose of measles-containing vaccine (preferably MMR to ensure immunity to mumps and rubella) for certain groups. Children with no documentation of live measles vaccination after the first birthday should receive two doses of live measles-containing vaccine not less than 1 mo apart. In addition, the following persons born in 1957 or later should have documentation of measles immunity (ie, 2 doses of measles-containing vaccine [at least one of which being MMR], physician-diagnosed measles, or laboratory evidence of measles immunity): (a) those entering post-high school educational settings; (b) those beginning employment in health care settings who will have direct patient contact; and (c) travelers to areas with endemic measles.
Killed Vaccines. In general, inactivated vaccines can be administered simultaneously at separate sites. However, when vaccines commonly associated with local or systemic side effects (eg, cholera, parenteral typhoid, and plague) are administered simultaneously, the side effects might be accentuated. When feasible, it is preferable to administer these vaccines on separate occasions.

Live Vaccines. The simultaneous administration of the most widely used live and inactivated vaccines has not resulted in impaired antibody responses or increased rates of adverse reactions. Administration of combined measles, mumps, and rubella (MMR) vaccine yields results similar to administration of the individual vaccines at different sites. Concern has been raised that oral live attenuated typhoid (Ty21a) vaccine theoretically might interfere with the immune response to OPV when OPV is administered simultaneously or soon after live oral typhoid vaccine, but no published data exist to support this theory.

Routine Childhood Vaccines. The simultaneous administration of routine childhood vaccines does not interfere with the immune response to these vaccines. When administered at the same time and at separate sites, DTP, OPV, and MMR have produced seroconversion rates and rates of side effects similar to those observed when the vaccines are administered separately. Simultaneous vaccination of infants with DTP, OPV (or IPV), and either Hib vaccine or hepatitis B vaccine has resulted in acceptable response to all antigens. Routine simultaneous administration of DTP (or DTaP), OPV (or IPV), Hib vaccine, MMR, and hepatitis B vaccine is encouraged for children who are the recommended age to receive these vaccines and for whom no specific contraindications exist at the time of the visit. Individual vaccines should not be mixed in the same syringe unless they are licensed for mixing by the U.S. Food and Drug Administration (FDA).

Other Vaccines. The simultaneous administration of pneumococcal polysaccharide vaccine and whole-virus influenza vaccine elicits satisfactory antibody responses without increasing the frequency or severity of adverse reactions in adults. Simultaneous administration of the pneumococcal vaccine and split-virus influenza vaccine also yields satisfactory results in both children and adults.

Hepatitis B vaccine administered with yellow fever vaccine is as safe and efficacious as when these vaccines are administered separately. Measles and yellow fever vaccines have been administered together safely and with full efficacy.

The antibody response to yellow fever and cholera vaccines is decreased if administered simultaneously or within a short time of each other. If possible, separate yellow fever and cholera vaccinations by at least 3 weeks. If time constraints exist and both vaccines are necessary, the injections can be administered simultaneously or within a 3-week period with the understanding that antibody response may not be optimal. Yellow fever vaccine is required by many countries and is highly effective in protecting against a disease with substantial mortality and for which no therapy exists. The currently used cholera vaccine provides limited protection of brief duration; few indications exist for its use.

Antimalarials and Vaccination. The antimalarial mefloquine (Lariam) could potentially affect the immune response to oral live attenuated typhoid (Ty21a) vac-
cine if both are taken simultaneously. To minimize this effect, it may be prudent to administer Ty21a typhoid vaccine at least 24 hours before or after a dose of mefloquine. Because chloroquine phosphate (and possibly other structurally related antimalarials, such as mefloquine) may interfere with the antibody response to human diploid cell rabies vaccine (HDCV) when HDCV is administered by the intradermal route, HDCV should be administered by the intramuscular route when chloroquine, mefloquine, or other structurally related antimalarials are used.

**Nonsimultaneous Administration.** Inactivated vaccines generally do not interfere with the immune response to other inactivated vaccines or to live vaccines except in certain instances (eg, yellow fever and cholera vaccines). In general, an inactivated vaccine can be administered either simultaneously or at any time before or after a different inactivated vaccine or a live vaccine. However, limited data indicate that prior or concurrent administration of DTP vaccine may enhance anti-PRP antibody response following vaccination with certain *Haemophilus influenzae* type b conjugate vaccines (ie, PRP-T, PRP-D, and HbOC). For infants, the immunogenicity of PRP-OMP appears to be unaffected by the absence of prior or concurrent DTP vaccination.

Theoretically, the immune response to one live-virus vaccine might be impaired if administered within 30 days of another live-virus vaccine. Whenever possible, live-virus vaccines administered on different days should be administered at least 30 days apart. However, OPV and MMR vaccines can be administered at any time before, with, or after each other, if indicated. Live-virus vaccines can interfere with the response to a tuberculin test. Tuberculin testing, if otherwise indicated, can be done either on the same day the live-virus vaccines are administered or 4–6 weeks later.

**IMMUNE GLOBULIN**

**Live Vaccines.** OPV and yellow fever vaccines can be administered at any time before, with, or after the administration of immune globulin or specific immune globulins (eg, hepatitis B immune globulin [HBIG], rabies immune globulin [RIG]). The concurrent administration of immune globulin should not interfere with the response to Ty21a typhoid vaccine. Recent evidence suggests that high doses of immune globulin can inhibit the immune response to measles vaccine for more than 3 months. Administration of immune globulin can also inhibit the response to rubella vaccine. The effect of immune globulin preparations on the response to mumps and varicella vaccines is unknown, but commercial immune globulin preparations contain antibodies to these viruses.

Blood (eg, whole blood, packed RBCs, and plasma) and other antibody-containing blood products (eg, immune globulin; specific immune globulins; and immune globulin, intravenous [IGIV]) can diminish the immune response to MMR or its individual component vaccines. Therefore, after an immune globulin preparation is received, these vaccines should not be administered before the recommended interval has passed. However, postpartum vaccination of rubella-susceptible women with rubella or MMR vaccine should not be delayed because anti-Rho(D) IG (human) or any other blood product was received during the last
trimester of pregnancy or at delivery. These women should be vaccinated immediately after delivery and, if possible, tested at least 3 months later to ensure immunity to rubella and, if necessary, to measles.

If administration of an immune globulin preparation becomes necessary because of imminent exposure to disease, MMR or its component vaccines can be administered simultaneously with the immunoglobulin preparation, although vaccine-induced immunity might be compromised. The vaccine should be administered at a site remote from that chosen for the immune globulin inoculation. Unless serologic testing indicates that specific antibodies have been produced, vaccination should be repeated after the recommended interval.

If administration of an immune globulin preparation becomes necessary after MMR or its individual component vaccines have been administered, interference can occur. Usually vaccine virus replication and stimulation of immunity occurs 1–2 weeks after vaccination. Thus, if the interval between administration of any of these vaccines and subsequent administration of an immune globulin preparation is less than 14 days, vaccination should be repeated after the recommended interval unless serologic testing indicates that antibodies were produced.

**Killed Vaccines.** Immune globulin preparations interact less with inactivated vaccines and toxoids than with live vaccines. Therefore, administration of inactivated vaccines simultaneously with or at any interval before or after receipt of immune globulins should not substantially impair the development of a protective antibody response. The vaccine or toxoid and immune globulin preparation should be administered at different sites.

**Interchangeability of Vaccines From Different Manufacturers.** When at least one dose of a hepatitis B vaccine produced by one manufacturer is followed by subsequent doses from a different manufacturer, the immune response has been shown to be comparable with that resulting from a full course of vaccination with a single vaccine.

Both HDCV and rabies vaccine, adsorbed (RVA) are considered equally efficacious and safe. When used as licensed and recommended, they are considered interchangeable during the vaccine series. **RVA should not be used intradermally.** The full 1 mL dose of either product, administered by IM injection, can be used for both pre-exposure and postexposure prophylaxis.

When administered according to their licensed indications, different diphtheria and tetanus toxoids and pertussis vaccines as single antigens or various combinations, as well as the live and inactivated polio vaccines, also can be used interchangeably.

Currently licensed Haemophilus influenzae type b conjugate vaccines (ie, PRP-OMP, PRP-T, HbOC, and combination DTP-Hib vaccines) have been shown to induce different temporal patterns of immunologic response in infants. Data suggest that infants who receive sequential doses of different vaccines produce a satisfactory antibody response after a complete series. The primary vaccine series should be completed with the same Hib vaccine, if feasible. However, if different vaccines are administered, a total of 3 doses of Hib vaccine is considered adequate for the primary series among infants, and any combination of Hib conjugate vaccines licensed for use among infants may be used. Any of the licensed
conjugate vaccines can be used for the recommended booster dose at 12–18 months of age.

**HYPERSENSITIVITY TO VACCINE COMPONENTS**

Vaccine components can cause allergic reactions in some recipients. These reactions can be local or systemic, and can include mild to severe anaphylaxis or anaphylactoid responses (e.g., generalized urticaria or hives, wheezing, swelling of the mouth and throat, difficulty breathing, hypotension, and shock). The responsible vaccine components can derive from vaccine antigen, animal protein, antibiotics, preservatives, and stabilizers.

**Egg Allergy.** The most common animal protein allergen is egg protein found in vaccines prepared using embryonated chicken eggs (e.g., influenza and yellow fever vaccines) or chicken embryo cell cultures (e.g., measles and mumps vaccines). Ordinarily, persons who are able to eat eggs or egg products safely can receive these vaccines; persons with histories of anaphylactic or anaphylactoid allergy to eggs or egg proteins should not. Asking persons whether they can eat eggs without adverse effects is a reasonable way to determine who might be at risk for allergic reactions. Protocols for testing and vaccinating those persons with anaphylactic reactions to egg ingestion or vaccinating children with egg hypersensitivity and severe asthma have been developed. Rubella vaccine is grown in human diploid cell cultures and can be safely administered to persons with histories of severe allergy to eggs or egg proteins.

**Antibiotic Allergy.** Some vaccines contain trace amounts of antibiotics to which patients may be hypersensitive. The information provided in the vaccine package insert should be carefully reviewed before deciding if the uncommon patient with such hypersensitivity should receive the vaccine(s). No currently recommended vaccine contains penicillin or penicillin derivatives. MMR and its individual component vaccines contain trace amounts of neomycin and, although the amount present is less than would usually be used for a skin test to determine hypersensitivity, persons who have experienced anaphylactic reactions to neomycin should not receive these vaccines. Most often, neomycin allergy is a contact dermatitis—a manifestation of a delayed-type (cell-mediated) immune response—rather than anaphylaxis. A history of delayed-type reactions to neomycin is not a contraindication for these vaccines.

**Thimerosal Allergy.** Exposure to vaccines containing the preservative thimerosal (e.g., DTP, DTaP, DT, Td, Hib, hepatitis B, influenza, and Japanese encephalitis) can lead to induction of hypersensitivity. However, most patients do not develop reactions to thimerosal given as a component of vaccines even when patch or intradermal tests for thimerosal indicate hypersensitivity, which usually consists of local delayed-type hypersensitivity reactions. Manufacturers are removing thimerosal from many products.

**Vaccine Allergy.** Certain parenteral bacterial vaccines (i.e., cholera, DTP, plague, and typhoid) are frequently associated with local or systemic adverse effects,
such as redness, soreness, and fever. These reactions are difficult to link with a specific sensitivity to vaccine components and appear to be toxic rather than hypersensitive. Urticarial or anaphylactic reactions in DTP, DT, or Td or tetanus toxoid recipients have been reported rarely. When these reactions are reported, appropriate skin tests should be performed to determine sensitivity to tetanus toxoid before its use is discontinued. Alternatively, serologic testing to determine immunity to tetanus can be performed to evaluate the need for a booster dose of tetanus toxoid.

Vaccination in Special Populations

Preterm Infants. Infants born prematurely, regardless of birth weight, should be vaccinated at the same chronologic age and according to the same schedule and precautions as full-term infants and children. Birthweight and size generally are not factors in deciding whether to postpone routine vaccination of a clinically stable premature infant. The full recommended dose of each vaccine should be used. To prevent the theoretical risk of poliovirus transmission in the hospital, the administration of OPV should be deferred until discharge.

Any premature infant born to a hepatitis B surface antigen (HBsAg)-positive mother should receive immunoprophylaxis with hepatitis B vaccine and HBIG beginning at or shortly after birth. For premature infants of HBsAg-negative mothers, the optimal timing of hepatitis B vaccination has not been determined. Some studies suggest that decreased conversion rates might occur in some premature infants with low birthweights (ie, <2000 g) following administration of hepatitis B vaccine at birth. Such low-birthweight premature infants of HBsAg-negative mothers should receive the hepatitis B vaccine series, which can be initiated at discharge from the nursery if the infant weighs at least 2000 g or at 2 months of age along with DTP, OPV, and Hib vaccine.

Breastfeeding and Vaccination. Neither killed nor live vaccines affect the safety of breastfeeding for mothers of infants. Breastfeeding does not adversely affect immunization and is not a contraindication for any vaccine. Breast-fed infants should be vaccinated according to routine recommended schedules.

Inactivated or killed vaccines do not multiply within the body. Therefore, they should pose no special risk for mothers who are breastfeeding or for their infants. Although live vaccines do multiply within the mother’s body, most are not excreted in breastmilk. Although rubella vaccine virus may be transmitted in breastmilk, the virus usually does not infect the infant, and, if it does, the infection is well tolerated. There is no contraindication for vaccinating breastfeeding mothers with yellow fever vaccine. Breastfeeding mothers can receive OPV without any interruption in feeding schedule.

Vaccination During Pregnancy. Risk from vaccination during pregnancy is largely theoretical. The benefit of vaccination among pregnant women usually outweighs the potential risk when the risk for disease is high, infection would pose a special risk to the mother or fetus, and the vaccine is unlikely to cause harm.

Combined tetanus and diphtheria toxoids are the only immunobiologic agents routinely indicated for susceptible pregnant women. Previously vaccinated
pregnant women who have not received a Td vaccination within the past 10 years should receive a booster dose. Pregnant women who are unimmunized or only partially immunized against tetanus should complete the primary series. Depending on when a woman seeks prenatal care and the required interval between doses, one or two doses of Td can be administered before delivery. Women for whom the vaccine is indicated but who have not completed the required three-dose series during pregnancy should be followed up after delivery to ensure they receive the doses necessary for protection.

There is no convincing evidence of risk from immunizing pregnant women with other inactivated virus or bacteria vaccines or toxoids. Hepatitis B vaccine is recommended for women at risk for hepatitis B infection, and influenza and pneumococcal vaccines are recommended for women at risk for infection and for complications of influenza and pneumococcal disease.

OPV can be administered to pregnant women who are at substantial risk of exposure to natural infection. Although OPV is preferred, IPV may be considered if the complete vaccination series can be administered before the anticipated exposure. Pregnant women who must travel to areas where the risk of yellow fever is high should receive yellow fever vaccine. The small theoretical risk from vaccination is far outweighed by the risk of yellow fever infection. Known pregnancy is a contraindication for rubella, measles, and mumps vaccines. Although a theoretical concern, no cases of congenital rubella syndrome or abnormalities attributable to rubella vaccine virus infection have been observed in infants born to susceptible mothers who received rubella vaccine during pregnancy.

Persons who receive measles, mumps, or rubella vaccines can shed these viruses, but generally do not transmit them. These vaccines can be administered safely to the children of pregnant women. Although live poliovirus is shed by persons recently immunized with OPV (particularly after the first dose), this vaccine can also be administered to the children of pregnant women because experience has not revealed any risk of polio vaccine virus to the fetus.

All pregnant women should be evaluated for immunity to rubella and tested for the presence of HBsAg. Women susceptible to rubella should be immunized immediately after delivery. A woman infected with hepatitis B virus should be followed carefully to ensure that the infant receives HBIG and begins the hepatitis B vaccine series shortly after delivery.

There is no known risk to the fetus from passive immunization of pregnant women with immune globulin. Further information regarding immunization of pregnant women is available in the American College of Obstetricians and Gynecologists Technical Bulletin Number 160, October 1991.7

Altered Immunocompetence. This section is a summary of the more extensive recommendations on vaccines and immune globulin preparations for immunocompromised persons. Additional information can be found in references 8 and 9.

Severe immunosuppression can be the result of congenital immunodeficiency. HIV infection, leukemia, lymphoma, generalized malignancy, or therapy with alkylating agents, antimetabolites, radiation, or large amounts of corticosteroids. Severe complications have followed vaccination of immunocompromised patients with live, attenuated-virus vaccines and with live-bacteria
vaccines. In general, these patients should not receive live vaccines except in certain circumstances that are noted below. In addition, OPV should not be administered to any household contact of a severely immunocompromised person. If polio immunization is indicated for immunosuppressed patients, their household members, or other close contacts, IPV should be administered. MMR is not contraindicated in close contacts of immunocompromised patients.

Killed or inactivated vaccines can be administered to all immunocompromised patients, although response to such vaccines may be suboptimal. All such childhood vaccines are recommended for immunocompromised persons in usual doses and schedules. Certain vaccines such as pneumococcal vaccine or Hib vaccine are recommended specifically for certain groups of immunocompromised patients, including those with functional or anatomic asplenia.

**HIV Infection.** Limited studies of MMR vaccination in HIV-infected patients have not documented serious or unusual adverse events. Because measles may cause severe illness in persons with HIV infection, MMR vaccine is recommended for all asymptomatic HIV-infected persons and should be considered for all symptomatic HIV-infected persons. HIV-infected persons on regular IGIV therapy may not respond to MMR or its individual component vaccines because of the continued presence of passively acquired antibody. However, because of the potential benefit, measles vaccination should be considered approximately 2 weeks before the next monthly dose of IGIV (if not otherwise contraindicated), although an optimal immune response is unlikely to occur. Unless serologic testing indicates that specific antibodies have been produced, vaccination should be repeated (if not otherwise contraindicated) after the recommended interval. An additional dose of IGIV should be considered for persons on routine IGIV therapy who are exposed to measles 3 or more weeks after administration of a standard dose (100–400 mg/kg) of IGIV.

**Chemotherapy or Radiation Therapy.** Vaccination during chemotherapy or radiation therapy should be avoided because antibody response is poor. Patients vaccinated while on immunosuppressive therapy or in the 2 weeks before starting therapy should be considered unimmunized and should be revaccinated at least 3 months after therapy is discontinued. Patients with leukemia in remission whose chemotherapy has been terminated for 3 months may receive live-virus vaccines.

**Corticosteroid Therapy.** The exact amount of systemically absorbed corticosteroids and the duration of administration needed to suppress the immune system of an otherwise healthy child are not well defined. Most experts agree that corticosteroid therapy usually does not contraindicate administration of live-virus vaccine when it is short term (ie, <2 weeks); low to moderate dose; long-term, alternate-day treatment with short-acting preparations; maintenance physiologic doses (replacement therapy); or administered topically (skin or eyes), by aerosol, or by intra-articular, bursal, or tendon injection. Although of recent theoretical concern, no evidence of increased severe reactions to live vaccines has been reported among persons receiving corticosteroid therapy by aerosol, and such therapy is not in itself a reason to delay vaccination. The immunosuppressive effects of corticosteroid treatment vary, but many clinicians consider a dose equivalent to...
a total of 20 mg/day of prednisone in adults as sufficiently immunosuppressive to raise concern about the safety of vaccination with live-virus vaccines. Corticosteroids used in greater than physiologic doses can also reduce the immune response to vaccines. Physicians should wait at least 3 months after discontinuation of therapy before administering a live-virus vaccine to patients who have received high systemically absorbed doses of corticosteroids for 2 or more weeks.

**Vaccination of Persons with Hemophilia.** Persons with bleeding disorders such as hemophilia have an increased risk of acquiring hepatitis B and at least the same risk as the general population of acquiring vaccine-preventable diseases. However, because of the risk of hematomas, intramuscular injections are often avoided among persons with bleeding disorders by using the subcutaneous or intradermal routes for vaccines that are normally administered by the intramuscular route. Hepatitis B vaccine administered intramuscularly to hemophiliacs using a 23-gauge needle, followed by steady pressure at the site for 1 to 2 minutes has resulted in a 4% bruising rate with no patients requiring clotting factor supplementation. Whether an antigen that produces more local reactions, such as pertussis, would produce an equally low rate of bruising is unknown.

When hepatitis B or any other intramuscular vaccine is indicated for a patient with a bleeding disorder, it should be administered intramuscularly if, in the opinion of a physician familiar with the patient’s bleeding risk, the vaccine can be administered with reasonable safety by this route. If the patient received antihemophilic or other similar therapy, intramuscular vaccination can be scheduled for shortly after such therapy is administered. A fine needle (≤23 gauge) can be used for the vaccination and firm pressure applied to the site (without rubbing) for at least 2 minutes. The patient or family should be instructed concerning the risk of hematoma from the injections.

### REFERENCES

The clinical management of medical emergencies is an area in which there continues to be some variability in treatment philosophy. Thus, the therapeutic approaches, drugs, and adult dosages given here are based on somewhat divergent and conflicting sources of information. In addition, some recommendations have been made based on the authors’ experience and suggestions from specialists and researchers in the field. As a result, the therapeutic concepts and dosages contained herein, although conforming to medical standards, may differ from those advocated by specific practitioners and institutions.

Anaphylaxis

William G. Troutman

Anaphylaxis is a systemic response to exposure to an allergen caused by rapid, IgE-mediated release of histamine and other mediators from tissue mast cells and circulating basophils. Symptoms usually occur within a few seconds or minutes of exposure but can be delayed or recur many hours after apparent resolution. The treatment of anaphylaxis is directed toward its three major presentations: skin manifestations (angioedema, urticaria), respiratory distress (wheezing, stridor and dyspnea from laryngeal edema, laryngospasm and bronchospasm), and hypotension. Upper airway obstruction and cardiovascular collapse are the most common causes of death in anaphylaxis. All specific treatment measures should be accompanied by basic resuscitative measures including clear airway, supplemental oxygen and IV access.

- GENERAL THERAPY AND SKIN MANIFESTATIONS

1. Epinephrine HCl, IM or SC, 0.3–0.5 mg (0.3–0.5 mL of 1:1000 soln), may repeat q 10–15 min. In children, 10 µg/kg up to 500 µg/dose (0.5 mL of 1:1000 soln).
2. Diphenhydramine, IV or IM, 1–2 mg/kg (up to 50 mg) over 5–10 min.
3. Cimetidine, IV, 300 mg over 5 min for urticaria or if hypotension does not respond to fluid replacement and pressors. In children, 3–5 mg/kg IV.
4. Although controversial, corticosteroids such as hydrocortisone phosphate or succinate, IV, 200 mg or methylprednisolone, IV, 1–2 mg/kg might reduce the risk of recurrent or prolonged anaphylaxis.

**RESPIRATORY DISTRESS**

1. Assure adequate oxygenation with supplemental oxygen by mask titrated to an oxygen saturation above 90%.
2. In addition to the general therapy described above, add albuterol, by nebulization, 2.5–5 mg q 20 min. In children, 0.15 mg/kg by nebulization q 20 min.
3. If response is inadequate after 3–4 doses of intermittent albuterol, consider albuterol, by continuous nebulization, 10–15 mg/hr. In children, 0.5 mg/kg/hr by continuous nebulization.

**HYPOTENSION**

1. If response to the general therapy described above is inadequate, give NS or lactated Ringer’s injection, IV, 500–1000 mL initially and continue at high flow rate. In children, 10–20 mL/kg IV initially.
2. Epinephrine HCl, IV continuous infusion, 1 µg/min (as a 1:10,000 or 1:100,000 soln), up to 10 µg/min.
3. Dopamine HCl, IV, 2–5 µg/kg/min, titrate to desired effect.
4. Patients taking β-adrenergic blockers may not respond adequately to epinephrine and fluid replacement and can be adversely affected by unopposed α-adrenergic stimulation from epinephrine. Glucagon, IV, 5–10 mg followed by 1–5 mg/hr by continuous infusion can increase myocardial contractility independent of β-receptors.

**REFERENCES**

Cardiac arrest is a medical emergency requiring a systematic approach. Early recognition must be followed by prompt, effective application of Basic Life Support (BLS) techniques to sustain the patient until Advanced Cardiac Life Support (ACLS) capabilities are available. The management of cardiac arrest is a four-step approach:

- **Recognition and Assessment**
- **Basic Life Support (BLS)**
- **Advanced Cardiovascular Life Support (ACLS)**
- **Postresuscitation Care**

## RECOGNITION AND ASSESSMENT

Verify that respiration and circulation have ceased:

1. Loss of consciousness.
2. Loss of functional ventilation (respiratory arrest or inadequate respiratory effort).

## BASIC LIFE SUPPORT (BLS)

The findings listed above are sufficient to justify the immediate application of BLS techniques. The goal in cardiac arrest is the restoration of spontaneous circulation (ROSC). The first step toward achieving ROSC is prompt initiation of BLS, where the goal is to rapidly and effectively perfuse the tissues with oxygenated blood. A delay in initiating BLS or providing ineffective BLS can result in irreversible hypoxic brain injury.

1. Summon help and resuscitation equipment.
2. Establish an adequate airway.
3. Provide rescue breathing by delivering two slow, deep breaths. Ventilate by mouth-to-mouth, mouth-to-mask, or bag-valve-mask techniques.
4. Check for pulse and other signs of circulation. Lay persons are not expected to perform a pulse check. Rather, they are instructed to look for other signs of circulation such as normal breathing, coughing, or movement. When available, assess heart rhythm with an automated external defibrillator or monophasic/biphasic defibrillator.
   - If ventricular tachycardia or ventricular fibrillation are documented, defibrillate with 200 joules of direct current shock.
• If the first shock fails to terminate the dysrhythmia, a second shock with 200–300 joules should be attempted. If the first two shocks fail, shock again with 360 joules.

5. Reassess cardiac rhythm and check for a pulse. If no pulse or other signs of circulation are present, initiate rescue breathing and chest compressions.

• For rescue breathing:
  —Give each breath slowly over 2 sec.
  —Deliver 10–12 breaths per minute or 1 breath q 4–5 sec.

• For external chest compressions:
  —Position patient supine on a firm surface.
  —Ensure proper placement of hands on sternum.
  —Depress sternum at a rate of 80–100 cycles per min (50% of cycle should be compression).
  —For every 15 chest compressions, give 2 breaths.

### ADVANCED CARDIOVASCULAR LIFE SUPPORT (ACLS)

**Note:** Only adult dosages are given in this section.

Trained personnel should attempt to maintain a patent airway, establish intravenous access for administration of fluids and drugs, establish an electrocardiographic diagnosis, and apply specific treatments to correct any recognized electrical and/or mechanical abnormalities.

#### DRUG THERAPY IN ACLS

**Ventricular Tachyarrhythmias.** In this category, and treated the same way, are unstable ventricular flutter and ventricular tachycardia (pulseless VT), and ventricular fibrillation (VF). All are associated with decreased cardiac output and hypotension.

1. **Electrical defibrillation** with 200 joules. If tachyarrhythmias persist, deliver subsequent shocks with 200, 300 and 360 joules, respectively. Any further shocks should be with 360 joules.
   - Class recommendation: I (excellent supporting evidence)
   - Defibrillation is the only treatment proven to decrease mortality in pulseless VT/VF. The objective is to shock soon and shock often. When drug administration is initiated, the sequence is CPR–drug–shock–repeat or CPR–drug–shock–shock–shock–repeat.

2. For pulseless VT/VF refractory to initial defibrillation, administration of medications should follow the sequence below:
   - **Epinephrine HCl, 1 mg IV push** (10 mL of 1:10,000 solution) q 3–5 min until the ROSC or **vasopressin 40 units** (2 mL of 20 units/mL vial) IV, one dose only. If after 5–10 min there is no response to vasopressin, administer epinephrine as instructed.
   - If IV access has not been established or has been lost, consider administering **epinephrine HCl via endotracheal tube** (2–2.5 times the intravenous dose; see Special Considerations), followed by 3 or 4 rapid
ventilations to aerosolize the drug. There is no evidence to support administration of vasopressin via endotracheal tube.

—Class recommendation: Indeterminate for epinephrine and vasopressin (insufficient data to support class recommendation).

—Epinephrine is used not as an aid to defibrillation but rather to increase perfusion and sustain blood pressure. Epinephrine stimulates α- and β-adrenergic receptors. Stimulation of α-receptors causes vasoconstriction, increasing systemic vascular resistance (SVR) and blood pressure. However, the β-agonist activity of epinephrine increases heart rate and contractility, increasing myocardial oxygen demand in resuscitated patients, and might precipitate or worsen myocardial ischemia.

—Vasopressin is an alternative to epinephrine (at least initially). It is an endogenous antidiuretic hormone that, at high doses (ie, ACLS doses), possesses considerable vasoconstrictor activity. Unlike epinephrine, vasopressin has no β-agonist activity and does not increase myocardial oxygen demand.

3. If pulseless VT/VF persists, the next step is to initiate antiarrhythmic drug therapy. Management has changed in that the initial antiarrhythmic of choice is now:

- **Amiodarone, 300 mg IV push** (6 mL of 50 mg/mL ampule diluted to 20–30 mL of NS or D5W). If pulseless VT/VF persists, give an additional 150 mg IV push. If ROSC occurs, initiate intravenous infusion (450 mg in 250 mL NS, 1.8 mg/mL) at 1 mg/min for 6 hr and then decrease to 0.5 mg/min. Maximum dose is 2.2 g in 24 hr.
  —Class recommendation: IIb (Fair to good supporting evidence).
  —Amiodarone, in addition to its sodium, potassium, and calcium channel blocking activity, possesses α- and β-antagonistic properties. The short-term side effects of amiodarone are bradycardia, hypotension, and QT prolongation. Hypotension, likely secondary to the polysorbate 80 diluent of the injectable formulation, can be prevented by slowing the rate of drug infusion. A polysorbate 80–free formulation of amiodarone is currently under investigation. Bradycardia and QT prolongation might respond to a dose reduction.
  —IV infusions of amiodarone should be admixed in glass bottles because drug adsorption to plastic containers is likely with prolonged exposure. This phenomenon was taken into account during clinical trials, so traditional PVC tubing for administration is acceptable.

4. If amiodarone fails to control the arrhythmia, consider:

- **Lidocaine HCl, 1.0-1.5 mg/kg IV push** (2.5–5 mL of 2% solution or 5–10 mL of 1% solution), may repeat in 3–5 min to a cumulative dose of 3 mg/kg. If the arrhythmia is controlled, initiate an intravenous infusion (1 g/250 mL D5W, 4 mg/mL) at 1–4 mg/min.
  —If IV access has not been established or has been lost, consider administering lidocaine HCl via endotracheal tube (2–2.5 times the intravenous dose; see Special Considerations), followed by 3 or 4 rapid ventilations to aerosolize the drug.
——Class recommendation: Indeterminate (insufficient data to support class recommendation).

Lidocaine is a class Ib antiarrhythmic agent that blocks cellular sodium ion channels and increases the electrical stipulation threshold of the heart. Lidocaine inhibits its own hepatic metabolism after 24–48 hr of therapy; therefore, it should be used with caution in the elderly and in patients with hepatic dysfunction. Signs of toxicity are mental status changes, muscle twitching, seizures, and bradycardia. If prolonged administration is likely, monitoring of serum concentrations might be helpful.

5. If amiodarone- and lidocaine-resistant dysrhythmias persist, consider the administration of:

- Procainamide HCl, 30 mg/min IV infusion, (1 g/250 NS, 4 mg/mL or 2 g/250 mL NS, 8 mg/mL) to a maximum dose of 17 mg/kg. If the arrhythmia terminates with procainamide, initiate an IV infusion at 1–4 mg/min.
  —Class recommendation: IIb for intermittent/recurrent VT/VF (Fair to good supporting evidence).

Procainamide is a class Ia antiarrhythmic agent that blocks the sodium ion channels of the heart. Avoid rapid administration (>30 mg/min) because this can lead to hypotension. Because procainamide must be administered slowly, it is not a first-line antiarrhythmic agent in the management of VT/VF. Serum concentrations of procainamide and its active metabolite N-acetylprocainamide, should be monitored and doses should be decreased in the presence of renal dysfunction. Procainamide also can prolong the QT interval; therefore, it should be avoided in patients with pre-existing QT prolongation and torsades de pointes.

6. If the rhythm is documented polymorphic VT (torsades de pointes) or secondary to hypomagnesemia, administer:

- Magnesium sulfate, 1–2 g IV infusion over 15–30 min. Rapid IV push administration can lead to hypotension, bradycardia, and asystole; therefore, it is not recommended. Consider a maintenance infusion of 0.5–1 g/hr if arrhythmia successfully terminates with magnesium.
  —Class recommendation: IIb (Fair to good supporting evidence).

7. Administering sodium bicarbonate during cardiac arrest has traditionally been a controversial issue. Its use in VT/VF arrests should be considered only after other accepted interventions (eg, defibrillation, intubation/ventilation, chest compressions, and vasopressors) have been ineffective. If desired, administer:

- Sodium bicarbonate, 1 mEq/kg slow IV push (50 mL of 8.4% solution, 1 mEq/mL).

8. Bretylium is no longer recommended by the American Heart Association because of a shortage of natural resources, limited product availability, high occurrence of side effects, and the availability of safer and at least as efficacious agents. It is not featured on the VT/VF algorithm but is still an appropriate choice for treatment after attempting lidocaine.
Bretylium tosylate, 5 mg/kg slow IV push, may repeat with 10 mg/kg q 5 min to a maximum dose of 30–35 mg/kg. If a response to the loading dose occurs, initiate an intravenous infusion (500 mg/250 mL D5W, 2 mg/mL) at 1–2 mg/min.

—Class recommendation: IIb (Fair to good supporting evidence).
—Bretylium is a class III antiarrhythmic that inhibits the potassium channel, prolonging action potential duration and refractoriness of the myocardium. Bretylium also causes a release of catecholamines shortly after administration but subsequently exhibits postganglionic adrenergic receptor blockade, frequently leading to the development of hypotension. Prolongation of the QT interval also can occur.

Pulseless Electrical Activity (PEA). PEA was previously known as electromechanical dissociation and is characterized by ineffective cardiac output (hypotension) in the face of ECG evidence of electrical myocardial activity. Etiologies of PEA can be remembered by the 5 Hs and 5 Ts:

- Hypovolemia
- Hypoxia
- Hydrogen ions (acidosis)
- Hypo/hyperkalemia
- Hypothermia
- Tablets (drugs)
- Tamponade (cardiac)
- Tension pneumothorax
- Thrombosis, coronary
- Thrombosis, pulmonary (embolism)

The most effective way to treat PEA is to correct the underlying cause. The methods discussed below are temporizing measures until the causative etiology is found and remedied.

1. Nonspecific treatment measures include administration of:
   - Epinephrine HCl, 1 mg IV push (10 mL of 1:10,000 solution) q 3–5 min.
   - If bradycardic, give atropine sulfate, 1 mg IV push (10 mL of 0.1 mg/mL solution) every 3–5 min to a maximum dose of 3 mg or 0.04 mg/kg. Atropine may be given via endotracheal tube at 2–2.5 times the intravenous dose (2–2.5 mg; see Special Considerations) followed by 3 or 4 rapid ventilations to aerosolize the drug.
2. The use of buffering agents is controversial. When clinical situations arise where alkalinization is necessary (see Class Recommendations, below), administer:
   - Sodium bicarbonate, 1 mEq/kg slow IV push (50 mL of 8.4% solution, 1 mEq/mL).
     —Class recommendation: I (Excellent supporting evidence) for documented hyperkalemia.
   - In addition to sodium bicarbonate, calcium is indicated for hyperkalemia with ECG changes. Calcium acts as a cardioprotectant and offsets the arrhythmogenic potential of excessive potassium. Administer calcium chloride, 0.5–1 g slow IV push (5–10 mL of
CARDIAC ARREST

10% solution = 6.8–13.6 mEq) or calcium gluconate 1–2 g slow IV push (10–20 mL of 10% solution = 4.7–9.4 mEq).

—Class recommendation: IIa (Good to very good supporting evidence) for bicarbonate-sensitive acidosis, tricyclic antidepressant overdose, or for urine alkalinization in aspirin and other drug overdoses.

—Class recommendation: IIb (Fair to good supporting evidence) following ROSC in mechanically ventilated patients after a prolonged arrest.

—Sodium bicarbonate can be harmful in hypercarbic acidosis; therefore, administration should be limited to those situations described above.

3. Hypovolemia is the most common underlying cause of PEA; therefore, rapid assessment of fluid status is crucial. In hypovolemic patients, fluid resuscitation using crystalloid (NS or lactated Ringer’s solution) or colloid (hetastarch or human albumin) products should be initiated immediately.

4. If volume is adequate and there is no evidence of cardiac tamponade, consider vasopressors for vasoconstrictor and inotropic/chronotropic effects.

• Dopamine HCl, start at 5 µg/kg/min IV infusion (400 mg/500 mL D5W, 800 µg/mL or 800 mg/500 mL D5W, 1600 µg/mL) and titrate to effect (BP and heart rate). Maximum dosage is 20 µg/kg/min. Dosages >20 µg/kg/min have no increased effect on BP and increase the risk for drug-induced tachyarrhythmias.

—Dopamine possesses dopaminergic and α- and β-adrenergic activity. At dosages <5 µg/kg/min, dopaminergic receptor activation causes an increase in renal and mesenteric blood flow. At dosages of 5–10 µg/kg/min, β-adrenergic receptor stimulation (β₁ > β₂) occurs, increasing heart rate and contractility. At dosages >10 µg/kg/min, α-receptor stimulation leads to an increase in SVR and elevation in BP.

• Norepinephrine bitartrate, start at 0.5–1 µg/min IV infusion (4 mg/250 mL D5W, 16 µg/mL, or 8 mg/250 mL D5W, 32 µg/mL) and titrate to effect (BP and heart rate). No maximum dose is noted.

—Norepinephrine stimulates α- and β-adrenergic receptors, increasing BP (secondary to increased SVR), heart rate, and contractility.

—Because increased doses of norepinephrine enhance β-agonist activity (especially in patients with prior cardiac disease), patients are at increased risk for drug-induced tachyarrhythmias.

Asystole. Asystole is characterized by cessation of cardiac muscular and electrical activities. It is important to note that true asystole, unless as a result of excessive vagal tone (bradyasystolic event), is frequently associated with irreversible cardiac damage. Like PEA, the most effective management of the asystolic patient is identifying and treating the underlying causes (see PEA management). However, many times a cause cannot be determined.
1. Initial management of asystole starts with transcutaneous or transvenous pacing, when the capability is available.

2. In conjunction with pacing, medications for managing asystole include:
   - **Epinephrine HCl, 1 mg IV push** (10 mL of 1:10,000 solution) q 3–5 min and **atropine sulfate, 1 mg IV push** (10 mL of 0.1 mg/mL solution) q 3–5 min to a maximum dose 3 mg or 0.04 mg/kg.

If asystole persists, the potential for a successful resuscitation should be evaluated and a decision made to continue or cease resuscitation efforts.

**Bradyarrhythmias.** Considered in this category, and treated the same way, are complete heart block, slow ventricular focus, sinus bradycardia, and agonal rhythm. In dealing with any of these symptomatic arrhythmias, transvenous pacing is the best long-term approach but is often not readily accessible. Therefore, drugs are used to enhance or initiate cardiac activity, at least until transcutaneous or transvenous pacing capabilities are available.

1. If symptomatic bradycardia occurs, initiate management with:
   - **Atropine sulfate, 1 mg IV push** (10 mL of 0.1 mg/mL solution) every 3–5 min to a maximum dose of 3 mg or 0.04 mg/kg.
     — Patients with denervated transplanted hearts will not respond to atropine; therefore, proceed immediately to transcutaneous pacing, administration of catecholamines, or both.

2. If capabilities are available, attempt:
   - **Transcutaneous pacing** to capture the slow rhythm and increase heart rate to a level at which symptoms disappear. If continued pacing is necessary, continue transcutaneous pacing until a transvenous pacer can be placed.

**SUPPORTIVE THERAPY**

**Management of Acidosis.** Severe acidosis can develop within 5 min after cardiac arrest and will continue unless BLS is provided. Acidosis can be respiratory and/or (to a lesser extent) metabolic in etiology.

1. Respiratory Acidosis
   - Secondary to hypoventilation and an accumulation of CO2.
   - Treat by providing adequate ventilation. There is no role for sodium bicarbonate in this situation.

2. Metabolic Acidosis
   - Due to tissue hypoxia and subsequent anaerobic metabolism that results in the slow accumulation of lactic acid.
   - Treat by adequate tissue perfusion and return to aerobic metabolism. Sodium bicarbonate administration is not indicated unless there is evidence of pre-existing acidosis, hyperkalemia, or TCA overdose. There is no evidence supporting routine use of bicarbonate and it should be limited to specific clinical situations.
• If sodium bicarbonate is to be given, the following guidelines should be followed:
  — If an arterial blood gas (ABG) is not available, empirically administer sodium bicarbonate, 1 mEq/kg slow IV push (50 mL of 8.4% solution, 1 mEq/mL).
  — If an ABG is available, the sodium bicarbonate dose can be calculated from the base deficit with the following equation:

\[
\text{NaHCO}_3 \text{ dose in mEq} = \text{base deficit (mEq/L)} \times 0.2 \times \text{body weight (kg)}
\]

■ POSTRESUSCITATION CARE

With the ROSC after cardiac arrest, cardiovascular and hemodynamic compromise is often considerable and can be manifested as different types of shock (hypovolemic, cardiogenic, and vasodilatory associated with systemic inflammatory response syndrome). If the patient is not already in an intensive care setting, transport to an intensive care unit should occur as soon as possible. Continuous monitoring, resuscitation equipment, and skilled nursing care are needed. Health care providers should be diligent in identifying the underlying causes and correcting them, if possible. The goal of postresuscitation care is to restore functional ventilation and maintain adequate tissue perfusion.

■ SPECIAL CONSIDERATIONS

• Time to Drug Effect
  — Systemic circulation times are grossly prolonged during external chest compressions. Remember to allow at least 2 min between the time of peripheral injection and anticipated response. To enhance the onset and activity of peripherally administered medications, give as a rapid bolus injection, followed by a 10–20 mL NS flush and, if possible, elevate the extremity.

• High-dose Epinephrine
  — It was once believed that high-dose epinephrine was more effective than standard ACLS doses. However, recent studies have found no advantage using high-dose over standard-dose epinephrine. There is some also preliminary evidence that higher doses of epinephrine can be harmful in resuscitated patients.

• Endotracheal Administration
  — Administration of epinephrine, lidocaine, and atropine can be done via endotracheal tube if IV access has not been established or has been lost. Doses are 2–2.5 times the IV dose. Undiluted drug (eg, epinephrine 1:1000) can be given, but it must be diluted to 10 mL with NS or followed by a 10-mL NS or sterile water flush. After administration of medications via the endotracheal route, 3 to 4 rapid ventilations should be performed to aerosolize the drug and maximize absorption. This route of administration may not be as effective as IV.
Intraosseous Administration
—Epinephrine, atropine, sodium bicarbonate, lidocaine, vasopressors, or calcium via the distal tibia can be used in situations in which IV access and endotracheal intubation have not been established. This route of administration is often reserved for pediatric patients but may be attempted in adults in rare situations.

Intracardiac Administration
—Administration of medication directly into the myocardium has no role in the modern management of cardiac arrest. Drugs do not work within the chambers of the heart but rather at the cellular level after delivery via the coronary circulation. Stopping BLS to attempt intracardiac injections only serves to interrupt vital CNS perfusion.

Physical Incompatibilities
—With many medications being given during a cardiac arrest (often through the same IV access site), it is important to recognize the likelihood of physical incompatibilities. Sodium bicarbonate inactivates catecholamines and can form a precipitate when mixed with calcium-containing solutions. Concomitant administration should be avoided, if possible. If sodium bicarbonate is administered through the same vascular access site, the line must be flushed before and after bicarbonate administration.

REFERENCES
Management of the poisoned patient involves procedures designed to prevent the absorption, minimize the toxicity, and hasten the elimination of the suspected toxin. The prompt employment of appropriate emergency management procedures often can prevent unnecessary morbidity and mortality.

A regional poison center is a practitioner’s best source of definitive treatment information and should be consulted in all poisonings, regardless of the apparent simplicity of the case. Contact the regional poison center in your area to learn of its staffing, resources, and capabilities before a need for its services arises. Well-qualified regional centers are certified by the American Association of Poison Control Centers.

In all cases, every attempt should be made to accurately identify the toxin, estimate the quantity involved, and determine the time that has passed since the exposure. These data, plus patient-specific parameters such as age, weight, sex, and underlying medical conditions or drug use, will assist you and the regional poison center in designing an appropriate therapeutic plan for the patient.

The techniques described below are intended for the initial management of the poisoned patient with the use of materials that should be readily available.

### TOPICAL EXPOSURES

1. **Immediately** irrigate affected areas with a copious amount of water; use soap only if a stubborn, oily substance is the contaminant. Skin should be gently washed, not scrubbed, and special attention should be given to the hair, skin folds, umbilicus, and other areas where the contaminant might be trapped.

2. If the patient’s clothes have been contaminated, remove them during the irrigation and clean them before they are worn again or destroy them. Clothing can interfere with the irrigation process and serve as a reservoir of toxic material.

3. Do not attempt to “neutralize” the contaminant with another chemical (eg, acids and alkalis). Attempts at neutralization waste valuable time, are of no benefit, and might be harmful.

4. Do not cover the affected area with emollients. These can trap unremoved contaminant against the skin. Severely damaged skin may be temporarily covered with a light, dry dressing.

5. Protect yourself from contamination. Gloves, aprons, or a change of clothes might be necessary.

6. After the irrigation is complete, contact a regional poison center for definitive treatment information.
**EYE EXPOSURES**

1. Immediately irrigate the eye; damage can occur within seconds. The stream of water from the tap or a pitcher should strike the patient on the forehead, temple, or bridge of the nose and then flow into the eye.
2. The eyelids should be open, with frequent blinking during the irrigation.
3. The irrigation should continue for at least 15 min (by the clock) to ensure adequate removal of the contaminant and normalization of the conjunctival pH. Body temperature water or saline may be substituted for tap water as the irrigation proceeds, but only if these can be obtained without interrupting the irrigation.
4. After the irrigation is complete, contact a regional poison center for definitive treatment information.

**INHALATION EXPOSURES**

1. Remove the patient from the suspected contaminated area, regardless of its apparent safety. Carbon monoxide, a common inhaled toxin, cannot be detected by sight, smell, or taste.
2. Institute artificial ventilation, if necessary, and provide supplementary humidified oxygen, if available and needed.
3. Protect yourself from contamination at all times.
4. Contact a regional poison center for definitive treatment information.

**INGESTIONS**

1. Remove any remaining contaminant from inside and around the mouth of the patient.
2. Give a small amount of water to clear the mouth and esophagus.
3. Contact a regional poison center for definitive treatment information.
4. In many cases, it will not be necessary to take additional steps. The following information can be used if additional care is recommended by the regional poison center.

**GASTROINTESTINAL DECONTAMINATION**

Gastrointestinal (GI) decontamination can be accomplished by the administration of activated charcoal, gastric lavage, ipecac-induced emesis, or whole-bowel irrigation. Indications for GI decontamination are ingestion of a known toxic dose, ingestion of an unknown dose of a known toxic substance, and ingestion of a substance of unknown toxicity. For all methods of GI decontamination, the value of the procedure diminishes rapidly with time. Some investigators now question the usefulness of gastric lavage or ipecac-induced emesis more than 1 hr after ingestion. None of these techniques should be presumed to provide complete removal or binding of the ingested toxin(s). Comparative experimental studies have shown only limited success with these techniques, and there is considerable interpatient variability in the results. In general, activated charcoal is the most useful agent for
preventing absorption of ingested toxic substances. Other methods of GI decontamination may be considered if the ingested contaminant is not adsorbed by activated charcoal or if circumstances do not permit its prompt administration.

**ACTIVATED CHARCOAL**

Activated charcoal is a nonspecific absorbent that binds unabsorbed toxins within the GI tract. There is limited experience using activated charcoal in the home setting. Activated charcoal is not effective for absorbing strong acids and alkalis, cyanide, ethanol, methanol, ethylene glycol, iron, or lithium.

1. Activated charcoal is administered orally or by gastric tube in doses that range from 30 to 120 g.
2. Activated charcoal is commercially supplied as a slurry in water or a concentrated solution of sorbitol. The water-based products are preferred because the large amount of sorbitol that accompanies a typical dose of activated charcoal can result in excessive sorbitol-induced catharsis, producing fluid and electrolyte imbalance. Gentle encouragement may be needed to make children swallow the charcoal. Having the child take the liquid through a drinking straw from an opaque container is sometimes helpful.
3. Activated charcoal administration is commonly followed by the administration of a cathartic (eg, sorbitol, magnesium citrate, or magnesium sulfate) to hasten the elimination of the activated charcoal–toxin complex. There is no evidence to support cathartic use.
4. Alert the patient that charcoal will cause the stools to turn black.
5. Repeated oral doses of activated charcoal (eg, 25 g q 2 hr) have been used to enhance the elimination of some drugs, most notably carbamazepine, dapsone, phenobarbital, quinine, or theophylline. Multiple-dose activated charcoal is suitable only for patients with active bowel sounds. Co-administration of a cathartic is not recommended during multiple-dose activated charcoal therapy.

**GASTRIC LAVAGE**

Gastric lavage can be used to remove toxic substances poorly adsorbed by activated charcoal. Lavage is contraindicated for patients who have ingested corrosives or aliphatic hydrocarbons (ie, gasoline) and for patients at risk for esophageal or gastric perforation due to underlying medical conditions (eg, esophageal varices).

1. If the patient’s gag reflex is weak or absent, the airway must be protected by the use of a cuffed endotracheal tube.
2. The largest possible orogastric tube should be used (26–28 F for children and 34–42 F for adults): the larger the tube diameter, the more efficient the lavage. The tube should be introduced through the mouth with the aid of a water-soluble lubricant. Nasogastric passage is not recommended.
3. Gastric lavage may be done with water, but a solution such as 0.45% NaCl may be used to minimize the risk of dilutional hyponatremia, especially in children. Aliquots of fluid up to 100 mL in children and 200 mL...
in adults are introduced through the tube and then removed by gravity or suction-assisted drainage. The lavage should be continued for several cycles after the returning fluid is clear. Warming the lavage fluid reduces the risk of hypothermia.

INDUCTION OF EMESIS

Do not induce emesis if the patient is experiencing or is at risk for CNS depression, seizures, or loss of gag reflex, or if the patient has ingested a caustic substance or a hydrocarbon with high aspiration potential (e.g., gasoline).

1. Induce emesis only with syrup of ipecac. Salt water, mustard water, other “home remedies,” or gagging have no place in the management of the poisoned patient. These techniques are ineffective and can be dangerous.

2. The usual initial dose of syrup of ipecac is 30 mL in persons older than 12 yr, 15 mL in children 1–12 yr old, and 10 mL in children between 6 months and 1 yr.

3. Give the patient additional water to drink: 125–250 mL (4–8 fluid ounces) in children, 250–500 mL (8–16 fluid ounces) in adults. Activated charcoal should not be given until after ipecac-induced emesis has occurred.

4. Emesis usually occurs within 15–20 min. If 30 min have passed without emesis, administer an additional dose of syrup of ipecac and more water.

5. Have the patient vomit into a bowl or other container so that the vomitus can be inspected for the presence of the ingested toxin.

WHOLE-BOWEL IRRIGATION

Whole-bowel irrigation with an orally administered polyethylene glycol electrolyte solution (e.g., GoLYTELY or CoLyte) is commonly used before bowel procedures. It has drawn attention as an alternative to other methods of GI decontamination in the management of acute poisoning. Results of studies are promising and the technique may have value in cases of ingestion of iron, enteric-coated or sustained-release products, foreign bodies, and drug-smuggling packets. Instillation rates have ranged from 500 mL/hr in children to 2 L/hr in adults. Typically, 4–6 L of fluid is administered. The endpoint is clearing of the rectal effluent. Contraindications to whole-bowel irrigation are persistent vomiting, adynamic ileus, bowel obstruction or perforation, and GI hemorrhage.

REFERENCES

Status Epilepticus

Brian K. Alldredge

Status epilepticus is a medical emergency in which prompt recognition and effective medical intervention are required to reduce the risk of permanent sequelae and death. Status epilepticus is defined as continuous seizures lasting at least 5 min, or two or more sequential seizures without full recovery of consciousness between seizures.

Status epilepticus can be categorized into two major types: convulsive and nonconvulsive. Convulsive status epilepticus is associated with the highest risk of morbidity and mortality, so this section focuses on the clinical features and management of this form of status epilepticus.

In about one-half of patients, status epilepticus is the first manifestation of seizures. The causes of status epilepticus are similar to those for new-onset seizures and include CNS infection, cerebral tumor, trauma, stroke, metabolic disorders, cardiopulmonary arrest, and drug toxicity. In the remainder of patients, status occurs in the setting of a pre-existing seizure disorder. Among persons with a history of epilepsy, antiepileptic drug withdrawal (usually noncompliance with prescribed therapy) is the most common cause of status epilepticus.

The primary determinant of patient outcome after status epilepticus is the underlying cause of the episode. In general, patients with status caused by an acute or progressive neurologic insult (eg, cardiopulmonary arrest, stroke) have poorer outcomes than patients in whom status epilepticus occurs in the setting of a more chronic or stable underlying condition (eg, antiepileptic drug withdrawal or medically refractory epilepsy). Nonetheless, aggressive medical intervention and administration of effective antiepileptic drug therapy are important to reduce status-related morbidity and mortality, regardless of the etiology.

Status epilepticus should be managed in an emergency department or an environment where continuous skilled medical and nursing support are available. The emergency management of status epilepticus should include the following:

- Ensure airway patency and adequate oxygenation.
- Obtain blood specimens for baseline laboratory measurements, including CBC, serum electrolytes (including calcium and magnesium), screen, and anticonvulsant serum levels.
- Establish IV access.
- Administer IV glucose (100 mg thiamine followed by 50 mL of 50% glucose in adults).
- Administer IV antiepileptic drugs.
- Monitor BP, respiratory rate and temperature. Treat hyperthermia with passive cooling.
- Obtain other diagnostic studies as needed.
- Treat precipitating factors.
DRUG THERAPY OF STATUS EPILEPTICUS

Adult doses only are given in this section.

If a treatable cause of status epilepticus can be identified rapidly, then drug therapy to terminate seizures might be unnecessary. In these situations, treatment of the underlying cause might be sufficient to stop status. Examples are status caused by an acute metabolic derangement (where correction of the underlying abnormality often stops seizures) or status after isoniazid overdose (where IV pyridoxine is usually effective). However, when a treatable cause is not known, drug therapy should begin immediately. The goal of drug treatment is to terminate seizures as rapidly as possible. Evidence from animal and human studies indicate that 60–120 min of status epilepticus is associated with neurologic sequelae and that the risk increases as status continues. Thus, it is important to have a clear, stepwise plan for the administration of effective drug therapy. Figure 4–1 shows an example of a status epilepticus treatment protocol. In addition, adequate support should be available to manage cardiac and respiratory complications that might occur during drug administration.

1. For rapid termination of seizures:
   - **Lorazepam, IV, 0.1 mg/kg (4–8 mg) at rate of 2 mg/min;** may repeat in 10 min if seizures continue (to maximum of 0.2 mg/kg). Lorazepam has a longer duration of anticonvulsant effect than diazepam and is often preferred for this reason.

![Figure 4–1. Timeline for administration of drug therapy for convulsive status epilepticus. Heavy bars (■) indicate duration (in minutes) of intravenous drug administration. PE = phenytoin equivalents.](ch14.qxd_8/13/2001_3:20_PM_Page_1016)
• Diazepam, IV, 0.2 mg/kg (5–10 mg) at rate of 5 mg/min; may repeat in 10 min if seizures continue (to maximum of 20 mg). Diazepam has a short duration of anticonvulsant effect (15–60 min) and must be immediately followed by a long-acting agent (eg, phenytoin).

2a. After benzodiazepine administration, give:
• Phenytoin, IV infusion, 20 mg/kg at rate of 50 mg/min or fosphenytoin 20 mg/kg phenytoin equivalents IV at a rate of 150 mg/min. Monitor BP and ECG during administration of phenytoin or fosphenytoin loading dose. Elderly and severely ill patients are predisposed to phenytoin-related hypotension.

2b. If status persists, then:
• Phenytoin or fosphenytoin, IV, up to 2 additional doses of 5 mg/kg, to a total dosage of 30 mg/kg.

2c. If seizures are not terminated after administration of phenytoin or fosphenytoin 30 mg/kg, then:
• Phenobarbital, IV, 20 mg/kg at rate of 100 mg/min. The risk of hypoventilation is increased markedly when phenobarbital is administered after a benzodiazepine; respiratory support is often required.

3. If seizures are not terminated after administration of phenytoin or fosphenytoin 30 mg/kg, then:
• Phenobarbital, IV, 20 mg/kg at rate of 100 mg/min. The risk of hypoventilation is increased markedly when phenobarbital is administered after a benzodiazepine; respiratory support is often required.

4. For patients who continue in status epilepticus despite the above recommendations, anesthetic doses of a benzodiazepine, barbiturate, or propofol are often required to suppress seizure activity. Ventilatory assistance and vasopressor drug therapy are often required; therefore, the patient should be admitted to the ICU and the following therapies considered:

4a. Midazolam, IV slow push, 200 µg/kg, then maintenance:
• Midazolam, IV infusion, 0.75–10 µg/kg/min. High-dose midazolam is probably associated with a lower risk of hypotension than high-dose pentobarbital; however, there is less experience with its use.

4b. Propofol, IV slow push, 1 mg/kg, then maintenance:
• Propofol, IV infusion 2–4 mg/kg/hr. Reduce dosage by one-half in elderly or hemodynamically unstable patients. The EEG should be monitored continuously during the first 1–2 hr of therapy, and infusion rates should be adjusted until suppression of electrographic seizures is evident. After seizures are terminated, the rate of the maintenance infusion can be slowed periodically to determine if status has remitted.

4c. Pentobarbital, IV infusion, 1–2 mg/kg/hr. Hypotension is a frequent complication of high-dose pentobarbital therapy; a vasopressor (eg, dopamine) may be required.

REFERENCES
Cytochrome P450 enzymes are found throughout the body and play an important role in the metabolism of many drugs by catalyzing α-hydroxylation, N-demethylation, ring oxidation, and more. Most substrates are metabolized by a specific enzyme, whereas each cytochrome P450 enzyme is generally capable of metabolizing many different compounds. Induction or inhibition of these enzymes can dramatically affect the outcome of drug therapy.

Cytochrome P450 enzymes are identified by the prefix “CYP” followed by an Arabic number identifying the family, although Roman numerals are still sometimes used. The three important enzyme families in humans are CYP1, CYP2, and CYP3. Subfamilies are given letters (eg, CYP2B, CYP2C) that are followed by numbers identifying the specific enzyme.

Although most concentrated in the liver, cytochrome P450 enzymes exist in all tissues of the human body. Intestinal mucosal cytochrome P450 enzymes appear to be primarily from the CYP3A family, probably CYP3A4 in humans. These enzymes affect the bioavailability of some drugs.

**INDUCTION AND INHIBITION**

When the amount of enzyme present in the body is increased by a drug or chemical, the enzyme is said to be “induced.” Although most inducers are P450 substrates, this is not always the case. Induction can increase the rate of clearance of a drug, decreasing its efficacy. It also can increase the rate of formation of an active or toxic metabolite, resulting in exaggeration of therapeutic effect or increased toxicity.

Theoretically, all substrates metabolized by the same enzyme can compete for the same binding site, causing competitive inhibition. However, the clinical relevance depends on the concentrations, relative affinities, and other elimination pathways of each substrate. Like inducers, not all inhibitors are enzyme substrates. Some drugs or their metabolites can form an inactive complex with a cytochrome P450 enzyme or its heme group. Inhibition can lead to increased toxic...
effects by causing drug accumulation, or it can lower toxic or therapeutic effects by decreasing the amount of toxic or active metabolite(s).

# DRUG INTERACTIONS

Knowing which drugs are metabolized by each cytochrome P450 enzyme and the drugs that influence those enzymes can help in predicting drug–drug interactions. However, there are additional points to consider when predicting drug interactions.

The effect of inhibition on drug elimination depends partly on whether a substrate has alternate elimination pathways. Inhibition of an enzyme might not be clinically important if there are alternative metabolic pathways. However, phenytoin, which is metabolized by CYP2C9 and CYP2C19, can interact with CYP2C9 and CYP2C19 inhibitors, resulting in phenytoin toxicity.

Therapeutic range also is important. If a drug has a wide therapeutic range, factors such as induction or inhibition might be clinically unimportant. The opposite is true for drugs with a narrow therapeutic range, such as tricyclic antidepressants and antiarrhythmics.\(^4\,5\)

Last, consider metabolites. Not only does inhibition and induction of cytochrome P450 enzymes influence the formation of active metabolites, the formation of active metabolites can enhance inhibition or induction. Fluoxetine, an inhibitor of CYP2D6, has an active metabolite norfluoxetine, which also inhibits CYP2D6.\(^6\,7\)

The following table is meant to serve as an aid in the prediction of drug–drug interactions. However, it is also important to consider many other parameters: whether the patient is a poor or extensive metabolizer, the affinity of the drug for the binding site, the concentration of drug in the liver, the presence of alternate elimination pathways, and the therapeutic range. Because research on P450 metabolism is currently being published at a rapid rate, the table is not complete. The absence of a drug from the table does not necessarily imply that it is not metabolized by one of the P450 enzymes. When using the table, consider the following principles:

- Inhibition of drug metabolism tends to be substrate independent. That is, a potent inhibitor of CYP2D6 is likely to inhibit the metabolism of any drug metabolized by CYP2D6.
- The magnitude of cytochrome P450 enzyme inhibition is usually dose related over the dosage range of the inhibitor. For example, fluconazole 100 mg/day is usually a modest inhibitor of CYP3A4, but at 400 mg/day it can substantially inhibit the isozyme.
- Some cytochrome P450 inhibitors affect more than one enzyme. For example, ritonavir inhibits both CYP2D6 and CYP3A4.
- Drug enantiomers can be metabolized by different cytochrome P450 isozymes. For example, (R)-warfarin is metabolized by CYP1A2 and CYP3A4, and the more potent (S)-warfarin is metabolized primarily by CYP2C9. Thus, CYP1A2 or CYP3A4 inhibitors tend to produce only small increases in the hypoprothrombinemic response to warfarin, and CYP2C9 inhibitors produce large increases in warfarin effect.
<table>
<thead>
<tr>
<th>SUBFAMILY SUBSTRATES</th>
<th>SUBFAMILY SUBSTRATES</th>
<th>INDUCERS</th>
<th>INHIBITORS</th>
</tr>
</thead>
<tbody>
<tr>
<td>1A2</td>
<td>acetaminophen, amitriptyline, antipyrine, caffeine, clomipramine, clozapine, imipramine, olanzapine, propranolol, tacrine, theophylline, (R)-warfarin, zileuton</td>
<td>charcoal-broiled food, omeprazole, smoking</td>
<td>ciprofloxacin, enoxacin, fluvoxamine, macrolides, mexiletine, tacrine, zileuton</td>
</tr>
<tr>
<td>2B6</td>
<td>cyclophosphamide, ifosfamide</td>
<td>phenobarbital, phenytoin</td>
<td></td>
</tr>
<tr>
<td>2C8</td>
<td>benzphetamine, cervastatin, diazepam, diclofenac, (R)-mephenytoin, paclitaxel, pioglitazone, rosiglitazone, tolbutamide</td>
<td>barbiturates, carbamazepine, phenytoin, primidone, rifampin</td>
<td>amiodarone, clopidogrel, disulfiram, efavirenz, fluconazole, fluoxetine, fluvastatin, metronidazole, miconazole (IV), ritonavir, sulfamethoxazole, sulfa-phenazole, sulfipyrazone, zafirlukast</td>
</tr>
<tr>
<td>2C9/10</td>
<td>celecoxib, diclofenac, dronabinol, flubiprofen, hexobarbital, ibuprofen, losartan, (R)-mephenytoin, montelukast, naproxen, phenytoin, piroxicam, tolbutamide, torsemide, (S)-warfarin</td>
<td></td>
<td>amiodarone, clopidogrel, disulfiram, efavirenz, fluconazole, fluoxetine, fluvastatin, metronidazole, miconazole (IV), ritonavir, sulfamethoxazole, sulfa-phenazole, sulfipyrazone, zafirlukast</td>
</tr>
<tr>
<td>2C18</td>
<td>cimetidine, (S)-mephenytoin, propranolol, retinoic acid</td>
<td>omeprazole, piroxicam</td>
<td></td>
</tr>
<tr>
<td>2C19</td>
<td>amitriptyline, clomipramine, diazepam, hexobarbital, imipramine, lansoprazole, mephenytoin, mepho- barbital, omeprazole, pantoprazole, phenytoin, propranolol, rabeprazole</td>
<td>rifampin</td>
<td>efavirenz, felbamate, fluoxetine, fluvoxamine, omeprazole, ritonavir, ticlopidine</td>
</tr>
<tr>
<td>2D6</td>
<td>chlorpheniramine, codeine, debrisoquine, dextromethorphan, flecainide, fluoxetine, galantamine haloperidol, hydrocodone, loratadine, metoprolol, mexiletine, paroxetine, perphenazine, propafenone, propranolol, risperidone, thiordazine, timolol, tramadol, trazodone, tricyclic antidepressants, venlafaxine, voriconazole</td>
<td></td>
<td>amiodarone, chloroquine, cimetidine, diphenhydramine, fluoxetine, haloperidol, paroxetine, perphenazine, propoxyphene, quinidine, ritonavir, SSRIs, terbinafine, thiordazine</td>
</tr>
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<td></td>
<td></td>
<td></td>
<td>(continued)</td>
</tr>
<tr>
<td>SUBFAMILY</td>
<td>SUBSTRATES</td>
<td>INDUCERS</td>
<td>INHIBITORS</td>
</tr>
<tr>
<td>-----------</td>
<td>------------</td>
<td>----------</td>
<td>------------</td>
</tr>
<tr>
<td>2E1</td>
<td>acetaminophen, alcohol, chlorzoxazone, dapsone, halothane, isoflurane, methoxyflurane, sevoflurane</td>
<td>alcohol (chronic), isoniazid</td>
<td>alcohol (acute intoxication), disulfiram</td>
</tr>
<tr>
<td>3A3/4</td>
<td>alfentanil, alprazolam, amiodarone, amitriptyline, amlodipine, androgens, astemizole, atorvastatin, benzphetamine, bepridil, bromocriptine, buspirone, carbamazepine, clofazimine, cisapride, clomipramine, clonazepam, cocaine, corticosteroids, cyclosporine, dapsone, dexamethasone, diazepam, diltiazem, disopyramide, doxorubicin, ergotamine, erythromycin, ethinyloestradiol, ethosuximide, etoposide, felodipine, fentanyl, fexofenadine, finasteride, galantamine, hydrocortisone, ifosfamide, imatinib, imipramine, indinavir, isradipine, itraconazole, ketoconazole, lidocaine, losartan, lovastatin, miconazole, midazolam, miltefosine, montelukast, nefazodone, nelfinavir, nicardipine, nifedipine, nimodipine, nisoldipine, nitrendipine, omeprazole, paclitaxel, pimozone, pioglitazone, progesterone, propafenone, quinidine, quinine, rifabutin, ritonavir, saquinavir, sertraline, sibutramine, sildenafil, simvastatin, sirolimus, tacrolimus, tamoxifen, teniposide, testosterone, theophylline, triazolam, troglitazone, verapamil, vinca alkaloids, voriconazole, (R)-warfarin, zopolrestat</td>
<td>aminoglutethimide, barbiturates, carbamazepine, corticosteroids, efavirenz, griseofulvin, phenytoin, primidone, rifabutin, rifampin, sulfipyrazone</td>
<td>cyclophosphamide, cyclosporine, delavirdine, diltiazem, fluconazole, fluoxetine, grapefruit juice, ifosfamide, indinavir, itraconazole, ketoconazole, macrolides, metronidazole, miconazole (IV), nefazodone, nelfinavir, nicardipine, nifedipine, quinidine, ritonavir, verapamil, zafirlukast</td>
</tr>
</tbody>
</table>

*CYP3A4 enzyme inhibition by macrolide antibiotics varies by drug: troglitazone > erythromycin > clarithromycin > azithromycin = dirithromycin = 0.*

*CYP2D6 enzyme inhibition by SSRI varies by drug: paroxetine = fluoxetine >> citalopram > fluvoxamine.*

*Also an inhibitor of p-glycoprotein.*

*Also a substrate of p-glycoprotein.*

Compiled from references 3, 8–36.
REFERENCES
23. Manufacturer’s Product Information.
Drug-Induced Discoloration of Feces and Urine

The drugs and drug classes in the following tables have been associated with the discoloration of feces or urine. Drugs and drug classes are listed generically.

### DRUGS THAT CAN DISCOLOR FECES

<table>
<thead>
<tr>
<th>DRUG/DRUG CLASS</th>
<th>COLOR PRODUCED</th>
</tr>
</thead>
<tbody>
<tr>
<td>Antacids, Aluminum Hydroxide Types</td>
<td>Whitish or speckling</td>
</tr>
<tr>
<td>Anthraquinones</td>
<td>Brownish staining of rectal mucosa</td>
</tr>
<tr>
<td>Antibiotics, Oral</td>
<td>Greenish gray</td>
</tr>
<tr>
<td>Anticoagulants</td>
<td>Pink to red or black</td>
</tr>
<tr>
<td>Bismuth Salts</td>
<td>Greenish black</td>
</tr>
<tr>
<td>Charcoal</td>
<td>Black</td>
</tr>
<tr>
<td>Clofazimine</td>
<td>Red to brownish black</td>
</tr>
<tr>
<td>Ferrous Salts</td>
<td>Black</td>
</tr>
<tr>
<td>Heparin</td>
<td>Pink to red or black</td>
</tr>
<tr>
<td>Indocyanine Green</td>
<td>Green</td>
</tr>
<tr>
<td>Indomethacin</td>
<td>Green because of biliverdinemia</td>
</tr>
<tr>
<td>NSAIDs</td>
<td>Pink to red or black</td>
</tr>
<tr>
<td>Omeprazole</td>
<td>Discoloration</td>
</tr>
<tr>
<td>Phenazopyridine</td>
<td>Orange-red</td>
</tr>
<tr>
<td>Pyrvinium Pamoate</td>
<td>Red</td>
</tr>
<tr>
<td>Rifampin</td>
<td>Red-orange</td>
</tr>
<tr>
<td>Risperidone</td>
<td>Discoloration</td>
</tr>
<tr>
<td>Salicylates (especially Aspirin)</td>
<td>Pink to red or black</td>
</tr>
</tbody>
</table>

*a*These colors can indicate intestinal bleeding.

### DRUGS THAT CAN DISCOLOR URINE

<table>
<thead>
<tr>
<th>DRUG/DRUG CLASS</th>
<th>COLOR PRODUCED</th>
</tr>
</thead>
<tbody>
<tr>
<td>Aminopyrine</td>
<td>Red</td>
</tr>
<tr>
<td>Aminosalicylic Acid</td>
<td>Discoloration; red in hypochlorite solution</td>
</tr>
</tbody>
</table>

(continued)
### DRUGS THAT CAN DISCOLOR URINE (continued)

<table>
<thead>
<tr>
<th>DRUG/DRUG CLASS</th>
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</tr>
</thead>
<tbody>
<tr>
<td>Amitriptyline</td>
<td>Blue-green</td>
</tr>
<tr>
<td>Anthraquinones</td>
<td>Yellow-brown in acid urine; yellow-pink-red in alkaline urine</td>
</tr>
<tr>
<td>Antipyrine</td>
<td>Red-brown</td>
</tr>
<tr>
<td>Azuresin</td>
<td>Blue or green</td>
</tr>
<tr>
<td>Chloroquine</td>
<td>Rust yellow to brown</td>
</tr>
<tr>
<td>Chlorzoxazone</td>
<td>Orange or purplish red</td>
</tr>
<tr>
<td>Cimetidine (injection)</td>
<td>Green</td>
</tr>
<tr>
<td>Clofazimine</td>
<td>Red to brownish black</td>
</tr>
<tr>
<td>Daunorubicin</td>
<td>Red</td>
</tr>
<tr>
<td>Deferoxamine</td>
<td>reddish</td>
</tr>
<tr>
<td>Doxorubicin</td>
<td>Red</td>
</tr>
<tr>
<td>Enacapone</td>
<td>Brownish-orange</td>
</tr>
<tr>
<td>Ethoxazene</td>
<td>Orange to orange-brown</td>
</tr>
<tr>
<td>Ferrous Salts</td>
<td>Black</td>
</tr>
<tr>
<td>Flutamide</td>
<td>Amber or yellow-green</td>
</tr>
<tr>
<td>Furozolidone</td>
<td>Brown</td>
</tr>
<tr>
<td>Idarubicin</td>
<td>Red</td>
</tr>
<tr>
<td>Indandiones</td>
<td>Orange-red in alkaline urine</td>
</tr>
<tr>
<td>Indomethacin</td>
<td>Green because of biliverdinemia</td>
</tr>
<tr>
<td>Iron Sorbitex</td>
<td>Brown-black</td>
</tr>
<tr>
<td>Levodopa</td>
<td>Red-brown; dark on standing in hypochlorite solution</td>
</tr>
<tr>
<td>Loratadine</td>
<td>Discoloration</td>
</tr>
<tr>
<td>Methocarbamol</td>
<td>Dark to brown, black or green on standing</td>
</tr>
<tr>
<td>Metyldopa</td>
<td>Dark on standing in hypochlorite solution</td>
</tr>
<tr>
<td>Methylene Blue</td>
<td>Blue or green</td>
</tr>
<tr>
<td>Metronidazole</td>
<td>Dark, brown</td>
</tr>
<tr>
<td>Mitoxantrone</td>
<td>Blue-green</td>
</tr>
<tr>
<td>Niacin</td>
<td>Dark</td>
</tr>
<tr>
<td>Nitrofurantoin</td>
<td>Rust yellow to brown</td>
</tr>
<tr>
<td>Pamaquine</td>
<td>Rust yellow to brown</td>
</tr>
<tr>
<td>Phenacelid</td>
<td>Dark brown to black on standing</td>
</tr>
<tr>
<td>Phenazopyridine</td>
<td>Orange to red</td>
</tr>
<tr>
<td>Phenolphthalein</td>
<td>Pink to purplish red in alkaline urine</td>
</tr>
<tr>
<td>Phenothiazines</td>
<td>Pink to red or red-brown</td>
</tr>
</tbody>
</table>

(continued)
**DRUGS THAT MIGHT DISCOLOR URINE (continued)**

<table>
<thead>
<tr>
<th>DRUG/DRUG CLASS</th>
<th>COLOR PRODUCED</th>
</tr>
</thead>
<tbody>
<tr>
<td>Phensuximide</td>
<td>Pink to red or red-brown</td>
</tr>
<tr>
<td>Phenytoin</td>
<td>Pink to red or red-brown</td>
</tr>
<tr>
<td>Primaquine</td>
<td>Rust yellow to brown</td>
</tr>
<tr>
<td>Promethazine (injection)</td>
<td>Green</td>
</tr>
<tr>
<td>Propofol (injection)</td>
<td>Green, white, pink, brown, or red-brown</td>
</tr>
<tr>
<td>Quinacrine</td>
<td>Deep yellow in acidic urine</td>
</tr>
<tr>
<td>Quinine</td>
<td>Brown to black</td>
</tr>
<tr>
<td>Resorcinol</td>
<td>Dark green</td>
</tr>
<tr>
<td>Riboflavin</td>
<td>Yellow fluorescence</td>
</tr>
<tr>
<td>Rifabutin</td>
<td>Discoloration</td>
</tr>
<tr>
<td>Rifampin</td>
<td>Red-orange</td>
</tr>
<tr>
<td>Sulfasalazine</td>
<td>Orange-yellow in alkaline urine</td>
</tr>
<tr>
<td>Sulfonamides, Antibacterial</td>
<td>Rust yellow to brown</td>
</tr>
<tr>
<td>Sulindac</td>
<td>Discoloration</td>
</tr>
<tr>
<td>Tolcapone</td>
<td>Bright yellow</td>
</tr>
<tr>
<td>Tolonium</td>
<td>Blue-green</td>
</tr>
<tr>
<td>Triamterene</td>
<td>Pale blue fluorescence</td>
</tr>
<tr>
<td>Warfarin</td>
<td>Orange</td>
</tr>
</tbody>
</table>

*Hypochlorite solution in toilet bowl from prior use of chlorine bleach.
*Caused by phenol as a preservative in the injectable formulation.

**REFERENCES**

Nutrition status is a major determinant of patients’ morbidity and mortality. Morbidity increases with malnutrition, as manifested by depressed immunocompetence and impaired wound healing. Conditions that indicate a possible need for nutrition support are inadequate oral nutrition for longer than 7 days, recent body weight loss >10%, an illness lasting longer than 3 weeks, recent major surgery, a lymphocyte count <1.2 × 10^9/L, serum albumin <3 g/dL, serum transferrin <150 mg/dL, and serum prealbumin <15 mg/dL. Sepsis, trauma, and other factors that induce hypermetabolism might intensify the need.

The term “nutrition support” can be applied to any nutrition regimen that is provided for conditions that preclude the use of regular foods. There are two broad categories of nutrition support, enteral and parenteral, determined by their route of administration. Enteral nutrition applies to regimens provided via any portion of the GI tract. Parenteral nutrition (PN), although implying all routes other than the GI tract, refers primarily to regimens that are provided directly by the intravenous route of administration. Less frequently used modes of PN such as intradialytic parenteral nutrition and intraperitoneal nutrition are not discussed in this chapter.

Whenever possible, maintenance rather than repletion should be the primary objective of nutrition support. Early provision of nutrition requirements without exceeding energy balance promotes the synthesis of lean body mass rather than adipose tissue.

**NUTRITION ASSESSMENT**

Nutrition assessment of the patient can aid in diagnosing malnutrition and determining its degree of severity, so that a proper nutrition support regimen can be formulated. The patient’s physical and dietary history should be obtained to establish baseline data. Clinical parameters for assessing the patient’s nutrition status can be evaluated through the use of an assessment form (Table 6–1). Because a patient’s nutrition status is best reflected by body protein, nutrition assessment should focus on the protein compartments. Protein compartments are classified into two types: somatic (muscle protein) and visceral (all other protein).
TABLE 6–1. NUTRITION ASSESSMENT

<table>
<thead>
<tr>
<th>NAME: AGE: HT (CM):</th>
<th>DEPLETION</th>
</tr>
</thead>
<tbody>
<tr>
<td>DATE: SEX: WT (KG):</td>
<td>STANDARD</td>
</tr>
<tr>
<td>TSF (mm)</td>
<td>M 12.5</td>
</tr>
<tr>
<td></td>
<td>F 16.5</td>
</tr>
<tr>
<td>Ideal Body Weight:</td>
<td></td>
</tr>
<tr>
<td>ABW × 100 =</td>
<td>100%</td>
</tr>
<tr>
<td>MUAC (cm)</td>
<td>M 29.3</td>
</tr>
<tr>
<td></td>
<td>F 28.5</td>
</tr>
<tr>
<td></td>
<td>M 25.3</td>
</tr>
<tr>
<td>MUAMC: MUAC (cm) − (0.314 × TSF [mm]) =</td>
<td>F 23.2</td>
</tr>
<tr>
<td>Creatinine/Height Index:</td>
<td></td>
</tr>
<tr>
<td>C₄H₅ × 100 =</td>
<td>100%</td>
</tr>
<tr>
<td>Serum Albumin (g/dL)</td>
<td>3.5–5.0</td>
</tr>
<tr>
<td>Serum Prealbumin (mg/dL)</td>
<td>&gt;20</td>
</tr>
<tr>
<td>Serum Transferrin (mg/dL)</td>
<td>200–400</td>
</tr>
<tr>
<td>Total Lymphocytes/μL:</td>
<td>1800–3000</td>
</tr>
</tbody>
</table>

*The standards specified represent those of healthy persons. Measurements in patients can be affected by nonnutritional and nutritional factors.

ABW, actual body weight; C₄H₅, urinary creatinine; IBW, ideal body weight; IC₄H₅, ideal urinary creatinine; MUAC, mid–upper arm circumference; MUAMC, mid-upper arm muscle circumference; TSF, triceps skinfold.

SOMATIC PROTEIN ASSESSMENT PARAMETERS

Percentage Ideal Body Weight. A simple initial measurement of a patient’s nutrition status is body weight expressed as a percentage of ideal body weight. (See Appendix 2, Anthropometrics.)

\[
\text{Percentage Ideal Body Weight} = \frac{\text{Actual Body Weight}}{\text{Ideal Body Weight}} \times 100
\]

Creatinine/Height Index. Creatinine/height index (CHI), when accurately obtained, is a more sensitive indicator of somatic protein and nutrition status than is percentage of ideal body weight. Creatinine, a product of muscle metabolism, is normally excreted in urine at a constant rate proportional to the amount of skeletal muscle and lean body mass catabolized. CHI is calculated from a 24-hr urinary
creatinine measurement, and the ideal urinary creatinine value found in Table 6–2, using the following formula:

\[
CHI = \frac{\text{Actual Urinary Creatinine}}{\text{Ideal Urinary Creatinine for Height}} \times 100
\]

It is important that the urine sample be an aliquot drawn from a 24-hr collection of urine rather than a random sample.

### Table 6–2. Ideal Urinary Creatinine

<table>
<thead>
<tr>
<th>Height (cm)</th>
<th>Ideal Creatinine (mg/24 hr)</th>
<th>Height (cm)</th>
<th>Ideal Creatinine (mg/24 hr)</th>
</tr>
</thead>
<tbody>
<tr>
<td>157.5</td>
<td>1288</td>
<td>147.3</td>
<td>830</td>
</tr>
<tr>
<td>160.0</td>
<td>1325</td>
<td>149.9</td>
<td>851</td>
</tr>
<tr>
<td>162.6</td>
<td>1359</td>
<td>152.4</td>
<td>875</td>
</tr>
<tr>
<td>165.1</td>
<td>1386</td>
<td>154.9</td>
<td>900</td>
</tr>
<tr>
<td>167.6</td>
<td>1426</td>
<td>157.5</td>
<td>925</td>
</tr>
<tr>
<td>170.2</td>
<td>1467</td>
<td>160.0</td>
<td>949</td>
</tr>
<tr>
<td>172.7</td>
<td>1513</td>
<td>162.6</td>
<td>977</td>
</tr>
<tr>
<td>175.3</td>
<td>1555</td>
<td>165.1</td>
<td>1006</td>
</tr>
<tr>
<td>177.8</td>
<td>1596</td>
<td>167.6</td>
<td>1044</td>
</tr>
<tr>
<td>180.3</td>
<td>1642</td>
<td>170.2</td>
<td>1076</td>
</tr>
<tr>
<td>182.9</td>
<td>1691</td>
<td>172.7</td>
<td>1109</td>
</tr>
<tr>
<td>185.4</td>
<td>1739</td>
<td>175.3</td>
<td>1141</td>
</tr>
<tr>
<td>188.0</td>
<td>1785</td>
<td>177.8</td>
<td>1174</td>
</tr>
<tr>
<td>190.5</td>
<td>1831</td>
<td>180.3</td>
<td>1206</td>
</tr>
<tr>
<td>193.0</td>
<td>1891</td>
<td>182.9</td>
<td>1240</td>
</tr>
</tbody>
</table>

\( ^a \)Creatinine coefficient (men) = 23 mg/kg of ideal body weight.

\( ^b \)Creatinine coefficient (women) = 18 mg/kg of ideal body weight.


There are limitations in using CHI as an indicator of malnutrition. Patients sometimes excrete amounts of creatinine and nitrogen that change with diet, medications, degree of renal function, conditions of illness, or stress. Certain drugs interfere with urine creatinine determinations. (See Drug–Laboratory Test Interferences, page 1070.)

**Anthropometric Measurements.** Anthropometric measurements can be of questionable value because of slow changes over time and interobserver variability. If used, the triceps skinfold (TSF) and mid–upper arm circumference (MUAC)
should be measured on the mid–upper portion of the nondominant arm by trained personnel. Detailed procedures and methods of measurement are available. TSF measurement with calipers is compared with the standards in Table 6–1 to give a reasonable estimate of subcutaneous fat reserves. TSF and MUAC, obtained with a tape measure, can be used to derive the mid upper arm muscle circumference MUAMC by the formula:

\[ \text{MUAMC} = \text{MUAC (cm)} - (0.314 \times \text{TSF [mm]}) \]

**VIScERAL PROTEIN ASSESSMENT PARAMETERS**

The status of visceral protein reflects the patient’s ability to respond to stress by means such as immunocompetence and wound healing. Visceral protein status can be determined by measurements of serum albumin, serum thyroxine–binding prealbumin (also referred to as transthyretin or prealbumin), and serum transferrin. These visceral protein indicators usually decrease after trauma or surgical procedures; however, consistently low levels for at least 1 week might indicate a degree of malnutrition. Serum albumin is unreliable as an assessment parameter in certain patients. Serum albumin can be elevated as a result of dehydration, shock, hemoconcentration, or administration of anabolic hormones or IV albumin. Decreased albumin levels can result from chronic illness, malabsorption, pregnancy, nephrotic syndrome, hepatic insufficiency, protein-losing enteropathy, overhydration, or severe burns.

Prealbumin and transferrin are visceral proteins with a more rapid turnover than albumin; they are effective assessment parameters with half-lives of approximately 2 and 8 days, respectively.

Visceral protein levels and nitrogen balance are expected to decline postoperatively. In a comparison between postoperative prealbumin and transferrin serum levels, the decline in prealbumin was much greater, and changes in transferrin were more closely correlated with changes in nitrogen balance. Transferrin levels can be elevated in patients who are iron deficient, pregnant, or taking estrogens or oral contraceptives. Serum albumin, prealbumin, and transferrin values indicative of different degrees of depletion are given in Table 6–1.

**PERIODIC REASSESSMENT**

An initial assessment can be made before beginning a nutrition support regimen. Periodic reassessment of the patient, using some or all of the previously mentioned parameters, can provide a means of objectively evaluating the efficacy of nutrition support. Additional parameters to consider during this stage of assessment are nitrogen balance and body weight.

**NITROGEN BALANCE**

Nitrogen balance determinations indicate the extent to which exogenous protein is being used and can serve as a method for evaluating the efficacy of nutrition support. Because nitrogen balance data are subject to errors of collection and other variables, they should be used only as a relative index of daily change and not an
absolute measure of depletion or improvement. Nitrogen balance is calculated for a 24-hr period with the following formula:

\[ \text{Nitrogen Balance} = \text{Total Nitrogen In} - \text{Total Nitrogen Out} \]

Urinary urea nitrogen (UUN), although a less sensitive indicator of nitrogen output than total urea nitrogen, is a simpler laboratory procedure and is therefore a more frequently used measurement to estimate nitrogen balance. Nitrogen balance is calculated as follows:

\[ \text{Nitrogen Balance} = \frac{\text{Protein Intake (in g)}}{6.25} - (\text{UUN [in g]} + 4) \]

UUN is usually reported in mg/dL; therefore, to derive the amount in grams for use in the above formula, the value must be multiplied by the total 24-hr volume of urine output. The urine sample sent to the laboratory should be an aliquot drawn from an accurate 24-hr urine collection. The factor 4 is added as an empirical number to account for nonurinary nitrogen such as that excreted in feces, sweat, and other normal losses. Excessive nitrogen losses that cannot be measured, such as nitrogen lost in exudates from severe burns or other fluid losses, render nitrogen balance data less reliable.

Positive nitrogen balance can indicate a retention of nitrogen as newly synthesized body protein tissue and nitrogen retained in body fluids. A positive nitrogen balance of 4–6 g/day is the maximum that should be expected; greater amounts are not considered efficient. Because only synthesized protein is of therapeutic interest, increments in BUN above baseline (in grams) should be subtracted from total nitrogen balance. This calculation is summarized as follows:

\[ \text{Corrected Nitrogen Balance} = \text{Nitrogen Balance} - \text{BUN Increment (g)} \]

To derive the BUN increment above baseline in grams, the total body water volume of the patient must be considered. Body water can be estimated to be 55% of total body weight (0.55 L/kg). A BUN of 10 mg/dL above baseline in a 70-kg patient represents a BUN increment of 3.85 g (70 kg × 0.55 L/kg × 100 mg/L = 3850 mg).

**BODY WEIGHT**

The weight difference between body water and tissue is indistinguishable unless water balance is measured. Body weight gain alone is therefore not a reliable maintenance assessment parameter. It is known, however, that weight gain in excess of 200 g/day is undesirable because patients cannot synthesize lean body tissue at a greater rate. Despite its shortcomings as a monitoring parameter, body weight should be measured throughout the support regimen at the same time each day, and intake and output should be considered in the interpretation of body weight changes.

**NUTRIENT REQUIREMENTS**

The nutrients required for enteral and parenteral nutrition are virtually the same. Either mode of nutrition support must consist of the basic components of a normal diet: water, carbohydrate, fat, protein, electrolytes, vitamins, and trace elements.
CALORIC REQUIREMENTS

Accurate estimation of caloric requirements is essential, particularly for the severely stressed or depleted patient, to avoid problems associated with overfeeding and underfeeding. Requirements can be calculated accurately by indirect calorimetry using instruments that measure respiratory gas exchange. When this is not possible, requirements can be estimated as a multiple of the patient’s basal energy expenditure (BEE). BEE is the amount of energy required to maintain basic metabolic functions in the resting state and can be derived from the Harris-Benedict equations:

\[
\text{BEE (Men): } 66 + (13.8 \times \text{wt in kg}) + (5 \times \text{ht in cm}) - (6.8 \times \text{age in yr})
\]

\[
\text{BEE (Women): } 655 + (9.6 \times \text{wt in kg}) + (1.8 \times \text{ht in cm}) - (4.7 \times \text{age in yr})
\]

Mechanically ventilated nonsurgical patients without stress or sepsis should receive a total caloric intake no greater than the calculated BEE. Trauma and sepsis increase energy and protein requirements, and the nutrition support regimen should be adjusted accordingly. One means of determining the severity of catabolism in stress conditions is by measurement of UUN excreted per 24 hr. Caloric requirements can then be estimated as a multiple of BEE, as shown in Table 6–3.

<table>
<thead>
<tr>
<th>24-HR UUN</th>
<th>DEGREE OF NET CATABOLISM</th>
<th>CALORIC REQUIREMENTS</th>
</tr>
</thead>
<tbody>
<tr>
<td>0–5 g</td>
<td>1° (normal)</td>
<td>1 × BEE</td>
</tr>
<tr>
<td>5–10 g</td>
<td>2° (mild)</td>
<td>1.5 × BEE</td>
</tr>
<tr>
<td>10–15 g</td>
<td>3° (moderate)</td>
<td>1.75 × BEE</td>
</tr>
<tr>
<td>&gt;15 g</td>
<td>4° (severe)</td>
<td>2 × BEE</td>
</tr>
</tbody>
</table>

BEE, basal energy expenditure; UUN, urinary urea nitrogen.

In estimating the calories to be provided by each substrate, yields may be considered as follows: dextrose, 3.4 kcal/g; fat, 9 kcal/g; and protein, 4 kcal/g. Although protein is considered a calorigenic substrate, it is not usually included in estimating caloric goals because the main role of protein is the preservation or synthesis of lean body mass.

PROTEIN REQUIREMENTS

The minimum requirement for protein is about 0.8 g/kg/day of a balanced mixture of amino acids (AAs), and can be as high as 2.5 g/kg/day in severely stressed or traumatized patients. For optimal synthesis of protein, concurrent provision of
nonprotein calories must be sufficient. To calculate the nonprotein calorie-to-nitrogen ratio, assume that the nitrogen content is 1 g/6.25 g of AAs. The optimal ratio of nonprotein calories to nitrogen for efficient nitrogen retention and nitrogen balance is not definite, but differs with the metabolic state of the patient. Nonprotein calorie-to-nitrogen ratios of standard enteral nutrition and PN formulas are typically about 150:1. Lower ratios are indicated for stress or trauma and higher ratios for nonstressed patients and those with impaired protein metabolism.

**ENTERAL NUTRITION**

For physiologic and economic reasons, the enteral route should be used whenever possible, but adequacy of the GI tract must be established before enteral nutrition is provided. The IV route should be strictly reserved for patients who cannot be adequately nourished by the enteral route.

Formulas for enteral nutrition are available for supplemental oral feeding or enteral feeding through different types of tubes. When the oral route is not feasible, transnasal passage of a feeding tube into the stomach (nasogastric) or intestine (nasoduodenal or nasojejunal) is the feeding route generally employed. Feeding ostomies, most commonly the gastrostomy, jejunostomy, or combination gastrostomy–jejunostomy, are generally indicated when insertion through the nares is not feasible or when long-term feeding is anticipated.

**FORMULA SELECTION**

The abundance of products and lack of an ideal system of categorization can cause confusion in selecting the most appropriate enteral formula for a patient. It is not within the scope of this chapter to fully describe criteria for formula selection or provide a complete list of formulas.

Some nutritionally complete, ready-to-use liquid enteral formulas that are suitable for a variety of patients are presented in Table 6–4. Carbohydrate, fat, and protein sources differ with products and can be important criteria for selecting a product. Because patients with abnormal intestinal function are usually lactose intolerant, only lactose-free products are included. Disease-specific formulas, such as those with high content of branched-chain amino acids (BCAAs) for liver disease or essential AAs for renal disease, might be nutritionally incomplete and are not included because of inadequate evidence of their superiority.

**ADMINISTRATION**

One of two types of feeding schedules can be employed, continuous or intermittent. Continuous drip infusion is the preferred method of administration, particularly for patients who have not eaten for a long time. Large 24-hr volumes may be given by infusion without challenging the GI tract, thereby allowing readaptation of the starved gut. Although gravity can be used, an infusion pump is recommended when initiating therapy. For most patients, it is recommended that the first day’s feeding be infused at a rate of 50 mL/hr using a lactose-free, nutrient-intact, isotonic formula of 1 kcal/mL. Many protocols recommend diluting the initial formula to one-half strength; however, this practice has been questioned.11
### TABLE 6–4. REPRESENTATIVE ENTERAL FORMULAS

<table>
<thead>
<tr>
<th>PRODUCT</th>
<th>CALORIES (per mL)</th>
<th>PROTEIN (g/L)</th>
<th>FAT (g/L)</th>
<th>CARBOHYDRATE (g/L)</th>
<th>NONPROTEIN CALORIES (cal/g nitrogen)</th>
<th>SODIUM (mEq/L)</th>
<th>POTASSIUM (mEq/L)</th>
<th>CALCIUM (mg/L)</th>
<th>PHOSPHORUS (mg/L)</th>
<th>OSMOLARITY (mOsm/L)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Compleat Modified</td>
<td>1.07</td>
<td>43</td>
<td>37</td>
<td>140</td>
<td>135</td>
<td>44</td>
<td>36</td>
<td>670</td>
<td>930</td>
<td>300</td>
</tr>
<tr>
<td>Criticare HN</td>
<td>1.06</td>
<td>38</td>
<td>5</td>
<td>220</td>
<td>152</td>
<td>27</td>
<td>34</td>
<td>530</td>
<td>530</td>
<td>650</td>
</tr>
<tr>
<td>Ensure Plus</td>
<td>1.5</td>
<td>54</td>
<td>53</td>
<td>197</td>
<td>146</td>
<td>45</td>
<td>49</td>
<td>705</td>
<td>705</td>
<td>690</td>
</tr>
<tr>
<td>Ensure Plus HN</td>
<td>1.5</td>
<td>62</td>
<td>49</td>
<td>197</td>
<td>125</td>
<td>51</td>
<td>46</td>
<td>1057</td>
<td>1057</td>
<td>650</td>
</tr>
<tr>
<td>Impact</td>
<td>1.0</td>
<td>56</td>
<td>28</td>
<td>130</td>
<td>86</td>
<td>48</td>
<td>33</td>
<td>800</td>
<td>800</td>
<td>375</td>
</tr>
<tr>
<td>Isocal</td>
<td>1.06</td>
<td>34</td>
<td>143</td>
<td>172</td>
<td>23</td>
<td>34</td>
<td>630</td>
<td>530</td>
<td>270</td>
<td>1000</td>
</tr>
<tr>
<td>Isocal HCN</td>
<td>2.0</td>
<td>75</td>
<td>102</td>
<td>200</td>
<td>143</td>
<td>35</td>
<td>43</td>
<td>1000</td>
<td>1000</td>
<td>640</td>
</tr>
<tr>
<td>Isosource HN</td>
<td>1.2</td>
<td>53</td>
<td>41</td>
<td>160</td>
<td>119</td>
<td>48</td>
<td>44</td>
<td>670</td>
<td>670</td>
<td>330</td>
</tr>
<tr>
<td>Jevity</td>
<td>1.06</td>
<td>44</td>
<td>35</td>
<td>151</td>
<td>130</td>
<td>40</td>
<td>40</td>
<td>910</td>
<td>760</td>
<td>300</td>
</tr>
<tr>
<td>Magnacal</td>
<td>2.0</td>
<td>70</td>
<td>80</td>
<td>250</td>
<td>154</td>
<td>43</td>
<td>32</td>
<td>1000</td>
<td>1000</td>
<td>590</td>
</tr>
<tr>
<td>Nitrolan</td>
<td>1.24</td>
<td>60</td>
<td>40</td>
<td>160</td>
<td>104</td>
<td>30</td>
<td>30</td>
<td>800</td>
<td>800</td>
<td>310</td>
</tr>
<tr>
<td>Osmolite</td>
<td>1.06</td>
<td>37</td>
<td>143</td>
<td>153</td>
<td>27</td>
<td>26</td>
<td>528</td>
<td>528</td>
<td>300</td>
<td>300</td>
</tr>
<tr>
<td>Osmolite HN</td>
<td>1.06</td>
<td>44</td>
<td>35</td>
<td>140</td>
<td>124</td>
<td>40</td>
<td>40</td>
<td>758</td>
<td>758</td>
<td>300</td>
</tr>
<tr>
<td>Pulmocare</td>
<td>1.5</td>
<td>62</td>
<td>92</td>
<td>104</td>
<td>125</td>
<td>56</td>
<td>44</td>
<td>1060</td>
<td>1060</td>
<td>475</td>
</tr>
<tr>
<td>Reabilan HN</td>
<td>1.3</td>
<td>58</td>
<td>52</td>
<td>158</td>
<td>119</td>
<td>44</td>
<td>42</td>
<td>500</td>
<td>500</td>
<td>490</td>
</tr>
<tr>
<td>Suplena</td>
<td>2.0</td>
<td>30</td>
<td>92</td>
<td>253</td>
<td>389</td>
<td>34</td>
<td>28</td>
<td>1386</td>
<td>728</td>
<td>600</td>
</tr>
<tr>
<td>Sustacal</td>
<td>1.0</td>
<td>61</td>
<td>23</td>
<td>138</td>
<td>78</td>
<td>40</td>
<td>52</td>
<td>1010</td>
<td>930</td>
<td>620</td>
</tr>
<tr>
<td>Sustacal Plus</td>
<td>1.5</td>
<td>61</td>
<td>53</td>
<td>200</td>
<td>131</td>
<td>55</td>
<td>53</td>
<td>850</td>
<td>850</td>
<td>600</td>
</tr>
<tr>
<td>TraumaCal</td>
<td>1.5</td>
<td>83</td>
<td>69</td>
<td>195</td>
<td>105</td>
<td>52</td>
<td>36</td>
<td>750</td>
<td>750</td>
<td>490</td>
</tr>
<tr>
<td>Ultrical</td>
<td>1.06</td>
<td>44</td>
<td>45</td>
<td>123</td>
<td>127</td>
<td>40</td>
<td>41</td>
<td>850</td>
<td>850</td>
<td>310</td>
</tr>
</tbody>
</table>

Incremental advances in rate and strength can be attempted daily until the desired rate of a full-strength formula is achieved. To minimize the risk of aspiration, proper placement of the tube must be confirmed, and the patient’s head and shoulders must be kept at a 30–45° angle during and for 1 hr after feeding. The stomach should be checked periodically for residual volumes during gastric feedings.

Once a patient has been stabilized on maintenance therapy, intermittent infusions can be used, allowing the patient to rest from feedings at selected hours. A volume of 250–400 mL may be administered 5–8 times/day. This method is preferred for ambulatory patients because it permits more freedom of movement than does continuous feeding.
Formulas should be given at room temperature and kept no longer than 12 hr after the time of preparation and 6 hr from the start of administration to avoid excessive bacterial growth. The delivery system, including bag and tubing, should be changed q 24 hr.

COMPLICATIONS

Mechanical and GI complications known to occur with tube feedings are summarized in Table 6–5. Metabolic complications that occur with enteral nutrition are similar to those with PN and are included in Table 6–12.

To prevent metabolic complications, monitoring of the patient as suggested in Table 6–13 is recommended.

### TABLE 6–5. TUBE FEEDING COMPLICATIONS AND MANAGEMENT

<table>
<thead>
<tr>
<th>COMPLICATION</th>
<th>PREVENTION OR MANAGEMENT</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Mechanical</strong></td>
<td></td>
</tr>
<tr>
<td>Clogged Tube</td>
<td>Flush with water, replace tube if necessary. Avoid passing crushed tablets through small-bore feeding tubes.</td>
</tr>
<tr>
<td>Nasal, Pharyngeal,</td>
<td>Use small-lumen flexible tube. Provide daily care of nose and mouth.</td>
</tr>
<tr>
<td>Esophageal Irritation</td>
<td>Ensure proper tube placement and verify location. Maintain patient’s head and shoulders at 30–45° in the upright position during and for 1 hr after feeding. Monitor for gastric reflux and abdominal distention. Stop infusion if vomiting occurs. Check residual gastric volume before and q 2–4 hr during infusion. Hold if the residual exceeds the hourly volume or 150 mL.</td>
</tr>
<tr>
<td>Aspiration</td>
<td>Verify tube location and mark tube at insertion site.</td>
</tr>
<tr>
<td>Dislocated Tube</td>
<td></td>
</tr>
<tr>
<td><strong>Gastrointestinal</strong></td>
<td></td>
</tr>
<tr>
<td>Diarrhea and Cramps</td>
<td>Reduce flow rate, dilute formula, or consider alternative formula. Rule out alternative causes. If persistent, add antidiarrheal agent.</td>
</tr>
<tr>
<td>Vomiting or Bloating</td>
<td>Check stool output and measure residual formula in gut q 2–4 hr if necessary, stop or reduce flow.</td>
</tr>
<tr>
<td>Constipation</td>
<td>Consider different formula or a laxative.</td>
</tr>
</tbody>
</table>

■ PARENTERAL NUTRITION

PN may be administered by one of two routes of access: peripheral veins or larger central veins. The peripheral route is indicated for those patients who require only short-term supplementation or supplementation in addition to enteral support or for those in whom the risks of central venous administration are too great. Peripheral veins are susceptible to thrombophlebitis, particularly when the osmolarity of the solution exceeds 600 mOsm/L. Therefore, it is recommended that formulas for peripheral administration not exceed final concentrations of 10% dextrose and
4.5% AAs plus electrolyte and vitamin additives. Many techniques to prevent or delay onset of peripheral vein thrombophlebitis have been reported. Addition of small amounts of hydrocortisone (5 mg/L) and heparin (1000 units/L) to PN formulas, as well as the topical use of agents such as nitroglycerin, have demonstrated success. Concurrent administration of IV fat emulsion, which is a concentrated, iso-osmotic calorie source, is vital because it increases the caloric content of a peripheral regimen and minimizes the risk of thrombophlebitis.

The complete nutrition needs of the malnourished or hypermetabolic patient are difficult to provide through peripheral vein for long periods. The concentrated, hyperosmolar solutions required by such patients for PN must be administered into a large central vein, such as the superior vena cava, where rapid dilution occurs.

**ADMINISTRATION**

Initiation of PN should be gradual, particularly in the malnourished patient, to avoid glucose intolerance and the dangers of refeeding syndrome. With high concentrations of dextrose and AAs intended for central vein administration, an initial rate of 40 mL/hr for the first 24 hr is suggested. Infusion rates may then be increased daily in accordance with assessment goals. Less concentrated formulas that are suited for peripheral vein administration do not warrant such slow initial rates of infusion.

Different catheters exist for infusion of PN formulas by central or peripheral vein. Use of an in-line filter is recommended to minimize adverse consequences in case precipitation occurs in the PN solution.

**PARENTERAL NUTRIENTS**

Each of the following nutrient substrate groups are required in formulas for effective PN.

**Water.** The average healthy adult can tolerate a fluid infusion volume of about 5 L/day. The patient who is fluid restricted might be limited to an intake of ≤ 2 L/day. This might be the deciding factor in selecting a hypertonic concentrated solution for infusion through a large central vein rather than a more dilute solution for peripheral administration.

**Carbohydrate.** The presently preferred carbohydrate substrate for PN is dextrose. The concentration of dextrose should be determined by the osmotic limitation of the administration route and the nonprotein calorie requirement of the patient. The concentrations of available dextrose solutions with their corresponding caloric concentrations and osmolarities are shown in Table 6–6. Dextrose remains the primary source of calories for PN through central vein, and the rate of infusion should be limited to its maximum rate of oxidation, which is 5 mg/kg/min or 7.2 g/kg/day. On a calorie-for-calorie basis, carbohydrate is more efficient than fat in sparing body protein during hypocaloric feedings. The inclusion of dextrose and fat is recommended in PN regimens, but the optimal proportion of each has not been established.

**Fat.** Fat is an important parenteral substrate for three major reasons: (1) it is a concentrated source of calories in an isotonic medium, which makes it useful for
peripheral administration; (2) it is a source of essential fatty acids (EFAs) required for prevention or treatment of EFA deficiency, which can develop during prolonged fat-free PN; and (3) it is a useful substitute for carbohydrate when dextrose calories must be limited because of glucose intolerance or diminished ventilatory capacity. When a patient’s ventilatory effort is hampered, it is important to avoid excessive calories of any type. In comparison with dextrose, the metabolism of fat increases heat production, decreases respiratory quotient (RQ), and increases oxygen consumption. Because it has a lower RQ, fat produces less CO₂ for a given number of calories, thereby minimizing the ventilatory effort required to eliminate CO₂. The RQ of fat is 0.7 compared with 1 for carbohydrate. An RQ in excess of 1 indicates net lipogenesis and is undesirable.

Fat is available as emulsions of 10, 20, or 30% soybean oil or 10 or 20% soybean–safflower oil mixtures. Clinical studies have not shown any major advantages of one lipid source over the other. The major differences between these products are their fatty acid contents, which are summarized in Table 6-7. The 20 and 30% emulsions are more readily cleared than the 10% one because of the lower proportion of phospholipid to triglyceride.

Fat emulsions that are currently marketed in the United States contain only long-chain triglycerides (LCTs); however, the use of fat emulsions that contain LCTs and medium-chain triglycerides (MCTs) is being investigated. MCTs are reported to be more rapidly cleared from the blood and more ketogenic than

### Table 6-6. IV Dextrose Solutions

<table>
<thead>
<tr>
<th>Concentration</th>
<th>kcal/L</th>
<th>mOsm/L</th>
</tr>
</thead>
<tbody>
<tr>
<td>5%</td>
<td>170</td>
<td>252</td>
</tr>
<tr>
<td>10%</td>
<td>340</td>
<td>505</td>
</tr>
<tr>
<td>20%</td>
<td>680</td>
<td>1010</td>
</tr>
<tr>
<td>40%</td>
<td>1360</td>
<td>2020</td>
</tr>
<tr>
<td>50%</td>
<td>1700</td>
<td>2520</td>
</tr>
<tr>
<td>60%</td>
<td>2040</td>
<td>3030</td>
</tr>
<tr>
<td>70%</td>
<td>2380</td>
<td>3530</td>
</tr>
</tbody>
</table>

### Table 6-7. IV Fat Emulsions

<table>
<thead>
<tr>
<th>Fatty Acid</th>
<th>Soybean Oil</th>
<th>Soybean Oil/Safflower Oil</th>
</tr>
</thead>
<tbody>
<tr>
<td>Linoleic Acid</td>
<td>54%</td>
<td>65.8%</td>
</tr>
<tr>
<td>Linolenic Acid</td>
<td>8%</td>
<td>4.2%</td>
</tr>
<tr>
<td>Oleic Acid</td>
<td>26%</td>
<td>17.7%</td>
</tr>
<tr>
<td>Palmitic Acid</td>
<td>9%</td>
<td>8.8%</td>
</tr>
<tr>
<td>Stearic Acid</td>
<td>2.5%</td>
<td>3.4%</td>
</tr>
</tbody>
</table>
LCTs, and emulsions containing MCTs and LCTs have greater protein-conserving properties than pure LCT emulsions. LCTs are required for their EFA content, however.

The caloric density of 10% fat emulsions is 1.1 kcal/mL, of which 1 kcal is supplied by lipid and 0.1 kcal by glycerol (carbohydrate); the 20% and 30% emulsions have caloric densities of 2 and 3 kcal/mL, respectively, of which 0.1 kcal/mL is glycerol. The average particle size (0.5 µ) is the same in all concentrations, and all are nearly iso-osmotic.

Fat emulsion can be infused concurrently with AA/dextrose solution through peripheral or central veins. The 10% or 20% emulsion may be infused separately or combined with AAs and dextrose in a single container to form a total nutrient (“3-in-1”) admixture (TNA). The 30% concentration is intended only for compounding TNA. Because lipid emulsion is iso-osmolar, it reduces the thrombophlebitic effect of hyperosmolar AA/dextrose solutions on the endothelium of peripheral veins when they are infused concurrently. For this reason and its potential adverse effect on the immune system, fat emulsion should be infused as slowly as possible. For further information on dosage, administration, and precautions of fat emulsion, the product literature should be consulted.

Protein. Various brands and concentrations of AA solutions are available as sources of protein for parenteral use. The AA profile differs in each; therefore, their nitrogen contents are not equivalent. A comparison of formulations is summarized in Table 6–8. AA solutions >3.5% concentration should be diluted to a lower final concentration with dextrose and other additives.

SPECIAL AMINO ACID SOLUTIONS
Special AA solutions are available for specific metabolic or disease states. Discretion is recommended in the use of these solutions because they are expensive and clinical benefit is not proved.

Protein Sparing. A low concentration of AAs infused with or without concurrent nonprotein calories conserves endogenous nitrogen more efficiently than the traditional 5% dextrose infusion alone. For a limited infusion of no more than 1 week’s duration in patients who are not severely catabolic, low-concentration AA formulas merit consideration. Low-concentration AA formulas are available with or without electrolytes and with or without a nonprotein calorie source (see Table 6–8).

Renal Failure. The objective of PN in patients with renal failure is to provide sufficient AAs and calories for protein synthesis without exceeding the renal capacity for excretion of fluid and metabolic wastes. Four parenteral products that contain primarily essential AAs have been developed for this purpose (see Table 6–8), but controversy exists regarding their use. Patients who undergo renal re-
TABLE 6–8. AMINO ACID SOLUTIONS COMPARISON CHART

<table>
<thead>
<tr>
<th>AA SOLUTION AND OSMOLARITY CONCENTRATION</th>
<th>TOTAL BCAAs (g/dL)</th>
<th>TOTAL ESSENTIAL AAs (g/dL)</th>
<th>TOTAL N (g/dL)</th>
<th>ELECTROLYTES (mEq/L)</th>
<th>OSMOLARITY (mOsm/L)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Na⁺</td>
<td>K⁺</td>
<td>Mg⁺⁺</td>
<td>Cl⁻</td>
<td>Ac⁻</td>
</tr>
<tr>
<td>FOR GENERAL PURPOSE</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Aminosyn 3.5%</td>
<td>0.86</td>
<td>1.65</td>
<td>0.55</td>
<td>7</td>
<td>—</td>
</tr>
<tr>
<td>Aminosyn 5%</td>
<td>1.23</td>
<td>2.35</td>
<td>0.79</td>
<td>—</td>
<td>5.4</td>
</tr>
<tr>
<td>Travasol 5.5% (with electrolytes)</td>
<td>0.86</td>
<td>2.15</td>
<td>0.93</td>
<td>70</td>
<td>60</td>
</tr>
<tr>
<td>Aminosyn 7% (with electrolytes)</td>
<td>1.73</td>
<td>3.32</td>
<td>1.1</td>
<td>—</td>
<td>5.4</td>
</tr>
<tr>
<td>Aminosyn 8.5% (with electrolytes)</td>
<td>2.11</td>
<td>4.06</td>
<td>1.34</td>
<td>76</td>
<td>66</td>
</tr>
<tr>
<td>Travasol 8.5% (with electrolytes)</td>
<td>1.32</td>
<td>3.34</td>
<td>1.43</td>
<td>—</td>
<td>—</td>
</tr>
<tr>
<td>FreAmine III 8.5% (with electrolytes)</td>
<td>1.92</td>
<td>3.94</td>
<td>1.43</td>
<td>70</td>
<td>60</td>
</tr>
<tr>
<td>Aminosyn 10%</td>
<td>2.46</td>
<td>4.7</td>
<td>1.57</td>
<td>—</td>
<td>5.4</td>
</tr>
<tr>
<td>Aminosyn II 10%</td>
<td>2.16</td>
<td>4.3</td>
<td>1.53</td>
<td>45</td>
<td>—</td>
</tr>
<tr>
<td>FreAmine III 10%</td>
<td>2.26</td>
<td>4.63</td>
<td>1.53</td>
<td>10</td>
<td>—</td>
</tr>
<tr>
<td>Travasol 10%</td>
<td>1.91</td>
<td>4.05</td>
<td>1.65</td>
<td>—</td>
<td>—</td>
</tr>
<tr>
<td>Novamine</td>
<td>2.09</td>
<td>5.11</td>
<td>1.8</td>
<td>—</td>
<td>—</td>
</tr>
<tr>
<td>AA SOLUTION AND OSMOLARITY CONCENTRATION</td>
<td>TOTAL BCAAs (g/dL)</td>
<td>TOTAL ESSENTIAL AAs (g/dL)</td>
<td>TOTAL N (g/dL)</td>
<td>ELECTROLYTES (mEq/L)</td>
<td>PO₄ (mmol/L)</td>
</tr>
<tr>
<td>------------------------------------------</td>
<td>------------------</td>
<td>-----------------------------</td>
<td>----------------</td>
<td>----------------------</td>
<td>-------------</td>
</tr>
<tr>
<td>Aminosyn II 15%</td>
<td>3.24</td>
<td>6.42</td>
<td>2.3</td>
<td>63</td>
<td>—</td>
</tr>
<tr>
<td>Novamine 15%</td>
<td>2.75</td>
<td>6.72</td>
<td>2.37</td>
<td>—</td>
<td>—</td>
</tr>
<tr>
<td><strong>FOR PROTEIN SPARING</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>ProcalAmine 3%b</td>
<td>0.68</td>
<td>1.4</td>
<td>0.46</td>
<td>35</td>
<td>24</td>
</tr>
<tr>
<td>FreAmine III 3% (with electrolytes)</td>
<td>0.68</td>
<td>1.4</td>
<td>0.46</td>
<td>35</td>
<td>24.5</td>
</tr>
<tr>
<td>Aminosyn 3.5% M&lt;sup&gt;a&lt;/sup&gt;</td>
<td>0.86</td>
<td>1.65</td>
<td>0.55</td>
<td>47</td>
<td>13</td>
</tr>
<tr>
<td>3.5% Travaasol (with electrolytes)</td>
<td>0.55</td>
<td>1.38</td>
<td>0.59</td>
<td>25</td>
<td>15</td>
</tr>
<tr>
<td><strong>FOR RENAL FAILURE</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Aminess 5.2%</td>
<td>1.95</td>
<td>5.18</td>
<td>0.66</td>
<td>—</td>
<td>—</td>
</tr>
<tr>
<td>Aminosyn RF 5.2%</td>
<td>1.72</td>
<td>4.83</td>
<td>0.79</td>
<td>—</td>
<td>5.4</td>
</tr>
<tr>
<td>NephrAmine 5.4%</td>
<td>2.08</td>
<td>5.33</td>
<td>0.65</td>
<td>5</td>
<td>—</td>
</tr>
<tr>
<td>RenAmin 6.5%</td>
<td>1.92</td>
<td>4.32</td>
<td>1.0</td>
<td>—</td>
<td>—</td>
</tr>
<tr>
<td><strong>FOR TRAUMA</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>BranchAmin 4%c,d&lt;sup&gt;c&lt;/sup&gt;</td>
<td>4.0</td>
<td>4.0&lt;sup&gt;d&lt;/sup&gt;</td>
<td>0.44</td>
<td>—</td>
<td>—</td>
</tr>
<tr>
<td>FreAmine HBC 6.9%c&lt;sup&gt;c&lt;/sup&gt;</td>
<td>3.01</td>
<td>4.28</td>
<td>0.97</td>
<td>10</td>
<td>—</td>
</tr>
<tr>
<td>Aminosyn HBC 7%c&lt;sup&gt;c&lt;/sup&gt;</td>
<td>3.15</td>
<td>4.21</td>
<td>1.12</td>
<td>7</td>
<td>40</td>
</tr>
</tbody>
</table>

(continued)
<table>
<thead>
<tr>
<th>AA SOLUTION AND OSMOLARITY CONCENTRATION</th>
<th>TOTAL BCAAs (g/dL)</th>
<th>TOTAL ESSENTIAL AAs (g/dL)</th>
<th>TOTAL N (g/dL)</th>
<th>ELECTROLYTES (mEq/L)</th>
<th>OSMOLARITY (mOsm/L)</th>
</tr>
</thead>
<tbody>
<tr>
<td>FOR LIVER DISEASE</td>
<td></td>
<td></td>
<td></td>
<td>Na⁺</td>
<td>K⁺</td>
</tr>
<tr>
<td>HepatAmine</td>
<td>2.84</td>
<td>4.17</td>
<td>1.2</td>
<td>10</td>
<td>—</td>
</tr>
<tr>
<td>FOR PEDIATRICS</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Aminosyn-PF 7%</td>
<td>1.82</td>
<td>3.2</td>
<td>1.07</td>
<td>3.4</td>
<td>—</td>
</tr>
<tr>
<td>Aminosyn-PF 10%</td>
<td>2.63</td>
<td>4.61</td>
<td>1.52</td>
<td>3.4</td>
<td>—</td>
</tr>
<tr>
<td>TrophAmine 6%</td>
<td>1.8</td>
<td>4.28</td>
<td>0.93</td>
<td>5</td>
<td>—</td>
</tr>
<tr>
<td>TrophAmine 10%</td>
<td>3.0</td>
<td>7.2</td>
<td>1.55</td>
<td>5</td>
<td>—</td>
</tr>
</tbody>
</table>

*Also available as Aminosyn II which contains glutamic and aspartic acids, and differs slightly in content of other amino acids, acetate, and chloride.

*Contains glycerol as a nonprotein calorie source.

*BCAA–enriched products. Each of these products has distinct indications for use and should not be interchanged.

*Contains only BCAA. Other essential AA are not included.

AA, amino acid; BCAA, branched-chain amino acid.
placement therapy such as peritoneal or hemodialysis require essential and nonessential AAs and should receive standard AA solutions.

**Hepatic Failure.** Patients with hepatic failure, in whom muscle breakdown and an altered serum and CNS AA profile might contribute to hepatic encephalopathy, can benefit from a special AA formula. This formula has relatively greater amounts of BCAAs (i.e., leucine, isoleucine, valine) and smaller amounts of the aromatic acids (i.e., phenylalanine, tyrosine, tryptophan) and methionine. One parenteral formula, HepatAmine, is currently available specifically for therapeutic and nutrition support of patients with liver disease (see Table 6–8).

**Stress and Trauma.** The hypermetabolism that occurs in response to stress and trauma presents difficulty in providing nutrition support. BCAAs, in addition to their useful effect in metabolic support of the patient with liver disease, are reported to be useful for patients with stress and trauma. Three BCAA-enriched products are available (see Table 6–8). FreAmine HBC and Aminosyn HBC are solutions of nonessential and essential AAs enriched with BCAAs. BranchAmin 4% is a solution of only BCAAs intended for use as a supplement to be admixed with a complete AA and a nonprotein caloric source. These products are indicated only for stress and trauma and should not be confused with the BCAA-enriched product that is indicated for hepatic encephalopathy.

**Pediatrics.** It is beyond the scope of this chapter to describe procedures for nutrition support of pediatric patients except for this brief mention of parenteral AA products. Crystalline AA solutions marketed for infants are based on the essentiality of certain AAs in these patients (see Table 6–8). Compared with adult AA formulations, these products contain taurine and glutamic and aspartic acids. Increased amounts of tyrosine and histidine and lower amounts of phenylalanine, methionine, and glycine are included. Although cysteine is also assumed to be essential for infants, adequate amounts cannot be included in AA formulas because of its limited solubility. A cysteine solution (50 mg/mL) is available separately for admixture to the formula before administration.

### ELECTROLYTES

Formulas also are available with standard electrolyte compositions that might be suitable for most patients, after the addition of certain additives. Electrolyte provision, however, should be based on close monitoring of patients’ laboratory values. Average daily requirements are summarized in Table 6–9.

### VITAMINS

Vitamin requirements for PN have been suggested in a report by an advisory group to the American Medical Association (AMA). Multiple vitamins are available in adult and pediatric formulations for once-daily IV administration (see Table 6–10). The usual daily dosage of the adult formulation is 10 mL to provide the amounts of vitamins specified in Table 6–10. The daily dosage of the pediatric formulation for infants who weigh <1 kg is 1.5 mL. For infants weighing 1–3 kg, the daily dosage is 3 mL. For infants and children weighing ≥3 kg up to 11 yr of...
# TABLE 6–9. ELECTROLYTES AND REQUIREMENTS

<table>
<thead>
<tr>
<th>ELECTROLYTES</th>
<th>AVERAGE DAILY REQUIREMENT</th>
<th>DOSAGE FORMS</th>
<th>COMMENTS</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>CATIONS</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Sodium</td>
<td>60–150 mEq</td>
<td>Sodium chloride concentrate (4 mEq/mL)</td>
<td>Requirements during parenteral nutrition should not differ from normal fluid therapy requirements unless there is excessive sodium loss. Lactate and bicarbonate salts of sodium should not be used.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Sodium acetate (2 mEq/mL)</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>Sodium phosphate (4 mEq Na⁺/mL)</td>
<td></td>
</tr>
<tr>
<td>Potassium</td>
<td>40–240 mEq</td>
<td>Potassium chloride (2 mEq/mL)</td>
<td>Requirements are related to glucose metabolism and therefore increase with higher concentrations of dextrose infused.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Potassium acetate (2 mEq/mL)</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>Potassium phosphate (4.4 mEq K⁺/mL)</td>
<td></td>
</tr>
<tr>
<td>Magnesium</td>
<td>10–45 mEq</td>
<td>Magnesium sulfate (4 mEq/mL)</td>
<td>Requirements increase with anabolism but with less variation than with potassium.</td>
</tr>
<tr>
<td>Calcium</td>
<td>5–30 mEq</td>
<td>Calcium gluconate 10% (4.5 mEq/10 mL)</td>
<td>Requirements increase only slightly during parenteral nutrition. Limited amounts of calcium and phosphate, as determined by compatibility references, may be combined in solutions that contain amino acids.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Calcium chloride 10% (13 mEq/10 mL)</td>
<td></td>
</tr>
<tr>
<td><strong>ANIONS</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Phosphate</td>
<td>10 mmol/1000 kcal</td>
<td>Potassium phosphate (3 mmol P/mL, Abbott)</td>
<td>Requirements increase with anabolism. Safe empirical dosage guidelines should be developed, taking into account the sodium or potassium content of the phosphate solution.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Sodium phosphate (3 mmol P/mL, Abbott) (other concentrations may vary according to manufacturer)</td>
<td></td>
</tr>
<tr>
<td>Acetate and Chloride</td>
<td>The amounts of acetate and chloride contained in each amino acid solution vary. (See Table 6–8.) Acetate is metabolized to bicarbonate.</td>
<td>Bicarbonate salts should not be added to PN solutions because of incompatibility.</td>
<td></td>
</tr>
</tbody>
</table>
age, the daily dosage is 5 mL. Vitamin K is included in the pediatric product only. Phytonadione 5 mg may be given to adults weekly in the PN formula, or by IM or SC administration, if needed.30

Fat emulsion contains vitamin K. Intralipid 10% contains about 0.31 mg/L and Liposyn II contains 0.13 mg/L; 20% products contain twice as much. Intralipid 20% 500 mL provides about 300 μg of vitamin K, an amount that exceeds maintenance recommendations and interferes with oral anticoagulant therapy.30

### TABLE 6–10. IV MULTIVITAMINS

<table>
<thead>
<tr>
<th>AMOUNT</th>
<th>TYPICAL FORMULA</th>
<th>ADULT (PER VIAL)</th>
<th>PEDIATRIC (5 mL)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ascorbic Acid (C)</td>
<td>100 mg</td>
<td>80 mg</td>
<td></td>
</tr>
<tr>
<td>Vitamin A</td>
<td>3300 IU</td>
<td>2300 IU</td>
<td></td>
</tr>
<tr>
<td>Vitamin D</td>
<td>200 IU</td>
<td>400 IU</td>
<td></td>
</tr>
<tr>
<td>Vitamin E</td>
<td>10 IU</td>
<td>7 IU</td>
<td></td>
</tr>
<tr>
<td>Thiamine (B₁)</td>
<td>3 mg</td>
<td>1.2 mg</td>
<td></td>
</tr>
<tr>
<td>Riboflavin (B₂)</td>
<td>3.6 mg</td>
<td>1.4 mg</td>
<td></td>
</tr>
<tr>
<td>Niacinamide (B₃)</td>
<td>40 mg</td>
<td>17 mg</td>
<td></td>
</tr>
<tr>
<td>Pantothenic Acid (B₅)</td>
<td>15 mg</td>
<td>5 mg</td>
<td></td>
</tr>
<tr>
<td>Pyridoxine (B₆)</td>
<td>4 mg</td>
<td>1 mg</td>
<td></td>
</tr>
<tr>
<td>Biotin</td>
<td>60 μg</td>
<td>20 μg</td>
<td></td>
</tr>
<tr>
<td>Folic Acid</td>
<td>400 μg</td>
<td>140 μg</td>
<td></td>
</tr>
<tr>
<td>Cyanocobalamin (B₁₂)</td>
<td>5 μg</td>
<td>1 μg</td>
<td></td>
</tr>
<tr>
<td>Phytonadione (K₁)</td>
<td>0</td>
<td>200 μg</td>
<td></td>
</tr>
</tbody>
</table>

### TRACE ELEMENTS

Solutions of individual trace elements are available in several concentrations from different manufacturers. Solutions of multiple trace elements also are commercially available in products containing 4, 5, 6, or 7 elements and in concentrations suitable for adult or pediatric use. Guidelines for the use of trace elements in PN have been reported in an AMA statement31 and the recommended daily dosages appear in Table 6–11. Although a need for molybdenum and iodine in long-term PN has been described, there are no officially recommended requirements for these elements.32–34

### IRON

Iron deficiency can occur in patients deprived of iron during long-term PN. Iron dextran is sometimes added to PN solutions, but the advisability of its routine use
and its compatibility with fat emulsion are questionable. Dosage recommendations by this route are 1–12.5 mg/day of iron.36

INSULIN

Many patients who receive PN become hyperglycemic. When feasible, the cause should be investigated and controlled by means other than insulin before insulin is employed (see Table 6–12). Although the efficacy of PN is reportedly enhanced by insulin,37 it should be used cautiously to avoid hypoglycemia and because it promotes deposition of fatty acids in body fat stores, making them less available for important biochemical pathways.38 When it is required, insulin may be provided separately by SC or IV administration or added to the PN formula. Until a patient is stabilized on a consistent dosage of insulin, it is more cost effective to provide insulin separately to avoid wasting of PN formulations that might be discarded if the insulin dosage needs to be changed.39 Human insulin is the least immunogenic and is therefore the insulin of choice. Guidelines for dosage are empirical; one-half to two-thirds of the previous day’s sliding scale requirements may be added as regular human insulin to the daily PN formula. Standardized admixture procedures should be used to minimize variations of insulin activity caused by adsorption loss.

<table>
<thead>
<tr>
<th>TRACE ELEMENT</th>
<th>PEDIATRIC PATIENTS (µG/kg)a</th>
<th>STABLE ADULT</th>
<th>ADULT IN ACUTE CATABOLIC STATEb</th>
<th>STABLE ADULT WITH INTESTINAL LOSSESb</th>
</tr>
</thead>
<tbody>
<tr>
<td>Zinc</td>
<td>400 (preterm)c</td>
<td>2.5–4 mg</td>
<td>Additional 2 mg</td>
<td>Add 12.2 mg/L of small-bowel fluid lost; 17.1 mg/kg of stool or ileostomy output.e</td>
</tr>
<tr>
<td></td>
<td>250 (&lt;3 months)d</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>100 (&gt;3 months–1 yr)d</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>50 (&gt;1 yr)c</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Copper</td>
<td>20</td>
<td>0.5–1.5 mg</td>
<td>—</td>
<td>—</td>
</tr>
<tr>
<td>Chromium</td>
<td>0.14–0.2</td>
<td>10–15 µg</td>
<td>—</td>
<td>20 µgj</td>
</tr>
<tr>
<td>Manganese</td>
<td>1</td>
<td>0.15–0.8 mg</td>
<td>—</td>
<td>—</td>
</tr>
<tr>
<td>Selenium</td>
<td>2</td>
<td>20–60 µg</td>
<td>—</td>
<td>—</td>
</tr>
</tbody>
</table>

*aLimited data are available for infants weighing <1500 g. Their requirements might be more than the recommendations because of their low body reserves and increased requirements for growth.
*bFrequent monitoring of plasma levels in these patients is essential to provide proper dosage.
*cPremature infants (weight <1500 g) up to 3 kg of body weight. Thereafter, the recommendations for full-term infants apply.
*dFull-term infants and children ≤5 yr old. Thereafter, the recommendations for adults apply, up to a maximum dosage of 4 mg/day.
*eValues derived by mathematical fitting of balance data from a 71 patient-week study in 24 patients.
*fMean from balance study.

Modified from references 31 and 35.
<table>
<thead>
<tr>
<th>COMPLICATION</th>
<th>FREQUENT CAUSES</th>
<th>MANAGEMENT</th>
</tr>
</thead>
<tbody>
<tr>
<td>Hyponatremia</td>
<td>Excessive GI or urinary sodium losses, or inadequate sodium intake.</td>
<td>Increase sodium provision.</td>
</tr>
<tr>
<td></td>
<td>Excessive water intake.</td>
<td>Limit free water.</td>
</tr>
<tr>
<td>Hypokalemia</td>
<td>Excessive GI or urinary potassium losses; deficit of potassium; or</td>
<td>Increase potassium provision.</td>
</tr>
<tr>
<td></td>
<td>large glucose infusion.</td>
<td></td>
</tr>
<tr>
<td>Hypocalcemia</td>
<td>Insufficient calcium.</td>
<td>Increase calcium provision.</td>
</tr>
<tr>
<td></td>
<td>Magnesium deficit.</td>
<td>Increase magnesium provision.</td>
</tr>
<tr>
<td>Hypomagnesemia</td>
<td>Insufficient magnesium; or excess GI or urinary losses.</td>
<td>Increase magnesium provision.</td>
</tr>
<tr>
<td>Hypophosphatemia</td>
<td>Inadequate phosphate.</td>
<td>Increase phosphate provision.</td>
</tr>
<tr>
<td></td>
<td>Refeeding syndrome.</td>
<td>Refeed gradually.</td>
</tr>
<tr>
<td>Hypoglycemia</td>
<td>Abrupt interruption of formula infusion.</td>
<td>Begin dextrose infusion and monitor blood</td>
</tr>
<tr>
<td></td>
<td></td>
<td>glucose and potassium.</td>
</tr>
<tr>
<td></td>
<td>Excessive insulin.</td>
<td>Decrease insulin.</td>
</tr>
<tr>
<td>Hyperglycemia</td>
<td>Deficit of potassium or phosphorus.</td>
<td>Increase potassium or phosphate provision.</td>
</tr>
<tr>
<td></td>
<td>Insufficient insulin.</td>
<td>Give insulin.</td>
</tr>
<tr>
<td></td>
<td>Corticosteroid use.</td>
<td>Reduce rate of glucose infusion.</td>
</tr>
<tr>
<td>Hypertriglyceridemia</td>
<td>Impaired clearance.</td>
<td>Hold IV lipid if serum triglycerides</td>
</tr>
<tr>
<td></td>
<td></td>
<td>&gt;400 mg/dL (4.5 mmol/L).</td>
</tr>
<tr>
<td>Elevated BUN</td>
<td>Dehydration.</td>
<td>Correct dehydration.</td>
</tr>
<tr>
<td></td>
<td>Renal dysfunction; or calorie: nitrogen ratio imbalance.</td>
<td>Increase nonprotein calorie:nitrogen ratio.</td>
</tr>
<tr>
<td>Elevated Liver</td>
<td>Underlying disease; lack of GI use; or GI bacterial overgrowth.</td>
<td>Attempt enteral feeding.</td>
</tr>
<tr>
<td>Function Tests</td>
<td>Essential fatty acid deficiency.</td>
<td>Provide lipid.</td>
</tr>
<tr>
<td></td>
<td>Excessive nutrients.</td>
<td>Decrease PN.</td>
</tr>
<tr>
<td>Metabolic Acidosis</td>
<td>Excessive GI or urinary losses of base.</td>
<td>Increase acetate provision.</td>
</tr>
<tr>
<td></td>
<td>Inadequate amount of base-producing substance in formula.</td>
<td>Decrease chloride in formula or increase</td>
</tr>
<tr>
<td></td>
<td></td>
<td>acetate provision.</td>
</tr>
<tr>
<td>Osmotic Diuresis</td>
<td>Failure to recognize initial hyperglycemia and increased glucose in urine.</td>
<td>Reduce infusion rate.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Give insulin to correct hyperglycemia.</td>
</tr>
</tbody>
</table>

(continued)
ALBUMIN
Albumin is compatible when admixed with PN formulas; however, its supply is too limited and its cost is too prohibitive for casual use. Although inclusion of albumin in PN is reported to rapidly increase serum albumin levels and enhance tolerance of enteral feedings, the clinical benefits of such treatment are not proved. For synthesis of endogenous protein, albumin is inferior to crystalline AAs as a parenteral source of nitrogen. If administration of albumin is necessary, it should not be included in the PN formula.

CARNITINE
Carnitine is a micronutrient that is vital to energy metabolism because of its role in transporting long-chain fatty acids across the mitochondrial membrane. Certain patients, such as those with chronic renal failure on dialysis and premature neonates, are at increased risk of developing carnitine deficiency, especially if they are receiving long-term PN. L-carnitine, the physiologically active form, is available for IV administration as a 1 g/5 mL solution that is stable when added to PN formulas. Consult the carnitine product information for detailed usage information.

MEDICATIONS
There may be advantages to the admixture of certain medications such as antibiotics, chemotherapeutic agents, and H2-receptor antagonists to PN, if there is compatibility reported with all components of the formula. Consult other sources for information regarding the stability and compatibility of medication/PN admixtures.

MONITORING THE PATIENT
Metabolic complications known to occur with enteral or parenteral nutrition are summarized in Table 6–12. Most of these can be avoided by proper precautions...
and close monitoring of the patient. Laboratory parameters for patient monitoring are summarized in Table 6–13.

<table>
<thead>
<tr>
<th>PARAMETER</th>
<th>FREQUENCYa</th>
</tr>
</thead>
<tbody>
<tr>
<td>Urinary glucose and specific gravity.</td>
<td>Every voided specimen until stable, then daily.</td>
</tr>
<tr>
<td>Finger stick glucose.</td>
<td>Every 6 hr until stable.</td>
</tr>
<tr>
<td>Vital signs, weight, intake, and output.</td>
<td>Daily.</td>
</tr>
<tr>
<td>Serum glucose, electrolytes, creatinine, and BUN</td>
<td>Daily until stable, then twice weekly.</td>
</tr>
<tr>
<td>Magnesium, calcium, and phosphorus.</td>
<td>Daily until stable, then once weekly.</td>
</tr>
<tr>
<td>CBC, hemoglobin, WBC, platelets, and prothrombin time.</td>
<td>Baseline, then weekly.</td>
</tr>
<tr>
<td>Serum protein, albumin, prealbumin, and liver functions.</td>
<td>Baseline, then weekly.</td>
</tr>
<tr>
<td>Serum cholesterol and triglycerides.</td>
<td>Baseline, then weekly.</td>
</tr>
<tr>
<td>Blood ammonia.</td>
<td>Baseline, then weekly in renal and hepatic patients.</td>
</tr>
</tbody>
</table>

*aFrequency should be increased in critically ill patients.

**FUTURE DEVELOPMENTS**

Technologic advancements in nutrition formulas and the means of preparing, providing, and monitoring their effects on patients continue to be made. These modifications enable safer and more cost-effective nutrition support of patients in the hospital or at home.

Body composition research is presenting innovative approaches to metabolic and nutrition assessments. Formulas with specialized AA mixtures continue to be investigated. The benefits of using BCAA-enriched formulas are reported for patients with hepatic encephalopathy or hypermetabolism but remain unproved in terms of morbidity and mortality. Recombinant human growth factors, arginine, and glutamine offer promise for their beneficial influences on protein synthesis rates, immunocompetence, and intestinal mucosal barrier protection, respectively.

In vitro and animal studies report an improvement in tissue protein synthesis and reduction in hypermetabolic response with the enteral use of structured lipids containing MCTs and omega-3 fish oil. Because of difficulties reported with the IV use of currently available LCT emulsions such as hepatic and pulmonary complications and immunosuppression, alternate shorter-chain lipid preparations have been investigated. MCTs continue to be explored for IV use as an obligate fuel and an important component of PN. Animal studies with short-chain triglyc-
erides such as triacetin show potential for better protein-sparing properties than MCTs, with less toxicity. Short-chain fatty acids also have been shown to be beneficial in inhibiting small-bowel mucosal atrophy when infused IV or intra-colonically.

New insights into the relationship between nutrition and immune function are emerging through advances with recombinant monokines and new discoveries concerning the involvement of interleukin-1 and tumor necrosis factor in energy metabolism. Although all of these are promising areas of research, they are not considered standard therapy in nutrition support.

REFERENCES

32. Lane HW et al. The effect of selenium supplementation on selenium status of patients receiving chronic total parenteral nutrition. JPN 1987;11:177–82.
PART III

Appendices

Principal Editor: William G. Troutman, PharmD

- Conversion Factors
- Anthropometrics
- Laboratory Indices
- Drug–Laboratory Test Interferences
- Pharmacokinetic Equations
SI UNITS

SI units (le Système International d’Unités) are being introduced in the United States to express clinical laboratory and serum drug concentration data. Instead of employing units of mass (such as micrograms), the SI system uses moles (mol) to represent the amount of a substance. A molar solution contains 1 mole (the molecular weight of the substance in grams) of the solute in 1 liter of solution. The following formula is used to convert units of mass to moles (\(\mu\)g/mL to \(\mu\)mol/L or, by substitution of terms, mg/mL to mmol/L or ng/mL to nmol/L).

\[
\text{Micromoles per Liter (}\mu\text{mol/L}) = \frac{\text{Drug concentration (}\mu\text{g/mL}) \times 1000}{\text{Molecular weight of drug (g/mol)}}
\]

MILLIEQUIVALENTS

An equivalent weight of a substance is that weight which will combine with or replace 1 g of hydrogen; a milliequivalent is 1/1000 of an equivalent weight.

\[
\text{Milliequivalents per Liter (mEq/L)} = \frac{\text{Weight of salt (g)} \times \text{Valence of ion} \times 1000}{\text{Molecular weight of salt}}
\]

\[
\text{Weight of salt (g)} = \frac{\text{mEq/L} \times \text{Molecular weight of salt}}{\text{Valence of ion} \times 1000}
\]

APPROXIMATE MILLIEQUIVALENTS—WEIGHTS OF SELECTED IONS

<table>
<thead>
<tr>
<th>SALT</th>
<th>mEq/g SALT</th>
<th>mg SALT/mEq</th>
</tr>
</thead>
<tbody>
<tr>
<td>Calcium Carbonate (CaCO₃)</td>
<td>20.0</td>
<td>50.0</td>
</tr>
<tr>
<td>Calcium Chloride (CaCl₂ · 2H₂O)</td>
<td>13.6</td>
<td>73.5</td>
</tr>
<tr>
<td>Calcium Gluceptate (Ca[C₇H₁₃O₈]₂)</td>
<td>4.1</td>
<td>245.2</td>
</tr>
<tr>
<td>Calcium Gluconate (Ca[C₆H₁₁O₇]₂ · H₂O)</td>
<td>4.5</td>
<td>224.1</td>
</tr>
<tr>
<td>Calcium Lactate (Ca[C₃H₅O₃]₂ · 5H₂O)</td>
<td>6.5</td>
<td>154.1</td>
</tr>
<tr>
<td>Magnesium Gluconate (Mg[C₆H₁₁O₇]₂ · H₂O)</td>
<td>4.6</td>
<td>216.3</td>
</tr>
<tr>
<td>Magnesium Oxide (MgO)</td>
<td>49.6</td>
<td>20.2</td>
</tr>
</tbody>
</table>

(continued)
The anion gap is the concentration of plasma anions not routinely measured by laboratory screening. It is useful in the evaluation of acid–base disorders. The anion gap is greater with increased plasma concentrations of endogenous (e.g., phosphate, sulfate, lactate, ketoacids) or exogenous (e.g., salicylate, penicillin, ethylene glycol, ethanol, methanol) species. The formulas for calculating the anion gap follow:
(A) Anion Gap = (Na\(^+\) + K\(^+\)) − (Cl\(^−\) + HCO\(_3^−\))

or

(B) Anion Gap = Na\(^+\) − (Cl\(^−\) + HCO\(_3^−\))

where

the expected normal value for A is 11–20 mmol/L;
the expected normal value for B is 7–16 mmol/L.

*Note that there is variation at the upper and lower limits of the normal range.

■ TEMPERATURE

Fahrenheit to Centigrade: \((°F − 32) \times \frac{5}{9} = °C\)
Centigrade to Fahrenheit: \((°C \times \frac{9}{5}) + 32 = °F\)
Centigrade to Kelvin: \(°C + 273 = °K\)

■ WEIGHTS AND MEASURES

**Metric Weight Equivalents**

1 kilogram (kg) = 1000 grams
1 gram (g) = 1000 milligrams
1 milligram (mg) = 0.001 gram
1 microgram (mcg, \(\mu g\)) = 0.001 milligram
1 nanogram (ng) = 0.001 microgram
1 picogram (pg) = 0.001 nanogram
1 femtogram (fg) = 0.001 picogram

**Metric Volume Equivalents**

1 liter (L) = 1000 milliliters
1 deciliter (dL) = 100 milliliters
1 milliliter (mL) = 0.001 liter
1 microliter (\(\mu L\)) = 0.001 milliliter
1 nanoliter (nL) = 0.001 microliter
1 picoliter (pL) = 0.001 nanoliter
1 femtoliter (fL) = 0.001 picoliter

**Apothecary Weight Equivalents**

1 scruple (\(\text{avity}\)) = 20 grains (gr)
60 grains (gr) = 1 dram (\(\text{dary}\))
8 drams (\(\text{dary}\)) = 1 ounce (\(\text{ounce}\))
1 ounce (\(\text{ounce}\)) = 480 grains
12 ounces (\(\text{ounce}\)) = 1 pound (lb)
Apothecary Volume Equivalents

60 minims (m) = 1 fluidram (fl D)
8 fluidrams (fl D) = 1 fluid ounce (fl O)
1 fluid ounce (fl D) = 480 minims
16 fluid ounces (fl D) = 1 pint (pt)

Avoirdupois Equivalents

1 ounce (oz) = 437.5 grains
16 ounces (oz) = 1 pound (lb)

Weight/Volume Equivalents

1 mg/dL = 10 µg/mL
1 mg/dL = 1 mg%
1 ppm = 1 mg/L

Conversion Equivalents

1 gram (g) = 15.43 grains
1 grain (gr) = 64.8 milligrams
1 ounce (oz) = 31.1 grams
1 ounce (oz) = 28.35 grams
1 pound (lb) = 453.6 grams
1 kilogram (kg) = 2.2 pounds
1 milliliter (mL) = 16.23 minims
1 minim (m) = 0.06 milliliter
1 fluid ounce (fl oz) = 29.57 mL

0.1 mg = 1/600 gr
0.12 mg = 1/500 gr
0.15 mg = 1/400 gr
0.2 mg = 1/300 gr
0.3 mg = 1/200 gr
0.4 mg = 1/150 gr
0.5 mg = 1/120 gr
0.6 mg = 1/100 gr
0.8 mg = 1/80 gr
1 mg = 1/65 gr
# CREATININE CLEARANCE FORMULAS

## FORMULAS FOR ESTIMATING CREATININE CLEARANCE IN PATIENTS WITH STABLE RENAL FUNCTION

### Adults [Age 18 Years and Older]¹

\[
\text{Clcr (Males)} = \frac{(140 - \text{Age}) \times (\text{Weight})}{\text{Cr}_s \times 72}
\]

\[
\text{Clcr (Females)} = 0.85 \times \text{Above value}^*
\]

where

- \(\text{Clcr}\) = creatinine clearance in mL/min
- \(\text{Cr}_s\) = serum creatinine in mg/dL
- \(\text{Age}\) is in years
- \(\text{Weight}\) is in kg.

*Some studies suggest that the predictive accuracy of this formula for women is better without the correction factor of 0.85.

### Children [Age 1–18 Years]²

\[
\text{Clcr} = \frac{0.48 \times (\text{Height}) \times (\text{BSA})}{\text{Cr}_s \times 1.73}
\]

where

- \(\text{BSA}\) = body surface area in m²
- \(\text{Clcr}\) = creatinine clearance in mL/min
- \(\text{Cr}_s\) = serum creatinine in mg/dL
- \(\text{Height}\) is in cm.

## FORMULA FOR ESTIMATING CREATININE CLEARANCE FROM A MEASURED URINE COLLECTION

\[
\text{Clcr (mL/min)} = \frac{U \times V^*}{P \times t}
\]

where

- \(U\) = concentration of creatinine in a urine specimen (in same units as \(P\))
- \(V\) = volume of urine in mL
P = concentration of creatinine in serum at the midpoint of the urine collection period (in same units as U)

\( t = \text{time of the urine collection period in minutes (eg, 6 hr = 360 min; 24 hr = 1440 min).} \)

*The product of \( U \times V \) equals the production of creatinine during the collection period and, at steady state, should equal 20–25 mg/kg/day ideal body weight (IBW) in males and 15–20 mg/kg/day IBW in females. If it is less than this, inadequate urine collection may have occurred and \( C_{\text{cr}} \) will be underestimated.

## IDEAL BODY WEIGHT

IBW is the weight expected for a nonobese person of a given height. The IBW formulas below and various life insurance tables can be used to estimate IBW. Most dosing methods described in the literature use IBW as a method in dosing obese patients.

### Adults [Age 18 years and Older]³

\[
\begin{align*}
\text{IBW (Males)} &= 50 + (2.3 \times \text{Height in inches over 5 feet}) \\
\text{IBW (Females)} &= 45.5 + (2.3 \times \text{Height in inches over 5 feet})
\end{align*}
\]

where \( \text{IBW} \) is in kg.

### Children [Age 1–18 Years]²

#### Under 5 Feet Tall:

\[
\text{IBW} = \frac{\text{Height}^2 \times 1.65}{1000}
\]

where

- \( \text{IBW} \) is in kg;
- Height is in cm.

#### 5 Feet or Taller:

\[
\begin{align*}
\text{IBW (Males)} &= 39 + (2.27 \times \text{Height in inches over 5 feet}) \\
\text{IBW (Females)} &= 42.2 + (2.27 \times \text{Height in inches over 5 feet})
\end{align*}
\]

where \( \text{IBW} \) is in kg;

## SURFACE AREA NOMOGRAMS

Nomograms represent the relationship between height, weight, and body surface area in infants and adults. To use a nomogram, a ruler is aligned with the height and weight on the two lateral axes. The point at which the centerline is intersected provides the corresponding value for body surface area.
NOMOGRAM FOR DETERMINATION OF BODY SURFACE AREA FROM HEIGHT AND WEIGHT (INFANTS)\(^4\)

\[ SA = W^{0.3378} \times H^{0.3964} \times 0.024265 \]

where

SA is in \( m^2 \)
Height (H) is in cm
Weight (W) is in kg.

Reproduced from reference 4, with permission.
NOMOGRAM FOR DETERMINATION OF BODY SURFACE AREA FROM HEIGHT AND WEIGHT (ADULTS)\textsuperscript{5}

\begin{align*}
SA & = W^{0.425} \times H^{0.725} \times 71.84
\end{align*}

where

- \( SA \) is in m\(^2\)
- Height (H) is in cm
- Weight (W) is in kg.

*Reproduced from reference 5, with permission.*

**REFERENCES**

The following table lists typical reference ranges for clinical laboratory tests in common use. Reference ranges for laboratory tests can vary widely among testing facilities, often as a result of methodologic differences. It is therefore always advisable to obtain reference ranges from the laboratory performing the analyses. Laboratory test results should never be accepted without correct identification of the units of measurement because most tests can be reported in several systems of measurement. The table presents conventional and international (usually the same as Système International, or SI) units.

The following abbreviations are used to identify the specimen:

(P) — Plasma
(S) — Serum
(U) — Urine
(WB) — Whole Blood
(WB, art) — Whole Blood, Arterial

The table begins on page 1062.
<table>
<thead>
<tr>
<th>TEST/SPECIMEN</th>
<th>AGE GROUP OR OTHER FACTOR</th>
<th>REFERENCE RANGE</th>
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<td>30–39 yr, M</td>
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<td>Iron (S)</td>
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<td>Lipase (S)</td>
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<td>&gt;60 yr</td>
<td>280–301</td>
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(continued)
**Test/Specimen** | **Age Group or Other Factor** | **Reference Range**
--- | --- | ---
**Osmolal Gap** | ≤10 | ≤10
Measured Osmolality = Calculated Osmolality
Calculated Osmolality = 2(Na⁺) + (Glucose/18) + (BUN/2.8)
**Oxygen, Partial Pressure (WB, art) (pO₂)** | 83–108 mm Hg 11.04–14.36 kPa
(Decreases with age and altitude)
**pH (WB, art)** | 7.35–7.45 | 7.35–7.45
**Phosphorus, Inorganic (S)** | mg/dL mmol/L
Child | 4.5–5.5 | 1.45–1.78
Adult | 2.7–4.5 | 0.87–1.45
>60 yr, M | 2.3–3.7 | 0.74–1.20
>60 yr, F | 2.8–4.1 | 0.90–1.32
**Potassium (S,P)** | mEq/L mmol/L
Child | 3.4–4.7 | 3.4–4.7
Adult | 3.5–5.1 | 3.5–5.1
**Protein, Total (S)** | g/dL g/L
Adult | | |
Ambulatory | 6.4–8.3 | 64–83
Recumbent | 6.0–7.8 | 60–78
>60 yr | lower by 0.2 | lower by 2
**Albumin** | g/dL g/L
Adult | 3.5–5.0 | 35–50
>60 yr | 3.7–4.7 | 37–47
**Globulins** | g/dL g/L
Adult | 2.3–3.5 | 23–35
**Prealbumin** | g/dL g/L
Adult | 10–40 mg/dL | 100–400 mg/L
**Sodium (S,P)** | mEq/L mmol/L
Child | 138–145 | 138–145
Adult | 136–146 | 136–146
**Thyroid-Stimulating Hormone (S,P)** | µunits/mL munits/L
(TSH) | µunits/mL munits/L
Child | 4.5 ± 3.6 | 4.5 ± 3.6
Adult | <10 | <10
>60 yr, M | 2–7.3 | 2–7.3
>60 yr, F | 2–16.8 | 2–16.8

(continued)
## BLOOD, SERUM, PLASMA CHEMISTRY (continued)

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<th>AGE GROUP OR OTHER FACTOR</th>
<th>REFERENCE RANGE</th>
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<td>Conventional</td>
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<td>Thyroxine, Total (S)</td>
<td>µg/dL</td>
<td>nmol/L</td>
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<td>5–10 yr</td>
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<td>5–12</td>
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<tr>
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<td>&gt;60 yr, M</td>
<td>5–10</td>
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<td>&gt;60 yr, F</td>
<td>5.5–10.5</td>
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<td>4–9 mo pregnant</td>
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<td>Transferrin (S)</td>
<td>mg/dL</td>
<td>g/L</td>
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<td>M</td>
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</tr>
<tr>
<td>12–15 yr</td>
<td>36–138</td>
<td>41–138</td>
</tr>
<tr>
<td>16–19 yr</td>
<td>40–163</td>
<td>40–128</td>
</tr>
<tr>
<td>20–29 yr</td>
<td>44–185</td>
<td>40–128</td>
</tr>
<tr>
<td>30–39 yr</td>
<td>49–284</td>
<td>38–160</td>
</tr>
<tr>
<td>40–49 yr</td>
<td>56–298</td>
<td>44–186</td>
</tr>
<tr>
<td>50–59 yr</td>
<td>62–288</td>
<td>55–247</td>
</tr>
<tr>
<td>Desired, Adult</td>
<td>&lt;150</td>
<td>&lt;150</td>
</tr>
<tr>
<td>Triiodothyronine Resin Uptake (S)</td>
<td>% of Total</td>
<td>Fraction of Total</td>
</tr>
<tr>
<td>(T₃RU)</td>
<td>Adult</td>
<td>24–34</td>
</tr>
<tr>
<td></td>
<td>&gt;60 yr, M</td>
<td>24–32</td>
</tr>
<tr>
<td></td>
<td>&gt;60 yr, F</td>
<td>22–32</td>
</tr>
<tr>
<td>Triiodothyronine, Total (S)</td>
<td>ng/dL</td>
<td>nmol/L</td>
</tr>
<tr>
<td>(T₃)</td>
<td>10–15 yr</td>
<td>80–210</td>
</tr>
<tr>
<td></td>
<td>Adult</td>
<td>120–195</td>
</tr>
<tr>
<td></td>
<td>&gt;60 yr, M</td>
<td>105–175</td>
</tr>
<tr>
<td></td>
<td>&gt;60 yr, F</td>
<td>108–205</td>
</tr>
<tr>
<td>Urea Nitrogen (S)</td>
<td>mg/dL</td>
<td>mmol/L urea</td>
</tr>
<tr>
<td>(BUN)</td>
<td>Child</td>
<td>5–18</td>
</tr>
<tr>
<td></td>
<td>Adult</td>
<td>7–18</td>
</tr>
<tr>
<td></td>
<td>&gt;60 yr</td>
<td>8–21</td>
</tr>
<tr>
<td>Uric Acid (S)</td>
<td>mg/dL</td>
<td>mmol/L</td>
</tr>
<tr>
<td>(Uricase Method)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Child</td>
<td>2.0–5.5</td>
<td>0.12–0.32</td>
</tr>
<tr>
<td>Adult, M</td>
<td>3.5–7.2</td>
<td>0.21–0.42</td>
</tr>
<tr>
<td>Adult, F</td>
<td>2.6–6.0</td>
<td>0.15–0.35</td>
</tr>
</tbody>
</table>
### URINE, RENAL FUNCTION TESTS

<table>
<thead>
<tr>
<th>TEST/SPECIMEN</th>
<th>AGE GROUP OR OTHER FACTOR</th>
<th>REFERENCE RANGE</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td><strong>Conventional</strong></td>
</tr>
<tr>
<td>Catecholamines, 24-hr (U)</td>
<td>&lt;110 µg</td>
<td>&lt;650 nmol</td>
</tr>
<tr>
<td>Creatinine, 24-hr (U)</td>
<td>mg/kg</td>
<td>µmol/kg</td>
</tr>
<tr>
<td>Child</td>
<td>8–22</td>
<td>71–195</td>
</tr>
<tr>
<td>Adolescent</td>
<td>8–30</td>
<td>71–265</td>
</tr>
<tr>
<td>Adult, M</td>
<td>14–26</td>
<td>124–230</td>
</tr>
<tr>
<td>Adult, F</td>
<td>11–20</td>
<td>97–177</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Decreases with age to 10 mg/kg/day at age 90.</td>
</tr>
<tr>
<td>Creatinine Clearance (S, P, and U)</td>
<td>mL/min/1.73 m²</td>
<td>mL/sec/m²</td>
</tr>
<tr>
<td>&lt;40 yr, M</td>
<td>97–137</td>
<td>0.93–1.32</td>
</tr>
<tr>
<td>&lt;40 yr, F</td>
<td>88–128</td>
<td>0.85–1.23</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Decreases with age &gt;40 yr.</td>
</tr>
<tr>
<td>Inulin Clearance (S and U)</td>
<td>mL/min/1.73 m²</td>
<td>mL/sec/m²</td>
</tr>
<tr>
<td></td>
<td>M</td>
<td>F</td>
</tr>
<tr>
<td>20–29 yr</td>
<td>90–174</td>
<td>84–156</td>
</tr>
<tr>
<td>30–39 yr</td>
<td>88–168</td>
<td>82–150</td>
</tr>
<tr>
<td>40–49 yr</td>
<td>78–162</td>
<td>82–146</td>
</tr>
<tr>
<td>50–59 yr</td>
<td>68–152</td>
<td>66–142</td>
</tr>
<tr>
<td>60–69 yr</td>
<td>57–137</td>
<td>58–130</td>
</tr>
<tr>
<td>70–79 yr</td>
<td>42–122</td>
<td>45–121</td>
</tr>
<tr>
<td>80–89 yr</td>
<td>39–105</td>
<td>39–105</td>
</tr>
<tr>
<td></td>
<td>M</td>
<td>F</td>
</tr>
<tr>
<td></td>
<td>0.87–1.68</td>
<td>0.81–1.50</td>
</tr>
<tr>
<td></td>
<td>0.85–1.62</td>
<td>0.79–1.44</td>
</tr>
<tr>
<td></td>
<td>0.75–1.56</td>
<td>0.79–1.41</td>
</tr>
<tr>
<td></td>
<td>0.65–1.46</td>
<td>0.63–1.37</td>
</tr>
<tr>
<td></td>
<td>0.55–1.32</td>
<td>0.56–1.25</td>
</tr>
<tr>
<td></td>
<td>0.40–1.17</td>
<td>0.43–1.17</td>
</tr>
<tr>
<td></td>
<td>0.38–1.01</td>
<td>0.38–1.01</td>
</tr>
<tr>
<td>pH (U)</td>
<td>4.5–8</td>
<td>4.5–8</td>
</tr>
<tr>
<td>Protein, Total (U)</td>
<td>1–14 mg/dL</td>
<td>10–140 mg/L</td>
</tr>
<tr>
<td></td>
<td>At Rest</td>
<td>50–80 mg/day</td>
</tr>
<tr>
<td>Specific Gravity, Random (U)</td>
<td>1.002–1.030</td>
<td>1.002–1.030</td>
</tr>
<tr>
<td>Uric Acid, 24-hr (U)</td>
<td>250–750 mg</td>
<td>1.48–4.43 mmol</td>
</tr>
<tr>
<td>TEST/SPECIMEN</td>
<td>AGE GROUP OR OTHER FACTOR</td>
<td>REFERENCE RANGE</td>
</tr>
<tr>
<td>-------------------------------</td>
<td>---------------------------</td>
<td>-----------------------------------</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Conventional</td>
</tr>
<tr>
<td></td>
<td></td>
<td>International Units</td>
</tr>
<tr>
<td>Bleeding time</td>
<td>3–9 min</td>
<td>180–540 sec</td>
</tr>
<tr>
<td>Erythrocyte Count (WB)</td>
<td>× $10^6/\mu$L</td>
<td>× $10^{12}/L$</td>
</tr>
<tr>
<td></td>
<td>M</td>
<td>4.6–6.2</td>
</tr>
<tr>
<td></td>
<td>F</td>
<td>4.2–5.4</td>
</tr>
<tr>
<td>Erythrocyte Indices (WB)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Mean Corpuscular Volume</td>
<td>80–96 µm³</td>
<td>80–96 fL</td>
</tr>
<tr>
<td>Mean Corpuscular Hemoglobin</td>
<td>27–31 pg</td>
<td>27–31 pg</td>
</tr>
<tr>
<td>Erythrocyte Sedimentation Rate (WB)</td>
<td>mm/hr</td>
<td>mm/hr</td>
</tr>
<tr>
<td></td>
<td>M</td>
<td>1–13</td>
</tr>
<tr>
<td></td>
<td>F</td>
<td>1–20</td>
</tr>
<tr>
<td>Fibrinogen (P)</td>
<td></td>
<td>200–400 mg/dL</td>
</tr>
<tr>
<td></td>
<td></td>
<td>2.00–4.00 g/L</td>
</tr>
<tr>
<td>Hematocrit (WB)</td>
<td>% Packed RBC Volume</td>
<td>Volume Fraction</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>6–12 yr</td>
<td>35–45</td>
<td>0.35–0.45</td>
</tr>
<tr>
<td>12–18 yr, M</td>
<td>37–49</td>
<td>0.37–0.49</td>
</tr>
<tr>
<td>12–18 yr, F</td>
<td>36–46</td>
<td>0.36–0.46</td>
</tr>
<tr>
<td>18–49 yr, M</td>
<td>41–53</td>
<td>0.41–0.53</td>
</tr>
<tr>
<td>18–49 yr, F</td>
<td>36–46</td>
<td>0.36–0.46</td>
</tr>
<tr>
<td>Hemoglobin (WB)</td>
<td>g/dL</td>
<td>mmol/L</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>6–12 yr</td>
<td>11.5–15.5</td>
<td>1.78–2.40</td>
</tr>
<tr>
<td>12–18 yr, M</td>
<td>13.0–16.0</td>
<td>2.02–2.48</td>
</tr>
<tr>
<td>12–18 yr, F</td>
<td>12.0–16.0</td>
<td>1.86–2.48</td>
</tr>
<tr>
<td>18–49 yr, M</td>
<td>13.5–17.5</td>
<td>2.09–2.71</td>
</tr>
<tr>
<td>18–49 yr, F</td>
<td>12.0–16.0</td>
<td>1.86–2.48</td>
</tr>
<tr>
<td>Hemoglobin A1C (WB)</td>
<td>5.3–7.5% of total Hb</td>
<td>0.053–0.075</td>
</tr>
<tr>
<td>Leukocyte Count (WB)</td>
<td>4.5–11 × $10^3/\mu$L</td>
<td>4.5–11 × $10^9/L$</td>
</tr>
<tr>
<td></td>
<td>Segs</td>
<td>31–71%</td>
</tr>
<tr>
<td></td>
<td>Bands</td>
<td>0–12%</td>
</tr>
<tr>
<td></td>
<td>Lymphocytes</td>
<td>15–50%</td>
</tr>
<tr>
<td></td>
<td>Monocytes</td>
<td>0–12%</td>
</tr>
<tr>
<td></td>
<td>Eosinophils</td>
<td>0–5%</td>
</tr>
<tr>
<td></td>
<td>Basophils</td>
<td>0–2%</td>
</tr>
</tbody>
</table>

(continued)
### HEMATOLOGY (continued)

<table>
<thead>
<tr>
<th>TEST/SPECIMEN</th>
<th>AGE GROUP OR OTHER FACTOR</th>
<th>REFERENCE RANGE</th>
</tr>
</thead>
<tbody>
<tr>
<td>Absolute Neutrophil Count (ANC)</td>
<td></td>
<td>ANC = (% Segs + % Bands) × Leukocyte Count</td>
</tr>
<tr>
<td>Partial Thromboplastin Time, Activated (WB) (aPTT)</td>
<td></td>
<td>25–37 sec</td>
</tr>
<tr>
<td>Platelets (WB)</td>
<td></td>
<td>150–440 × 10^3/µL, 0.15–0.44 × 10^{12}/L</td>
</tr>
<tr>
<td>Prothrombin Time (WB)</td>
<td></td>
<td>Less than 2-sec deviation from control.</td>
</tr>
<tr>
<td>Reticulocytes (WB)</td>
<td></td>
<td>0.5–1.5%, 0.005–0.015 of erythrocytes</td>
</tr>
</tbody>
</table>

#### REFERENCES

The following table lists common clinical laboratory tests and drugs that can interfere with those tests. Drugs can interfere with laboratory tests through pharmacological or toxic effects or through actual chemical interference with the testing process. Either effect can lead to an altered value of the laboratory test, resulting in an inappropriate diagnosis or treatment. It is therefore essential that clinicians recognize possible drug–laboratory interactions and use this information in the overall assessment of a patient’s clinical status.

The table lists drug interferences with the most common laboratory tests. For detailed information on laboratory tests not covered here, see references 1–3 at the end of this section. Also, it should be noted that drugs can interfere with laboratory tests by many different mechanisms. The reader should refer to the references cited in the table and other relevant sources to obtain more information about a specific test.

The following abbreviations are used in the table:

| (B) | — Blood |
| (CSF) | — Cerebrospinal Fluid |
| (I) | — Analytical Interference of Drug |
| (P) | — Pharmacological/Toxic Effect of Drug |
| (S) | — Serum |

### DRUGS THAT CAN AFFECT RESULTS AND CAUSE OF INTERFERENCE

**BLOOD, SERUM, PLASMA CHEMISTRY**

**Alkaline Phosphatase (S).** Elevated by acetaminophen (P), acetoheamidamid (P), albumin (I), alphetretinoin (P), allopurinol (P), aluminum salts (P), aminoglycosides (P), amiodarone (P), amphotericin B (P), anabolic steroids (P), azathioprine (P), barbiturates (P), bromocriptine (P), carbamazepine (P), cephalosporins (P), cheno-

diol (P), clofibrate (P), cyclophosphamide (P), cyclosporine (P), cytarabine (P),
danzol (P), dantrolene (P), dapsone (P), disulfiram (P), docetaxel (P), eryth-
romycin (P), estrogens (P), filgrastim (P), flucytosine (P), glycopyrrolate (P), gold salts (P), griseofulvin (P), haloperidol (P), hepatotoxic drugs (P), HMG-CoA
reductase inhibitors (P), hydralazine (P), ibuprofen (I,P), isoniazid (P), isotretinoin (P), ketoconazole (P), lithium salts (P), meprobamate (P), mercaptopurine (P),
metyldopa (P), mitomycin (P), nafarelin (P), niacin (P), nitrofurantoin (P), non-steroidal anti-inflammatory drugs (P), papaverine (P), penicillamine (P), penicillins (P), phenazopyridine (P), phenothiazines (P), phenytoin (P), pindolol (I), probenecid (P), propylthiouracil (P), pyrazinamide (P), quinidine (P), rifampin (P), sulfonamides (P), sulfonymylease (P), tetracyclines (P), thiabendazole (P), ticlopidine (P), topotecan (P), trimethoprim (P), troleandomycin (P), valproic acid (P), zidovudine (P).

Decreased by bisphosphonates (P), calcitriol (P), carvedilol (P), citrate salts (I), clofibrate (P), cyclosporine (P), danazol (P), EDTA (I), estrogens (P), fluoride salts (I), phosphate salts (I), prednisolone (P), prednisone (P), tamoxifen (P), theophylline (I), tricyclic antidepressants (P), ursodiol (P), zinc (I).

Aminotransferases (ALT [SGOT] or AST [SGPT]) (S). Elevated by abacavir (P), acarbose (P), acetaminophen (I,P), acitretin (P), allopurinol (P), amiodarone (P), ampicillin (I), anabolic steroids (P), anastrozole (P), aspirin (P), azathioprine (P), aztreonam (P), bisephosphonates (P), carbamazepine (P), cephalosporins (P), chloramphenicol (P), chloridazepoxide (I,P), cholestyramine (P), cholestyramine (P), cholinergic agents (P), clotidogrel (P), COX-2 inhibitors (P), cyclophosphamide (P), cytarabine (P), danazol (P), dantrolene (P), delavirdine (P), denileukin diftitox (P), disulfiram (P), diuretics (thiazide) (P), docetaxel (P), efavirenz (P), erythromycin (I,P), estrogens (P), etoposide (P), fenofibrate (P), fluconazole (P), flucytosine (P), flutamide (P), fomepizole (P), ganciclovir (P), gemcitabine (P), gentamicin (P), glucose-6-phosphate (P), gold salts (P), griseofulvin (P), haloperidol (P), heparin (P), hepatotoxic drugs (P), HMG-CoA reductase inhibitors (P), hydralazine (P), interferons (P), indinavir (P), interferon alfa-2a (P), interferon beta-1a (P), interferon beta-1b (P), irinotecan (P), isoniazid (P), isotretinoin (I), ketotifen (P), ketoconazole (P), levodopa (I,P), meprobamate (P), mercaptopurine (P), metronidazole (P), methylxophosphonate (P), naltrexone (P), narcotics (I,P), nevirapine (P), niacin (P), nilutamide (P), nitrofurantoin (P), nonsteroidal anti-inflammatory drugs (P), olanzapine (P), penicillamine (P), penicillins (I,P), pentosan polysulfate sodium (P), phenazopyridine (P), phenothiazines (P), porfimer (P), probenecid (P), propylthiouracil (P), pyrazinamide (P), quetiapine (P), quinupristin/dalfopristin (P), rifabutin (P), rifampin (P), rifampicin (P), salicylates (P), sulfonamides (P), sulfonymylease (P), tacrine (P), temozolomide (P), tetracyclines (P), thiabendazole (P), ticlopidine (P), tolcapone (P), total parenteral nutrition (P), troleandomycin (P), valproic acid (P), vidarabine (P), vinorelbine (P), vitamin C, zafirlukast (P), zalcitabine (P), zidovudine (P).

Decreased by acetaminophen (I), aspirin (I), cyclosporine (P), fluoride salts (I), interferons (P), metronidazole (I), naltrexone (P), pindolol (I), rifampin (I), tricyclic antidepressants (P), ursodiol (P), vitamin C (I), zalcitabine (P).

Ammonia (B). Elevated by acetazolamide (P), alcohol (P), ammonium chloride (P), aspirin (I), barbiturates (P), benzamidazepine (P), diuretics (loop, thiazide) (P), metronidazole (I), naltrexone (P), nicotine (P), tobacco (P), valproic acid (P), vitamin C (I), zalcitabine (P).
Decreased by cefotaxime (I), kanamycin, oral (P), *Lactobacillus acidophilus* (P), lactulose (P), MAO inhibitors (P), neomycin, oral (P), phosphate salts (I), potassium salts (P), tetracycline (P).1,2,4

**Amylase (S).** Elevated by alcohol (P), angiotensin II receptor blockers (P), ACE inhibitors (P), asparaginase (P), azathioprine (P), chloride salts (I), cholinergic agents (P), cisplatin (P), contraceptives, oral (P), corticosteroids (P), denileukin diftitox (P), didanosine (P), diuretics, loop and thiazide (P), erythromycin (P), estrogens (P), fluoride salts (I), indinavir (P), lamivudine (P), metronidazole (P), narcotics (P), nitrofurantoin (P), opioids (P), pancreatotoxins (P), potassium iodide (P), rifampin (P), ritonavir (P), sulfonamides (P), valproic acid (P), vinorelbine (P), zalcitabine (P).1,3,5,7

Decreased by anabolic steroids (P), cefotaxime (I), citrate salts (P), fluoride salts (I), somatostatin (P).1

**Bilirubin, Total (S).** Elevated by acarbose (P), acetaminophen (P), acetohexamide (P), allopurinol (P), amiodarone (P), amphotericin B (I,P), anabolic steroids (P), asparaginase (P), azathioprine (P), barbiturates (P), capceticabine (P), carbamazepine (P), cephalosporins (P), chloramphenicol (P), cholinergics (P), colchicine (P), cyclophosphamide (P), cyclosporine (P), cytarabine (P), danazol (P), dantrolene (P), dapsone (P), dextan (I), disulfiram (P), diuretics, thiazide and loop (P), docetaxel (P), epinephrine (I), erythromycin (P), estrrogens (P), etoposide (P), fluotamide (P), gemtuzumab ozogamicin (P), glycopyrrolate (P), gold salts (P), haloperidol (P), hemolytic agents (P), hepatotoxic drugs (P), HMG-CoA reductase inhibitors (P), hydralazine (P), indinavir (P), interferon beta-1b (P), irinotecan (P), isoniazid (P), isoproterenol (I), isotretinoin (P), ketoconazole (P), levodopa (I), meprobamate (P), mercaptopurine (P), methimazole (P), methotrexate (L,P), methylotulpa (I,P), narcotics (I), niacin (P), nitrofurantoin (L,P), nonsteroidal anti-inflammatory drugs (P), papaverine (P), penicillamine (P), penicillins (P), phenazopyridine (I), phenothiazines (P), phenytoin (P), pranobecid (P), propranolol (I), propylthiouracil (P), pyrazinamide (P), quinidine (P), quinupristin/dalfopristin (P), rifampin (I,P), rifampent (P), riluzole (P), salicylates (I,P), sulfonamides (P), sulfonylureas (P), theophylline (I), thiabendazole (P), topotecan (P), troleandomycin (P), valproic acid (P), vitamin C (I), zafirlukast (P), zidovudine (P).1–3,5

Decreased by amikacin (I), barbiturates (especially in newborns) (P), carbamazepine (P), corticosteroids (P), cyclosporine (P), fexofenadine (P), isotretinoin (P), levodopa (I), nitrofurantoin (I), phenazopyridine (I), phenytoin (P), pindolol (I), sulfonamides (P), temozolomide (P), theophylline (I), ursodiol (P), vitamin C (I).3,4

**Calcium (S).** Elevated by alitretinoin (P), amifostine (P), anabolic steroids (P), androgens (P), basiliximab (P), calcitriol (P), calcium salts (P), cefotaxime (I), chlorpropamide (I), diuretics, thiazide (P), estrogens (P), hydralazine (I), interferons (I), iron salts (I), lithium salts (P), magnesium salts (I), phenobarbital (P), progesterins (P), sevelamer (P), tamoxifen (P), toremifene (P), thyroid (P), vitamin A (P), vitamin D (P).1–5,7
Decreased by acetazolamide (P), albuterol (P), asparaginase (P), aspirin (I), bisphosphonates (P), calcitonin (P), carbamazepine (P), cisplatin (P), citrate salts (P), contraceptives, oral (P), corticosteroids (P), diuretics, loop (P), EDTA (I), ethanol (P), fluoride salts (LP), fosfomycin (P), glucagon (P), heparin (I), laxatives (P), magnesium salts (P), phenobarbital (P), phenytoin (P), phosphate salts (P), plicamycin (P), sodium polystyrene sulfonate (P), sulfisoxazole (I), zalcitabine (P).

**Carbon Dioxide (B). Elevated by** bicarbonate salts (P), diuretics (loop, thiazide) (P), respiratory depressants (P).

Decreased by acetazolamide (P), aspirin overdose (P), nephrotoxic drugs (P), theophylline (P).

**Chloride (S). Elevated by** acetazolamide (P), anabolic steroids (P), aspirin (LP), carbamazepine (I), cefotaxime (I), cholestyramine (P), corticosteroids (by salt retention) (P), COX-2 inhibitors (P), cyclosporine (P), diuretics, carbonic anhydrase inhibitor, thiazide—chronically by alkalosis (P), estrogens (P), guanethidine (P), halogens (eg, bromides, fluorides) (I), methyl dopa (P), nonsteroidal anti-inflammatory drugs (P), sodium phenylbutyrate (P).

Decreased by allopurinol (I), bicarbonates (P), cefotaxime (metabolite) (I), chloropropamide (P), corticosteroids (by alkalosis) (P), diuretics, loop, thiazide—by acute diuresis (P), fluoride salts (I), laxatives, long-term use (P), mannitol (P), mineralocorticoids (by alkalosis) (P), trimethoprim (P).

**Cholesterol, Total (S). Elevated by** acetohexamide (P), β-adrenergic blocking agents (P), alitretinoin (P), amiodarone (P), amphotericin B (I), amphenavir (P), anabolic steroids (by cholestasis) (P), aspirin (I), basiliximab (P), carbamazepine (P), cefotaxime (I), chenodiol (P), clopidogrel (P), contraceptives, oral (P), corticosteroids (LP), cyclosporine (P), danazol (P), dextran (I), diclofenac (P), disulfiram (P), diuretics, loop, thiazide (P), ethanol (P), fribates (P), gold salts (P), hepatotoxic drugs (cholestatic effect) (P), ibuprofen (P), imipramine (P), isorotinoin (P), meprobamate (P), methotrexate (I), mirtazapine (P), mycophenolate (P), nafarelin (P), phenobarbital (P), phenothiazines (LP), phentoyin (LP), protease inhibitors (P), quetiapine (P), ritonavir (P), rosiglitazone (P), sirolimus (P), smoking (P), sorbitol (P), sotalol (P), spironolactone (P), sulfadiazine (P), tamoxifen (P), tetracycline (I), thibendazole (P), ticlopidine (P), vitamin A (I), vitamin C (LP), vitamin D (LP).

Decreased by acarbose (P), acebutolol (P), α-adrenergic blockers (P), allopurinol (LP), aluminum salts (P), amiloride (P), amiodarone (P), ampicillin (I), anabolic steroids (by inhibiting synthesis) (P), ACE inhibitors (P), asparaginase (P), azathioprine (P), calcium channel blockers (P), carvedilol (P), chlorpropamide (P), cholestyramine (P), citrate salts (I), clofibrate (P), clomiphene (P), colchicine (P), colestipol (P), diuretics, thiazide (P), estrogens (P), fenofibrate (P), fluoride salts (I), haloperidol (P), hepatotoxic drugs (decreased synthesis) (P), HMG-CoA reductase inhibitors (P), hydroxychloroquine (P), insulin (P), isoniazid (P), isorotinoin (P), kanamycin, oral (P), ketoconazole (P), levothyroixne (P), MAOIs (P), metformin (P), methyl dopa (LP), metronidazole (P), neomycin, oral (P), niacin (P), nitrates (I), orlistat (P), penicillamine (I), pentamidine (P), phenytoin (P), pindolol (P), psyllium (P), raloxifene (P), rifampin (I), sevelamer (P) tamox-
ifen (P), tetracyclines (P), thyroid (P), ursodiol (P), valproic acid (P), vitamin C (I,P).1–5

**Coombs’ [Direct] (S).** Positive by aztreonam (P), captopril (P), cephalosporins (P), chlorpromazine (P), chlorpropamide (P), ethosuximide (P), hemolytic agents (P), hydralazine (P), imipenem/cilastatin (P), indomethacin (P), isoniazid (P), levo-
dopa (P), mefenamic acid (P), mephalan (P), methyl dopa (P), nitrofurantoin (P),
penicillamine (P), penicillins (P), phenyltoin (P), procainamide (P), quinine (P), quinine (P), rifampin (P), sulfasalazine (P), sulfonamides (P), sulfonylureas (P), tetracyclines (P), tolmetin (I).1,3,4,6

**Creatine Kinase (S).** Elevated by alcohol (chronic) (P), aminocaproic acid (P), amphotericin B (P), barbiturates (P), cefotaxime (I), clofibrate (P), cyclosporine (P), danazol (P), fenoibrate (P), HMG-CoA reductase inhibitors (P), gemfibrozil (P), IM injections (P), lithium salts (P), niacin (P), succimer (I), saquinavir (P), zidovudine (P).1,2,4,5

Decreased by amikacin (I), anesthetic agents (P), ascorbic acid (I), aspirin (I), dantrolene (P), phenothiazines (P), pindolol (I), succinylcholine (P), sul-famethoxazole (P), zalcitabine (P).1

**Creatinine (S).** Elevated by acebutolol (P), acetaminophen (I,P), acetohexamide (I), acyclovir (P), aminoglycosides (P), amiodarone (P), amphotericin B (P), ACE inhibitors (P), antacids (P), asparaginase (P), aztreonam (P), carvedilol (P), cephalosporins (Jaffe method) (I,P), chloroquine (P), cidofovir (P), cimetidine (P), cisplatin (P), clofibrate (P), colistin (P), co-trimoxazole (P), cy-
closporine (P), demeclocycline (P), denileukin difitox (P), dextran (P), diuretics (P), dopamine (I), doxycycline (P), fluycytosine (I,P), foscarinet (P), furosemide (I), ganciclovir (P), hydroxychloroquine (P), lactulose (I), levodopa (I), lidocaine (I), lithium (I,P), methicillin (P), methyl dopa (I), mitomycin (P), nalidixic acid (P), nephrotoxic drugs (P), nifedipine (P), nitrofurantoin (I), nonsteroidal anti-
inflammatory drugs (P), penicillamine (P), penicillin (I), pentamidine (P), phosphate salts (P), radiocontrast agents (P), ritonavir (P), salicylates (P), sirolimus (P), sulfactam (I), sulfamethoxazole (I), tacrolimus (P), tetracycline (P), vanco-
mycin (P), vitamin C (I), vitamin D (P).1–4,7

Decreased by amikacin (I), cephalosporins (I), citrate salts (I), dopamine (I), ibuprofen (I), interferon alfa-2a (P), methyldopa (I), sulfonylureas (P), vitamin C (I).1,5

**Glucose (S).** Elevated by abacavir (P), acetaminophen (SMA 12/60 method) (I), acet-
etazolamide (P), β-adrenergic blocking agents (also mask hypoglycemia) (P), al-
buterol (P), amiodarone (P), antidepressants (heterocyclic) (P), asparaginase (P), basiliximab (P), bicalutamide (P), cefotaxime (I), cholestyramine (P), citrate salts (I), clonidine (P), clozapine (P), corticosteroids (P), cyclosporine (P), daclizumab (P), dextran (I), dextroamphetamine (P), diazoxide (P), diclofenac (I), diltiazem (P), diuretics, loop and thiazide (P), epinephrine (I,P), ephedrine (P), estrogens (P), fos-
phenytoin (P), gemfibrozil (P), glucagon (P), interferon alfa-2a (P), iron dextran (I), isoniazid (P), isoproterenol (I), labetalol (I), lactose (I), levodopa (SMA 12/60
method) (I), lipids (P), lithium salts (P), mercaptopurine (I), methyldopa (I), metronidazole (I), mycophenolate (P), naldixic acid (I), niacin (LP), nifedipine (P), octreotide (P), olanzapine (P), pentamidine (IV, paradoxical effect) (P), perphenazine (P), phenothiazines (P), phenytoin (P), pravastatin (P), progestins (P), propranolol (P), propylthiouracil (I), protease inhibitors (P), reserpine (P), rifampin (LP), salicylates (acute toxicity) (LP), somatostatin (P), sorbitol (P), tacrolimus (P), terbutaline (P), tetracyclines (P), thiabendazole (P), thyroid (P), tolbutamide (P), vitamin C (neocuproin method) (I), zalcitabine (P).

Decreased by acarbose (P), acetaminophen (GOD-Perid method) (LP), acetazolamide (P), β-adrenergic blocking agents (nonselective) (P), alcohol (P), allopurinol (P), amikacin (I), anabolic steroids (P), anesthetics (P), chloroquine (P), chlorpropamide (LP), cimetidine (P), clonidine (P), doxazosin (P), erythromycin (P), estrogen (P), fenofibrate (P), interferon beta-1b (P), hydralazine (I), insulin (P), isoniazid (I), levodopa (glucose oxidase and other methods) (I), lipids (I), MAO inhibitors (P), metformin (P), methyldopa (I), metronidazole (I), miglitol (P), niacin (P), octreotide (P), pentamidine (IV) (P), phenazopyridine (I), phosphorus (P), psyllium (P), salicylates (acute and chronic toxicity) (P), saquinavir (P), SSRIs (P), sulfonamides (P), sulfonylureas (P), tetracyclines (I), thiabendazole (P), tolbutamide (I), tolbutamide (LP), verapamil (P), vitamin C (GOD-Perid method) (LP).

Iron (S). Elevated by cefotaxime (I), chloramphenicol (P), contraceptives (oral) (P), estrogens (P), ferrous salts (I), iron, parenteral (LP), methyldopa (P), miglitol (P), rifampin (I).

Decreased by allopurinol (P), aspirin (large doses) (P), clofibrate (P), cimetidine (P), disopyramide (P), doxazosin (P), erythromycin (P), estrogen (P), fenofibrate (P), interferon beta-1b (P), hydralazine (I), insulin (P), isoniazid (I), levodopa (glucose oxidase and other methods) (I), lipids (I), MAO inhibitors (P), metformin (P), methyldopa (I), metronidazole (I), miglitol (P), niacin (P), octreotide (P), pentamidine (IV) (P), phenazopyridine (I), phosphorus (P), psyllium (P), salicylates (acute and chronic toxicity) (P), saquinavir (P), SSRIs (P), sulfonamides (P), sulfonylureas (P), tetracyclines (I), thiabendazole (P), tolbutamide (I), tolbutamide (LP), verapamil (P), vitamin C (GOD-Perid method) (LP).

Iron Binding Capacity, Total (S). Elevated by contraceptives, oral (P), propylthiouracil (P).

Decreased by chloramphenicol (P), corticotropin (P), corticosteroids (P).

Magnesium (S). Elevated by cefotaxime (I), diuretics, potassium-sparing (P), lithium salts (P), magnesium salts (P), metformin (P).

Decreased by albuterol (P), alcohol (P), amifostine (P), aminoglycosides (P), amphotericin B (P), bisphosphonates (P), calcium salts (I), cefotaxime (I), cisdiphenyl (P), citrate salts (I), contraceptive, oral (P), cyclosporine (P), digitalis (toxic concentrations) (P), diuretics, loop, and thiazide (P), foscarnet (P), glucagon (P), insulin (P), tacrolimus (P).

Osmolality (S). Elevated by alcohol (ADH suppression) (P), citrate salts (I), corticosteroids (P), demeclocycline (ADH inhibition) (P), glucose (I), lithium salts (ADH inhibition) (P), mannitol (LP).

Decreased by antidepressants, tricyclic (P), carbamazepine (P), chlorpropamide (P), clonidine (P), cyclophosphamide (P), cytarabine (P), diuretics, thiazide (P), haloperidol (P), interferon alfa (I), MAOIs (P), phenothiazines (P), SSRIs (P), sulfonylureas (P), vasopressin (P), vinca alkaloids (P).
**Phosphate (S).** Elevated by anabolic steroids (P), basiliximab (P), contraceptives, oral (P), foscarnet (P), mannitol (I), methicillin (LP), pindolol (P), rifampin (LP), sodium phenylbutyrate (P), vitamin D (excessive) (P).\(^1,4\)

Decreased by acetazolamide (P), antacids (phosphate binding; eg, aluminum, calcium, and magnesium salts) (P), bisphosphonates (P), calcitonin (P), carbamazepine (P), cidofovir (P), citrate salts (I), foscarnet (P), insulin (P), lithium salts (P), mannitol (I), mycophenolate (P), parenteral nutrition (P), phenobarbital (P), phenothiazines (P), phenytoin (P), sevelamer (P), sirolimus (P), sorbitol (P), sucralfate (P), tacrolimus (P).\(^1,4–6\)

**Potassium (S).** Elevated by aminocaproic acid (P), angiotensin II receptor blockers (P), ACE inhibitors (P), \(\beta\)-adrenergic blockers (P), antineoplastic agents (cytotoxic effect) (P), basiliximab (P), cefotaxime (I), cyclosporine (P), COX-2 inhibitors (P), diuretics, potassium-sparing (P), flunazole (P), fluoride salts (I), heparins (P), iodide salts (I), isoniazid (P), lithium salts (P), low-molecular-weight heparins (P), mannitol (P), mycophenolate (P), nephrotoxic drugs (P), nonsteroidal anti-inflammatory drugs (primarily indomethacin) (P), pentamidine (P), potassium penicillin (P), procainamide (I), salt substitutes (P), succinylcholine (P), tacrolimus (P), trimethoprim (P), tromethamine (P).\(^1–6\)

Decreased by acetazolamide (P), \(\beta\)-adrenergic agonists (P), aminoglycosides (P), ammonium chloride (P), amphotericin B (P), basiliximab (P), bicarbonate salts (P), bisphosphonates (P), cisplatin (P), cyclosporine (P), diuretics (loop, thiazide) (P), fenoldopam (P), foscarnet (P), glucose (P), insulin (P), laxatives (P), levodopa (P), mineralocorticoids (P), mycophenolate (P), ondansetron (P), penicillins (extended-spectrum) (P), phosphate salts (P), salicylates (P), sirolimus (P), sorbitol (P), sodium polystyrene sulfonate (P), sodium phenylbutyrate (P), sulfasalazine (P), tacrolimus (P).\(^1–6\)

**Protein, Total.** Elevated by anabolic steroids \(\{S\}\) (P), aspirin \{CSF\} (I), cephalothin \{S\} (I), chloramphenicol \{S\} (I), corticosteroids \{S\} (P), dextran \{CSF/S\} (I), imipramine \{CSF\} (I), lidocaine \{CSF\} (I), mannitol \{CSF\} (I), methotrexate \{CSF\} (I), penicillins \{S/CSF\} (I), phenazopyridine \{S\} (I), phenothiazines \{CSF\} (I), progestins \{S\} (I), radiopaque agents \{S\} (I), rifampin \{S\} (I), sulfonamides \{CSF\} (I), tetracyclines \{CSF\} (I), thyroid \{S\} (P), vancomycin \{CSF\} (I), vitamin C \{CSF\} (I).\(^1–3\)

Decreased by acetaminophen \{CSF\} (I), cefotaxime \{CSF\} (I), contraceptives, oral (from estrogen) \{S\} (P), cytarabine \{CSF\} (P/I), dexamethasone \{CSF\} (P), dextran \{S\} (I/P), estrogens \{S\} (P), hepatotoxic drugs \{S\} (P), pyrazinamide \{S\} (P), rifampin \{S\} (P).\(^1–3\)

**Sodium (S).** Elevated by anabolic steroids (P), bicarbonate salts (P), carbamazepine (LP), cefotaxime (I), clonidine (P), contraceptives, oral (P), corticosteroids (P), COX-2 inhibitors (P), diazoxide (P), estrogens (P), fluoride salts (I), lactulose (P), mannitol (P), methyldopa (P), mineralocorticoids (P), nitrofurantoin (P), nonsteroidal anti-inflammatory drugs (P), sodium phenylbutyrate (P), tetracycline (P).\(^1,3,5\)

Decreased by acetazolamide (P), ammonium chloride (P), amphotericin B (P), antidepressants, tricyclic (P), bicarbonate salts (I), carbamazepine (P), chlor-
propamide (P), cisplatin (P), clonidine (P), cyclophosphamide (P), cytarabine (P), diuretics, loop and thiazide (P), haloperidol (P), indomethacin (P), interferons (P), laxatives (P), lithium salts (P), mannitol (P), MAOIs inhibitors (P), miconazole (P), nifedipine (P), nonsteroidal anti-inflammatory drugs (P), phenothiazines (P), SSRIs (P), somatostatin (P), spironolactone (P), sulfonureas (P), vasopressin and analogues (P), trimethoprim (P), vinca alkaloids (P).1,3–5,7

**Thyroxine (S).** *Elevated* by amiodarone (LP), clofibrate (P), contraceptives, oral (from estrogen) (P), estrogens (P), fluourouracil (P), heparin (I), insulin (P), levodopa (P), prazosin (P), propranolol (P), propylthiouracil (P), prostaglandins (P), radiocontrast agents (LP), tamoxifen (P).1,3–5

*Decreased* by anabolic steroids (P), asparaginase (P), barbiturates (P), carbamazepine (P), chloropropamide (P), cholestyramine (P), clofibrate (P), colestipol (P), corticosteroids (P), danazol (LP), diazepam (P), heparin (I), interferon alfa-2a (P), iodide salts (P), iron salts (P), lithium salts (P), penicillin (P), phenytoin (P), propylthiouracil (P), reserpine (P), salicylates (P), sulfonamides (P), sulfonyleureas (P), thyroid (P).1–3

**Triglycerides (S).** *Elevated* by β-adrenergic blockers (P), alitretinoin (P), amiodarone (P), amphenavir (P), aspirin (LP), clofibrate (P), contraceptives, oral (P), cyclosporine (P), danazol (P), didanosine (P), diuretics, loop and thiazide (P), estrogens (P), fomepizole (P), interferon alfa-2a (P), isoretinoin (P), itraconazole (P), lipids (P), low-molecular-weight heparins (P), HMG-CoA reductase inhibitors (P), mirtazapine (P), niraglycerin (I), olanzapine (P), protease inhibitors (P), quinine (P), ivermectin (P), sirolimus (P), tamoxifen (P).2–5

*Decreased* by acarbose (P), α-adrenergic blockers (P), amiodarone (P), ACE inhibitors (P), asparaginase (P), aspirin (I), chenodiol (P), citrate salts (I), clofibrate (P), danazol (P), fenofibrate (P), gemfibrozil (P), HMG-CoA reductase inhibitors (P), hydroxychloroquine (P), hydroxyurea (I), ketoconazole (P), metformin (P), methotrexate (I), methylprednisolone (P), metronidazole (I), niacin (P), nifedipine (P), orlistat (P), probucol (P), psyllium (P), rifampin (I), spironolactone (P), sulfonyleureas (P), verapamil (P), vitamin C (LP).1–4

**Urea Nitrogen (S).** *Elevated* by ACE inhibitors (P), acetylsalicylic acid (P), acetohexam ide (I), aminoglycosides (P), anabolic steroids (P), antacids (prolonged use) (P), asparaginase (P), busulfan (P), carbamazepine (P), chloral hydrate (I), chloramphenicol (Nesslerization method) (I), cisplatin (P), clonidine (P), colistin (P), co-trimoxazole (P), cyclosporine (P), dexamethasone (P), dextran (LP), diuretics, loop and thiazide (P), fluoxymesterone (P), gold salts (P), hydralazine (P), hydroxyurea (P), ifosfamide (P), iron salts (P), methotrexate (P), methylprednisolone (P), mithrysergide (P), mitomycin (P), nalidixic acid (P), nephrotoxic drugs (P), nitrofurantoin (P), nonsteroidal anti-inflammatory drugs (P), penicillamine (P), pentamidine (P), radiocontrast agents (P), salicylates (P), sulfonamides (I), tacrolimus (P), tetracyclines (LP), vancomycin (P), vitamin D (P).1,3–5

*Decreased* by amikacin (I), ascorbic acid (I), cefotaxime (I), chloramphenicol (Berthelot method) (I), fluoride salts (I), levodopa (P), phenothiazines (P), streptomycin (I).1,3–5
Uric Acid (S). Elevated by acetaminophen (I), acetzolamide (P), anabolic steroids (P), antineoplastics (P), azathioprine (P), basiliximab (P), caffeine (Bittner method) (I), cisplatin (P), citrate salts (P), cyclosporine (P), cytarabine (P), diazoxide (P), diuretics (carbonic anhydrase inhibitor, loop, thiazide) (P), epinephrine (I), ethambutol (P), filgrastim (P), hydralazine (I), isoniazid (I), levodopa (LP), mercaptopurine (P), methylxanthines (P), pyrazinamide (P), propranolol (P), propylthiouracil (P), rifampin (I), ritonavir (P), salicylates (low doses) (LP), sodium phenylbutyrate (P), spiranolactone (P), tacrolimus (P), theophylline (LP), triamterene (P), vitamin C (I), zalcitabine (P).2–5

Decreased by acetohexamide (P), allopurinol (P), cefotaxime (I), cidofovir (P), clofibrate (P), corticosteroids (P), diflunisal (P), fenofibrate (P), glucose infusions (P), griseofulvin (P), guaifenesin (P), hydralazine (I), indomethacin (P), levodopa (I), lithium (P), losartan (P), mannitol (P), methylxanthines (P), nonsteroidal anti-inflammatory drugs (P), phenothiazines (P), radiocontrast agents (P), salicylates (large doses) (P), spiranolactone (P), sulfonamides (P), uricosurics (eg, probenecid, sulfinpyrazone) (P), verapamil (P), vitamin C (by Seralyzer) (LP).2–5

URINE TESTS

Bilirubin. Elevated by acetohexamide (P), etodolac (I), hepatotoxic drugs (P), mfenamic acid (I), phenazopyridine (I), phenothiazines (I,P).1,5

Catecholamines. Elevated by acetaminophen (I), α1-adrenergic blockers (P), alcohol (P), aspirin (I), atenolol (P), caffeine (P), chloral hydrate (I), chlorpromazine (I), dopamine (I,P), epinephrine (I), erythromycin (I), hydralazine (I), insulin (P), isoproterenol (I), labetalol (I), levodopa (I), methenamine (I), methylxanthines (I), nifedipine (P), nitroglycerin (P), prochlorperazine (P), quinidine (I), reserpine (P), tetracyclines (I,P), triamterene (P).1,2,4,5

Decreased by α2-adrenergic blockers (P), bromocriptine (P), clonidine (P), disulfiram (P), guanethidine (P), methenamine (destroys catecholamines in bladder urine) (P), radiocontrast agents (I), reserpine (P).1,2,4

Color. (See Drug-Induced Discoloration of Feces and Urine, page 1024.)

Creatinine. Elevated by anabolic steroid (increased muscle mass) (P), asparaginase (I), cephalosporins, except cefotaxime and ceftazidime; (Jaffe method) (I), corticosteroids (P), levodopa (I), methotrexate (P), methylxanthines (P), nephrotoxic drugs (P), nitrofurantoin (I), reserpine (P), vitamin C (I).1,4,5

Decreased by anabolic steroids (anabolic effect) (P), captopril (P), cimetidine (P), diuretics, thiazide (P).1,4,5

Glucose. Elevated or False Positive by aspirin (copper reduction) (I,P), aminosalicylic acid (copper reduction) (I), cephalosporins, except cefotaxime (copper reduction) (I), chloral hydrate (copper reduction) (I), cidofovir (P), corticosteroids (P), dextroamphetamine (P), diuretics, loop and thiazide (P), glucagon (P), isoniazid (P), levodopa (copper reduction) (I), lithium salts (P), nalidixic acid (I), niacin (P), penicillins (I), pentamidine (P), phenazopyridine (Tes-Tape) (I), phenothiazines (P), probenecid (I), reserpine (P), sulfonamides (I), vitamin C (copper reduction).1–6
Decreased or False Negative by aspirin (glucose oxidase) (I,P), bisacodyl (I), chloral hydrate (glucose oxidase) (I), diazepam (I), digoxin (I), ferrous salts (I), flurazepam (I), furosemide (P), levodopa (glucose oxidase) (I), phenazopyridine (glucose oxidase) (I), phenobarbital (P), prednisone (glucose oxidase) (I), secobarbital (I), tetracycline (I), vitamin C (glucose oxidase).1–6

Gonadotropins (Pregnancy Test). False Positive by methadone (I), phenothiazines (I).1,5

Ketones. Elevated by acetylcysteine (I), albuterol (P), captopril (I), cephalosporins (I), dimercaprol (I), insulin (P), isoniazid (P), levodopa (Labstix) (I), mesna (I), metformin (I), methylprednisolone (I), phenazopyridine (I), phenothiazines (I), pyrazinamide (I), salicylates (acidotic effect) (I,P), succimer (I), valproic acid (I).1,2,4,5

Decreased by aspirin (oxidation of ketone bodies) (P), phenazopyridine (I).1

Protein. Elevated by acetaminophen (P), acetylsalicylic acid (I,P), aspirin (I), bacitracin (P), biperiden (I), captopril (P), carbamazepine (P), cephalosporins (I,P), chlorpromazine (I), cimetidine (P), ciprofloxacin (I), cytochrome P450 inhibitors (I), dextrose (P), dexamethasone (P), digoxin (I), flurbiprofen (I), furosemide (I), insulin (P), levodopa (P), metformin (I), metoprolol (I), methylprednisolone (I), phenazopyridine (I), phenothiazines (I), pyrazinamide (I), salicylates (acidotic effect) (I,P), succimer (I), valproic acid (I).1

Decreased by acebutolol (P), cyclosporine (P), diltiazem (P), interferon beta-1b (P), iron salts (P), isoniazid (P), lithium salts (P), nephrotoxic drugs (P), nonsteroidal anti-inflammatory drugs (P), penicillin (P), penicillins (I), penicillinase (I), phenazopyridine (I), radiographic agents (I), rifampin (P), salicylates (I), sulfonamides (I), tetracycline (P), tolbutamide (I), vancomycin (P), vitamin C (I).1,5

Specific Gravity. Elevated by dextran (P), diuretics (P), isotretinoin (P), mannitol (P), radiographic agents (P), sucrose (P).1,4,5

Decreased by colistin (P), lithium (P).1

HEMATOLOGY

Erythrocyte Sedimentation Rate (B). Elevated by contraceptives, oral (P), cyclosporine (P), dextran (P), isoniazid (P), methylprednisolone (P), methylprednisolone (P), nitrofurantoin (P), procainamide (P), theophylline (P), vitamin A (P).1,5

Decreased by corticosteroids (P), cyclophosphamide (P), infliximab (P), fluoride salts (I), gold salts (P), methotrexate (P), nonsteroidal anti-inflammatory drugs (P), penicillamine (P), quinine (P), salicylates (P), sulfasalazine (P), tamoxifen (P), trimethoprim (P), drugs that cause hyperglycemia (P).1,5

Prothrombin Time (B) [Does not include anticoagulants or drugs which potentiate or antagonize them]. Elevated by acetaminophen (P), antibiotics (gut sterilizing) (P), asparaginase (P), cephalosporins (P), chloramphenicol (P), chloral hydrate (P), chlorpromazine (P), chlorpropamide (P), cholestyramine (P), colistin (P), cyclophosphamide (P), hepatotoxic drugs (P), laxatives (P), mercaptopurine
(P), metronidazole (P), niacin (P), propylthiouracil (P), quinidine (P), quinine (P), salicylates (P), sulfonamides (P).1,4,5

Decreased by anabolic steroids (P), azathioprine (P), estrogens (P), vitamin K (P).1,3,5

REFERENCES
Abbreviations used in this appendix:

Alb_{meas} = measured serum albumin
Alb_{nl} = normal serum albumin
α = fraction of drug unbound to albumin
C_{adj} = adjusted serum concentration
C_{des} = desired serum concentration
Cl = serum drug clearance
C_{meas} = measured serum concentration
ΔC_p = desired increase in serum concentration
C_{p_t} = serum concentration at time t
C_{ss} = steady-state serum concentration
C_{ss \_ave} = average steady-state serum concentration
C_{ss \_max} = maximum (peak) steady-state serum concentration
C_{ss \_min} = minimum (trough) steady-state serum concentration
C_{0} = initial serum concentration
D = dose
F = fraction of dose absorbed
k_{0} = infusion rate (dose/t_{inf})
ka = absorption rate constant
kd = elimination rate constant
km = Michaelis–Menten constant in mg/L
S = salt fraction
τ = dosage interval in hours
t_{1/2} = elimination half-life
t_{inf} = duration of infusion
t_{max} = time of peak serum concentration
\text{t}_{max \_ss} = time of peak serum concentration at steady state
V_d = apparent volume of distribution
V_m = maximum rate of metabolism in mg/day
ONE-COMPARTMENT EQUATIONS

\[ k_d = \frac{\text{Cl}}{V_d} \]

\[ t_{1/2} = \frac{0.693 \times V_d}{\text{Cl}} \]

\[ k_d = \frac{0.693}{t_{1/2}} \]

\[ k_d = \frac{\ln (C_{p1}/C_{p2})}{t} \]

Where \( t \) = time between serum concentrations \( C_{p1} \) and \( C_{p2} \).

SINGLE-DOSE EQUATIONS

Concentration at time \( t \) (IV bolus):

\[ C_{p_t} = \frac{S \times D}{V_d} \times e^{-k_d \times t} \]

Concentration at time \( t \) (during IV infusion):

\[ C_{p_t} = \frac{S \times k_0}{\text{Cl}} \times (1 - e^{-k_d \times t_{\text{inf}}}) \]

Concentration at time \( t \) (after the end of an IV infusion):

\[ C_{p_t} = \frac{S \times k_0}{\text{Cl}} \times (1 - e^{-k_d \times t_{\text{inf}}}) \times e^{(-k_d \times t_{\text{inf}})} \]

Concentration at time \( t \) (PO or IM):

\[ C_{p_t} = \frac{S \times F \times D \times k_a}{V_d \times (k_a - k_d)} \times (e^{-k_d \times t} - e^{-k_a \times t}) \]

Time to Peak (PO or IM):

\[ t_{\text{max}} = \frac{\ln (k_a/k_d)}{k_a - k_d} \]

Loading Dose (negligible drug loss during administration):

\[ LD = \frac{\Delta C_p \times V_d}{S \times F} \]
PHARMACOKINETIC EQUATIONS

Loading Dose (IV loading dose when drug is lost during administration):

\[
LD = \frac{\text{Cl} \times t_{\text{inf}} \times (C_{\text{des}} - [C_{\min} \times e^{-kd \times t_{\text{inf}}}] )}{S \times (1 - e^{-kd \times t_{\text{inf}}})}
\]

Loading Dose (IM or PO):

\[
LD = \frac{\text{Vd} \times (C_{\text{des}} - [C_{\min} \times e^{-kd \times t_{\text{max}}}] )}{S \times F \times (1 - e^{-kd \times t_{\text{max}}})}
\]

STEADY-STATE EQUATIONS

Peak Concentration (IV):

\[
C_{ss \max} = S \times k_0 \times \frac{1 - e^{-kd \times t_{\text{inf}}}}{\text{Cl} \times (1 - e^{-kd \times \text{t}})}
\]

Peak Concentration (PO or IM):

\[
C_{ss \max} = \frac{S \times F \times D \times e^{-kd \times t_{\text{max ss}}}}{\text{Vd} \times (1 - e^{-kd \times \text{t}})}
\]

Trough Concentration (IV):

\[
C_{ss \min} = C_{ss \max} \times e^{-kd \times (\text{t} - t_{\text{inf}})}
\]

Trough Concentration (PO or IM):

\[
C_{ss \min} = \frac{S \times F \times D \times ka}{\text{Vd} \times (ka - kd)} \times \left[ \frac{e^{-kd \times \text{t}}}{1 - e^{-kd \times \text{t}}} - \frac{e^{-ka \times \text{t}}}{1 - e^{-ka \times \text{t}}} \right]
\]

Average Concentration (IV, PO, or IM):

\[
C_{ss \text{ ave}} = \frac{S \times F \times D}{\text{Cl} \times \text{t}}
\]

Time to Peak (PO or IM):

\[
t_{\text{max ss}} = \frac{\ln (ka \times (1 - e^{-kd \times \text{t}})) / (kd \times (1 - e^{-ka \times \text{t}})})}{ka - kd}
\]

Dosage Interval:

\[
\text{Interval} = T + \frac{(\ln \left[ \frac{C_{\max}}{C_{\min}} \right])}{kd}
\]

where T = infusion time for IV doses and \( \text{t}_{\text{max}} \) for PO and IM doses.
MICHAELIS–MENTEN EQUATIONS

\[
\text{Daily Dosage} = \frac{V_m \times C_{\text{ss,ave}}}{S \times F \times (\text{km} + C_{\text{ss,ave}})}
\]

\[
C_{\text{adj}} = \frac{C_{\text{meas}}}{[1 - \alpha] \times \left(\frac{\text{Alb}_{\text{meas}}}{\text{Alb}_{\text{al}}}\right) + \alpha}
\]

\[
C_{\text{ss,ave}} = \frac{S \times F \times \text{Dosage/day} \times \text{km}}{V_m - (S \times F \times \text{daily dosage})}
\]
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USE OF THE INDEX. Index terms in bold are US Nonproprietary Names (ie, generic names). Nonbolded terms are US Proprietary Names (ie, trade names), unless they are designated as (BAN) which indicates the British Approved Name (ie, generic name) or (Can) which indicates a Canadian Brand Name. Terms in small capital letters are DRUG CLASSES. Bold page numbers indicate the page(s) of a drug monograph. Page numbers in italics indicate the page(s) of a comparison chart. To ensure location of complete information, check a drug’s entries for both its Nonproprietary Name and its DRUG CLASS.

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### ACRONYMS AND ABBREVIATIONS USED IN THE BOOK

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<tbody>
<tr>
<td>ACE</td>
<td>angiotensin converting enzyme</td>
</tr>
<tr>
<td>ALL</td>
<td>acute lymphocytic leukemia</td>
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<tr>
<td>ALT</td>
<td>alanine aminotransferase</td>
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<tr>
<td>AML</td>
<td>acute myelocytic leukemia</td>
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<tr>
<td>ANA</td>
<td>antinuclear antibody</td>
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<tr>
<td>ANC</td>
<td>absolute neutrophil count</td>
</tr>
<tr>
<td>aPTT</td>
<td>activated partial thromboplastin time</td>
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<tr>
<td>AST</td>
<td>aspartate aminotransferase</td>
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<tr>
<td>AUC</td>
<td>area under the serum concentration-time curve</td>
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<tr>
<td>AV</td>
<td>atrioventricular</td>
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<tr>
<td>bid</td>
<td>twice daily</td>
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<tr>
<td>BP</td>
<td>blood pressure</td>
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<td>BPH</td>
<td>benign prostatic hypertrophy</td>
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<tr>
<td>BSA</td>
<td>body surface area</td>
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<tr>
<td>BUN</td>
<td>blood (serum) urea nitrogen</td>
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<td>Cap</td>
<td>capsule</td>
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<tr>
<td>CBC</td>
<td>compete blood (cell) count</td>
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<tr>
<td>Chew</td>
<td>chewable</td>
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<td>CHD</td>
<td>coronary heart disease</td>
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<td>congestive heart failure</td>
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<td>CI</td>
<td>clearance</td>
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<tr>
<td>Clr</td>
<td>creatinine clearance</td>
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<td>chronic lymphoblastic leukemia</td>
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<tr>
<td>CML</td>
<td>chronic myelocytic leukemia</td>
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<td>CMV</td>
<td>cytomegalovirus</td>
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<td>CNS</td>
<td>central nervous system</td>
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<tr>
<td>COMT</td>
<td>catechol-O-methyltransferase</td>
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<td>chronic obstructive pulmonary disease</td>
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<td>CPK</td>
<td>creatine phosphokinase</td>
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<td>Crm</td>
<td>cream</td>
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<tr>
<td>Cre</td>
<td>serum creatinine concentration</td>
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<tr>
<td>CSF</td>
<td>cerebrospinal fluid</td>
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<td>CVA</td>
<td>cerebrovascular accident</td>
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<td>CVP</td>
<td>central venous pressure</td>
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<tr>
<td>CYP</td>
<td>cytochrome P450</td>
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<td>D5W</td>
<td>5% dextrose solution</td>
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<td>DHL</td>
<td>diffuse histiocytic lymphoma</td>
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<td>DMSO</td>
<td>dimethyl sulfoxide</td>
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<td>Drp</td>
<td>drop(s)</td>
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<td>DVT</td>
<td>deep vein thrombosis</td>
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<td>EC</td>
<td>enteric coated</td>
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<td>EC50</td>
<td>mean effective concentration</td>
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<td>ECG</td>
<td>electrocardiogram</td>
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<td>Elxr</td>
<td>elixir</td>
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<td>FDA</td>
<td>U.S. Food &amp; Drug Administration</td>
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<td>G-6-PD</td>
<td>glucose-6-phosphate dehydrogenase</td>
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<td>GABA</td>
<td>γ-aminobutyric acid</td>
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<tr>
<td>GC-MS</td>
<td>gas chromatography-mass spectroscopy</td>
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<tr>
<td>GFR</td>
<td>glomerular filtration rate</td>
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<td>GI</td>
<td>gastrointestinal</td>
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<td>Gran</td>
<td>granules</td>
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<td>GU</td>
<td>genitourinary</td>
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<tr>
<td>GVHD</td>
<td>graft versus host disease</td>
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<tr>
<td>Hb</td>
<td>hemoglobin</td>
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<tr>
<td>HDL</td>
<td>high-density lipoprotein</td>
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<td>HDLc</td>
<td>high-density lipoprotein cholesterol</td>
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<td>HIC</td>
<td>mean inhibitory concentration</td>
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<td>IgG</td>
<td>immunoglobulin G</td>
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<td>IM</td>
<td>intramuscular(ly)</td>
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<td>Inhal</td>
<td>inhalation</td>
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<td>Inj</td>
<td>injection</td>
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<td>INR</td>
<td>international normalized ratio</td>
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<td>intraperitoneal(ly)</td>
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<td>IT</td>
<td>intrathecal(ly)</td>
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<td>IU</td>
<td>international units</td>
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<td>IUD</td>
<td>intrauterine device</td>
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<td>LBW</td>
<td>lean body weight</td>
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<td>ACRONYMS AND ABBREVIATIONS (cont’d)</td>
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<td>LDH</td>
<td>lactate dehydrogenase</td>
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<td>LDL</td>
<td>low-density lipoprotein</td>
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<tr>
<td>LDLc</td>
<td>low-density lipoprotein cholesterol</td>
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<tr>
<td>LFT</td>
<td>liver function test</td>
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<tr>
<td>Lot</td>
<td>lotion</td>
</tr>
<tr>
<td>LR</td>
<td>lactated Ringer’s solution</td>
</tr>
<tr>
<td>MAO</td>
<td>monoamine oxidase</td>
</tr>
<tr>
<td>MAOI</td>
<td>monoamine oxidase inhibitor</td>
</tr>
<tr>
<td>MAP</td>
<td>mean arterial pressure</td>
</tr>
<tr>
<td>MCH</td>
<td>mean corpuscular hemoglobin</td>
</tr>
<tr>
<td>MCV</td>
<td>mean corpuscular volume</td>
</tr>
<tr>
<td>MI</td>
<td>myocardial infarction</td>
</tr>
<tr>
<td>MIC</td>
<td>minimum inhibitory concentration</td>
</tr>
<tr>
<td>mol</td>
<td>mole(s)</td>
</tr>
<tr>
<td>NG</td>
<td>nasogastric</td>
</tr>
<tr>
<td>NPO</td>
<td>nothing by mouth, fasting</td>
</tr>
<tr>
<td>NS</td>
<td>normal saline, 0.9% NaCl solution</td>
</tr>
<tr>
<td>NSAID</td>
<td>nonsteroidal anti-inflammatory drug</td>
</tr>
<tr>
<td>OCD</td>
<td>obsessive-compulsive disorder</td>
</tr>
<tr>
<td>Oint</td>
<td>ointment</td>
</tr>
<tr>
<td>Ophth</td>
<td>ophthalmic</td>
</tr>
<tr>
<td>OTC</td>
<td>over the counter (nonprescription)</td>
</tr>
<tr>
<td>PaO₂</td>
<td>partial alveolar oxygen</td>
</tr>
<tr>
<td>PCA</td>
<td>patient-controlled analgesia</td>
</tr>
<tr>
<td>PCP</td>
<td><em>Pneumocystis carinii</em> pneumonia</td>
</tr>
<tr>
<td>PE</td>
<td>pulmonary embolism</td>
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<tr>
<td>PN</td>
<td>parenteral nutrition</td>
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<tr>
<td>PO</td>
<td>oral(ly)</td>
</tr>
<tr>
<td>PR</td>
<td>rectal(ly)</td>
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<tr>
<td>PSA</td>
<td>prostate-specific antigen</td>
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<tr>
<td>PT</td>
<td>prothrombin time</td>
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<tr>
<td>PTT</td>
<td>partial thromboplastin time</td>
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<tr>
<td>PVC</td>
<td>premature ventricular contraction</td>
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<tr>
<td>Pwdr</td>
<td>powder</td>
</tr>
<tr>
<td>q</td>
<td>every</td>
</tr>
<tr>
<td>qd</td>
<td>daily; every day</td>
</tr>
<tr>
<td>qid</td>
<td>four times daily</td>
</tr>
<tr>
<td>RBC</td>
<td>red blood cell</td>
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<tr>
<td>RDA</td>
<td>recommended dietary allowance</td>
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<td>RR</td>
<td>respiratory rate</td>
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<tr>
<td>SA</td>
<td>sinoatrial</td>
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<tr>
<td>SC</td>
<td>subcutaneous(ly)</td>
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<tr>
<td>SIADH</td>
<td>syndrome of inappropriate antidiuretic hormone</td>
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<tr>
<td>SL</td>
<td>sublingual(ly)</td>
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<tr>
<td>SLE</td>
<td>systemic lupus erythematosus</td>
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<tr>
<td>Soln</td>
<td>solution</td>
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<tr>
<td>SR</td>
<td>sustained-release</td>
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<tr>
<td>SSRI</td>
<td>selective serotonin reuptake inhibitor</td>
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<tr>
<td>Supp</td>
<td>suppository</td>
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<td>Susp</td>
<td>suspension</td>
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<td>tablet</td>
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<tr>
<td>TBW</td>
<td>total body weight</td>
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<tr>
<td>TIA</td>
<td>transient ischemic attack</td>
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<tr>
<td>tid</td>
<td>three times daily</td>
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<tr>
<td>Top</td>
<td>topical(ly)</td>
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<tr>
<td>TSH</td>
<td>thyroid-stimulating hormone</td>
</tr>
<tr>
<td>UTI</td>
<td>urinary tract infection</td>
</tr>
<tr>
<td>Vag</td>
<td>vaginal(ly)</td>
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<tr>
<td>Vₐₑ</td>
<td>apparent volume of distribution of the central compartment</td>
</tr>
<tr>
<td>Vₐₚₑ</td>
<td>apparent volume of distribution (one-compartment)</td>
</tr>
<tr>
<td>Vₑₚ</td>
<td>apparent volume of distribution of the β phase</td>
</tr>
<tr>
<td>Vₛₚₑ</td>
<td>steady-state apparent volume of distribution</td>
</tr>
<tr>
<td>VLDL</td>
<td>very low-density lipoprotein</td>
</tr>
<tr>
<td>WBC</td>
<td>white blood cell</td>
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</table>