

CURRENT REPORT—HIV

Treatment of AIDS and HIV-Related Conditions—1999

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During the last 2 years human immunodeficiency virus (HIV) disease and acquired immunodeficiency syndrome (AIDS) have been transformed from progressive, fatal diseases to ones in which the disease process can be improved. The striking effectiveness of combination antiretroviral therapy, especially regimens that include the protease inhibitor class of drugs, is largely responsible for this dramatic turnabout. Although some HIV-infected patients do not respond favorably to these therapies, and resistance to combination therapy can occur with time, there is now hope that if antiretroviral therapies continue to improve, HIV disease might eventually become another treatable chronic disease.

This Current Report — HIV updates our annual treatment guidelines.¹ It is based on a review of the medical literature, our experience at San Francisco General Hospital, and experience gained from answering calls to our National HIV Telephone Consultation Service (Warmline). These guidelines are intended to provide treatment recommendations for antiretroviral therapy and associated medical problems of adults and adolescents with HIV disease and AIDS.

Antiretroviral Therapy

Guidelines for antiretroviral therapy are readily available. The United States Public Health Service publishes guidelines² that are updated continually at <http://www.cdc.gov> or <http://hivatis.org>. Recommendations from the International AIDS Soci-

ety,³ the British HIV Association,⁴ and related articles add to the range of approaches to antiretroviral therapy.⁵⁻⁸

Monitoring HIV Progression and Antiretroviral Therapy

Refinements in viral load testing have revolutionized HIV care. The ultrasensitive assays of HIV RNA (viral load) levels can detect fewer than 50 or even 20 copies/mL, allowing the clinician and patient to estimate activity of HIV disease and the degree of effectiveness of antiretroviral therapy. Although suppression of the viral load to less than the detectable level of the assay is the optimal goal of therapy, it is not always possible. Viral loads of less than about 500 copies/mL are desirable. Decreases in viral load generally occur within 1 month after therapy begins, but maximal suppression can take up to 4 to 6 months of therapy. Antiretroviral therapy that does not suppress the viral load by at least 1 log (10-fold) should be considered ineffective.

The CD4+ lymphocyte count remains an important surrogate marker in assessing HIV disease. Opportunistic infections rarely occur in patients with a CD4+ cell count of more than 200/ μ L, and the conditions associated with severe disease generally occur when the CD4+ cell count is less than 50 to 100/ μ L. The CD4+ count usually rises with effective antiretroviral therapy.

Antiretroviral Drugs

The three principal classes of antiretroviral drugs (Table 1) are the nucleoside reverse transcriptase inhibitors (NRTIs), the protease inhibitors, and the nonnucleoside reverse transcriptase inhibitors (NNRTIs). Combination antiretroviral therapy is usually based upon a foundation of two NRTIs plus protease inhibitor therapy. Monotherapy is never recommended because drug resistance will develop rapidly.

The most common combinations of NRTIs are zidovudine plus lamivudine, didanosine, or

Submitted, revised, 3 September 1998.

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Supported in part by the Pacific AIDS Education and Training Center, Grant No. 2 U69 PE00118-05, with the Bureau of Health Professions, Health Resources and Services Administration, Department of Health and Human Services.

zalcitabine; and stavudine plus lamivudine or didanosine. Zidovudine and stavudine should not be used together because antagonism can occur. Didanosine and zalcitabine should not be used together because of additive toxicity.

The protease inhibitor class of drugs⁹ is most potent in decreasing viral load. When used in the absence of effective NRTI therapy, however, resistance usually occurs, rendering the protease inhibitors ineffective. Cross-resistance among protease inhibitors is common; when resistance develops to one protease inhibitor, resistance to other protease inhibitors can occur. Cross-resistance between ritonavir and indinavir is nearly 100 percent. Combination ritonavir plus saquinavir therapy can offer some benefits in persons who have protease inhibitor resistance to one other protease inhibitor. Protease inhibitors are associated with substantial side effects and important drug interactions, as listed in Table 1.

The NNRTIs can also be potent inhibitors of HIV replication. Without concomitant and effective NRTI therapy, however, the virus will rapidly develop complete resistance to NNRTI therapy. NNRTI resistance occurs across this class of drugs, so that resistance to one NNRTI results in resistance to the others. Currently, the most common role of the NNRTIs is to substitute for a protease inhibitor when toxicity or resistance to protease inhibitor therapy has occurred, but some providers use a combination of NRTIs plus protease inhibitor plus NNRTI as initial therapy or even NRTIs plus NNRTI as initial therapy. Long-term studies are needed to evaluate these strategies.

Hydroxyurea¹⁰ has been used in combination with didanosine with apparent benefit in both early HIV infection and in salvage therapy. Whether hydroxyurea has long-term benefits and whether it is effective in combination with other antiretroviral agents remains unknown. The role of the nucleotide analog adefovir is unclear at this time. Adefovir has been used with apparent effectiveness as an additional agent in salvage therapy in patients who have failed other regimens.

Pseudo-Cushing syndrome, which includes the development of "buffalo hump," abnormal fat deposition, systemic dyslipidemias, and glucose intolerance and insulin resistance, has been described as a side effect of effective antiretroviral therapy.^{11,12} These complications occur more commonly with combination therapy that includes protease in-

hibitors than with other combination regimens, but the syndrome also can occur in the absence of protease inhibitors. The pathophysiology of these side effects of therapy is not understood, although various hypotheses have been suggested.^{11,13} It is possible that the renourishment of patients that results from effective combination antiretroviral therapy might be a cause of this syndrome.¹¹ These abnormalities are important for both cosmetic and possible long-term cardiovascular reasons.

Initiating Antiretroviral Therapy

The optimal time to initiate antiretroviral therapy remains controversial.²⁻⁴ The prevailing philosophy in the United States is to begin highly potent combination (three-drug) antiretroviral therapy as early as possible for patients with detectable virus who are prepared to commit to a lifetime of antiretroviral therapy. Reducing viral burden results in fewer infected cells, less immune system destruction, and decreased disease progression. By maximally suppressing viral replication, it is hoped that resistance to antiretroviral drugs will also be minimized. The Public Health Service recommends beginning therapy for patients with a CD4⁺ cell count of fewer than 500/ μ L or HIV RNA levels of greater than 10,000 copies/mL (bDNA assay) or 20,000 copies/mL (RT-PCR assay).² The International AIDS Society uses a threshold of 5000 to 10,000 copies/mL.³ The British HIV Association Guidelines emphasize that therapy should be started before irreversible damage of the immune system has occurred, specifically at a CD4⁺ cell count of more than 350/ μ L.⁴

Adherence

Close adherence to antiretroviral drug regimens is notoriously difficult, yet critical to treatment success. Interruption of therapy can result not only in ineffective therapy but in the rapid induction of drug resistance. Failure of early treatment can result in the emergence of a virus that is resistant not only to the drugs being taken but to that entire class of drugs. Second- or third-choice (salvage) regimens are generally less effective than initial regimens. The patient who is not prepared or cannot adhere to lifelong complicated medical regimens could be harmed, rather than helped, by initiation of combination antiretroviral therapy,¹⁴ a fact that should weigh heavily on primary clinicians when recommending antiretroviral therapy,

especially for asymptomatic persons who have a low probability of developing AIDS within a few years. Although it is difficult for the clinician to withhold potentially beneficial therapy, in some cases this strategy can be the most prudent until the clinician and patient believe regimen adherence can be maintained.

Changing Antiretroviral Regimens

Antiretroviral regimens are considered to be failed regimens if patients develop progressive opportunistic conditions of advancing AIDS, when the viral load increases substantially (at least 0.5 log), or when the CD4+ count falls. When the viral load has been undetectable, development of new detectable virus in the serum is also considered to be a sign of failed therapy. Long-term studies will be needed to determine the proper time to change therapy, as clinical improvement sometimes can be sustained beyond the time when surrogate markers (viral load and CD4+ count) deteriorate.

When changing antiretroviral drugs, at least two drugs should be changed. Optimally, an entirely new regimen with at least three new drugs is used.²⁻⁴ When a protease inhibitor-containing regimen has failed, changing from that protease inhibitor to NNRTI therapy or possibly to combination protease inhibitor therapy (in addition to changing two NRTIs) is needed. Changing between ritonavir and indinavir and between the NNRTIs should be avoided because of cross-resistance.

Effects of Antiretroviral Therapy on AIDS-Related Conditions

Effective antiretroviral therapy can have profound effects on AIDS-related conditions as well as the progression of HIV disease. Both the incidence and the natural history of such conditions as *Pneumocystis carinii* pneumonia, Kaposi sarcoma, cytomegalovirus disease, progressive multifocal leukoencephalopathy, mycobacterial disease, and AIDS dementia have been altered favorably by antiretroviral therapy.¹⁵ With reconstitution of elements of the immune system, however, there are also reports of flare-ups of underlying disease¹⁶ as the improved immune system stimulates a more vigorous response against these infections than had occurred during the more immune-depleted state. Although these flare-ups, such as retinitis and inflammatory responses to indolent mycobacterial

disease, are usually temporary, clinicians will need to be alert to these unexpected manifestations.

Prophylaxis against opportunistic infections¹⁷ remains a mainstay in HIV primary care. Most important is primary prophylaxis against *P. carinii* pneumonia, which is recommended for all HIV-infected persons with symptomatic HIV disease or AIDS, including those with CD4+ cell counts of less than 200/ μ L. Prophylaxis against *Mycobacterium avium* complex (MAC) disease has been recommended after CD4+ cell counts decrease to less than 50/ μ L. Primary prophylaxis against candidal and other fungal diseases, herpes infections, and cytomegalovirus disease is not routinely recommended.

CD4+ count thresholds for opportunistic infection prophylaxis are based on the CD4+ count nadir. For patients whose CD4+ counts rise above those thresholds after initiation of antiretroviral therapy, prophylaxis against opportunistic infections usually should be continued. Studies are in progress to determine whether less intensive prophylaxis regimens might be adequate in persons receiving effective antiretroviral therapy. Some studies indicate these strategies can be effective.^{18,19} Effective antiretroviral therapy will ultimately result in changes in recommendations for both primary prophylaxis and maintenance therapy after treatment of acute infections, but until more definitive long-term studies show that prophylaxis regimens can be altered, current recommendations should remain in effect.

Tuberculosis

Tuberculosis prophylaxis is indicated for HIV-infected persons who have an induration of 5 mm or greater on the standard tuberculin skin test. Isoniazid prophylaxis does not require changing antiretroviral regimens, but the shorter rifampin- or rifabutin-based regimens require special drug-interaction considerations. In addition, treatment of active tuberculosis will result in potential drug interactions with antiretroviral medications. Consultation with a tuberculosis expert and adherence to new guidelines²⁰ can avoid serious treatment problems.

The Table

Table 1 gives our recommendations for treating specific diseases and the major symptoms of HIV infection and AIDS. The recommendations are

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Table 1. Treatment Regimens for HIV Disease**General/Systemic p. 74****Ophthalmologic p. 83****Gastrointestinal p. 86****Skin/Mucocutaneous p. 81****Oral Cavity p. 84****Pulmonary p. 87****Hematologic p. 83****Esophageal p. 85****Central Nervous System p. 90**

System, Problem, and Drug Regimen	Duration	Adverse Effects/Drug Interactions	Comments
GENERAL/SYSTEMIC			
Antiretroviral (Anti-HIV)			
<i>Combination therapies</i>	<i>Combination therapy is always recommended, as monotherapy is ineffective and promotes drug resistance. Most common regimens include two nucleoside reverse transcriptase inhibitors (NRTIs) along with protease inhibitor (PI) therapy. Nonnucleoside reverse transcriptase inhibitor (NNRTI) therapy can substitute for protease inhibitor therapy for patients who cannot take or are resistant to protease inhibitor therapy, or can be added to NRTI plus PI regimens when necessary. Cross-resistance among PIs is common, as is cross-resistance among NNRTIs. NRTI combinations are: zidovudine plus lamivudine, didanosine, or zalcitabine; or stavudine plus lamivudine or didanosine. Zidovudine and stavudine should not be used in combination. PI therapy is usually with nelfinavir, indinavir, ritonavir, saquinavir soft-gel capsules, or ritonavir plus saquinavir. Indinavir and saquinavir should not be used in combination. See text for further discussion.</i>		
<i>Nucleoside reverse transcriptase inhibitors (NRTIs)</i>			
Zidovudine (AZT, Retrovir) 200 mg po tid or 300 mg po bid; lower dosages (eg, 100 mg 3 times daily) for patients unable to tolerate higher dosages and patients with renal failure or cirrhosis. Available as liquid formulation. Available also as fixed-dose combination (Combivir) consisting of zidovudine (300 mg) with lamivudine (150 mg) given as 1 capsule bid	Until efficacy wanes or toxicity occurs	Malaise, headache, nausea, insomnia, seizures, myalgias. Anemia, granulocytopenia, thrombocytopenia; macrocytosis is an expected effect of zidovudine therapy and requires no intervention. Toxic myopathy (with elevated creatine phosphokinase [CPK]) with long-term use. Lactic acidosis. Aminotransferase elevations (alanine transaminase [ALT], aspartate transaminase [AST]), hepatomegaly with steatosis. Blue to black discoloration of nails and skin in pigmented races <i>Drug interactions</i> Careful monitoring required when used with other myelosuppressive drugs (ie, trimethoprim-sulfamethoxazole, ganciclovir). Probenecid can increase levels of zidovudine. Acetaminophen (Tylenol) administration does not increase zidovudine toxicity	Monitor for signs of zidovudine toxicity and reduce dosage if required. Transfusions or erythropoietin (if endogenous erythropoietin level < 500 IU/L) therapy can be used if anemia (eg, hemoglobin < 8.0 g/dL) occurs in patients who require zidovudine therapy. Decrease dosage or interrupt for absolute neutrophil count (ANC) < 500/ μ L; consider granulocyte colony-stimulating factor (G-CSF). Transfusions and erythropoietin and G-CSF therapies are expensive; changing to alternate agent preferred High-dosage (1200 mg po qd) zidovudine therapy can be considered for HIV dementia and thrombocytopenia. Toxicity of high-dosage zidovudine can be substantial
Didanosine (ddI, Videx) 200 mg po bid as 2 100-mg tablets or 250 mg po bid powder for patients > 60 kg; 125 mg (tablets) or 167 mg (powder) po bid for patients < 60 kg. Once-daily dosing (300 mg po qd) under investigation. Dosage reduction (ie, 200 mg/d) in renal failure	Until efficacy wanes or toxicity occurs	Pancreatitis; painful peripheral neuropathy (dosage related, reversible); nausea, abdominal cramps, diarrhea related to antacid in formulation; rash; hyperglycemia; hyperuricemia; aminotransferase elevations; headache, insomnia, seizures; elevated triglyceride and amylase levels; thrombocytopenia; retinal atrophy <i>Drug interactions</i> Concomitant administration of H ₂ antagonists, antacids, and omeprazole (Prilosec) and other proton pump inhibitors can increase didanosine absorption, resulting in toxicity. Avoid alcohol and other pancreatic toxins (eg, systemic pentamidine). Avoid concomitant neurotoxic drugs (eg, zalcitabine, vinca alkaloids, oral ganciclovir). Decreases absorption of drugs whose absorption is impaired by buffered products (eg, ketoconazole, itraconazole, indinavir, delavirdine, ritonavir, tetracyclines, quinolone antibiotics). Oral and intravenous ganciclovir increase didanosine toxicity	Monitor for signs of neuropathy. Two tablets must be given per dose to provide adequate buffer for absorption. Can be difficult to chew; tablets do not dissolve readily in water, can be crushed manually Administer didanosine on empty stomach 2 hours apart from antacids, H ₂ antagonists, and drugs (eg, ketoconazole, itraconazole, indinavir, ritonavir, tetracyclines, delavirdine, quinolone antibiotics) whose absorption is impaired by buffered products

Table 1. Continued

System, Problem, and Drug Regimen	Duration	Adverse Effects/Drug Interactions	Comments
GENERAL/SYSTEMIC			
Antiretroviral (Anti-HIV) (cont.)			
Zalcitabine (ddC, Hivid) 0.75 mg po tid; 0.375 mg po tid for patients < 30 kg. Dosage reduction in renal failure	Until efficacy wanes or toxicity occurs	Painful peripheral neuropathy (dosage related, reversible); rash; stomatitis, aphthous ulcers; pancreatitis; esophageal ulceration; seizures; aminotransferase elevations; cardiomyopathy <i>Drug interactions</i> Avoid alcohol and other pancreatic toxins (eg, systemic pentamidine). Avoid concomitant neurotoxic drugs (eg, didanosine, stavudine, isoniazid, vinca alkaloids, oral ganciclovir)	Zalcitabine might be less potent than other NRTIs
Stavudine (d4T, Zerit) 20 - 40 mg po bid for patients > 60 kg; 15 - 30 mg po bid for patients 40 - 60 kg; reduce dosage for patients < 40 kg and for patients with renal failure. Available as liquid formulation	Until efficacy wanes or toxicity occurs	Painful peripheral neuropathy; aminotransferase elevations; anemia, macrocytosis; psychological disturbances, insomnia, anxiety, panic attacks <i>Drug interactions</i> Avoid concomitant use of drugs that can cause neurotoxicity or pancreatic toxicity. See didanosine	Dosage range in this table is lower than standard dosage (40 mg po bid), as studies suggest these lower dosages are associated with equivalent efficacy and a lower incidence of peripheral neuropathy Do not use in combination with zidovudine
Lamivudine (3TC, Epivir) 150 mg po bid; 2 mg/kg po bid for patients < 50 kg. Dosage reduction in renal failure. Available as liquid formulation. Available also as fixed dose combination (Combivir) consisting of zidovudine (300 mg) with lamivudine (150 mg) given as 1 capsule bid	Until efficacy wanes or toxicity occurs	Headache, fatigue, insomnia; peripheral neuropathy, muscle aches; rash; aphthous ulcers; nausea; rare neutropenia, thrombocytopenia; paronychia	Provides some efficacy against hepatitis B. Once-daily dosing (300 mg po qd) under investigation
Abacavir (1592U89, Ziagen) 300 mg po bid with or without food	Until efficacy wanes or toxicity occurs	Nausea, headache, malaise; abdominal pain, diarrhea, rash. Hypersensitivity reaction (2% - 5%, usually in first 4 weeks): flu-like symptoms, fever, malaise, abdominal cramping, nausea, vomiting, diarrhea, morbilliform rash, elevations in transaminases and creatine kinase levels. Symptoms resolve if drug stopped. Do not rechallenge, as anaphylactic reactions and deaths reported	
Protease inhibitors (PIs)			
Nelfinavir (Viracept) 750 mg po tid with meals or 1250 mg po bid with meals. Available as liquid formulation	Until efficacy wanes or toxicity occurs	Diarrhea; hypertriglyceridemia; hypercholesterolemia, abnormal fat accumulation, hyperglycemia <i>Drug interactions</i> P-450 enzyme inhibitor. Avoid concomitant use with rifampin, rifabutin (or decrease rifabutin dosage to 150 mg po qd), astemizole, terfenadine, and cisapride. Benzodiazepine interactions under investigation. Avoid simvastatin or lovastatin because of rhabdomyolysis; can use pravastatin or fluvastatin	Resistance develops slowly; resistant strains might be sensitive to other PIs

Continued

Table 1. Continued

System, Problem, and Drug Regimen	Duration	Adverse Effects/Drug Interactions	Comments
GENERAL/SYSTEMIC Antiretroviral (Anti-HIV) (cont.)			
Indinavir (Crixivan) 800 mg po q 8 h on empty stomach or with skim milk, juice, coffee, tea, toast; dosage adjustment to 600 mg po q 8 h in hepatic disease	Until efficacy wanes or toxicity occurs	Nephrolithiasis, crystalluria, interstitial nephritis; hypertriglyceridemia, hypercholesterolemia, abnormal fat accumulation, hyperglycemia; nausea, vomiting, diarrhea, abdominal pain; asymptomatic hyperbilirubinemia, aminotransferase elevations; rash; insomnia, headache, dizziness, taste disturbances; thrombocytopenia <i>Drug interactions</i> Avoid concomitant use of indinavir with rifampin, rifabutin (or decrease rifabutin dosage to 150 mg po qod), astemizole (Hismanal), cisapride (Propulsid), triazolam (Halcion), or midazolam (Versed). Decrease indinavir dosage to 600 mg po q 8 h when given with ketoconazole or delavirdine. Indinavir administration must be at least 1 hour apart from didanosine or antacid administration. Avoid simvastatin or lovastatin because of rhabdomyolysis; can use pravastatin or fluvastatin	Take with at least 6 glasses of noncaffeinated liquid daily to avoid nephrolithiasis Must be taken every 8 hours, not 3 times daily
Ritonavir (Norvir) 600 mg po bid with meals; can increase from 300 mg po bid to 600 mg po bid over 4 - 7 days to minimize gastrointestinal symptoms. Available as liquid formulation. Combination PI therapy with saquinavir permits lower dosages of each (see below)	Until efficacy wanes or toxicity occurs	Nausea, vomiting, diarrhea, anorexia in more than 50% of patients; aminotransferase elevations; hypertriglyceridemia, hypercholesterolemia, abnormal fat accumulation, hyperglycemia; fatigue, weakness, headache, dizziness, circumoral paresthesias; hyperuricemia, increased creatine kinase; taste disturbances <i>Drug interactions</i> Potent hepatic P-450 enzyme inhibitor. Avoid concomitant use with rifampin, rifabutin (or decrease rifabutin dosage to 150 mg po qod), astemizole, cisapride, and benzodiazepines except lorazepam and temazepam. Dosages of desipramine and other antidepressants, narcotics, and oral contraceptives might need adjustment. Avoid simvastatin or lovastatin because of rhabdomyolysis; can use pravastatin or fluvastatin	Capsules must be refrigerated; solution is stable at room temperature for 30 days
Saquinavir soft-gel capsules (Fortovase) 1200 mg po tid with meals Combination PI therapy with zidovudine permits lower dosages of each (see below)	Until efficacy wanes or toxicity occurs	Headache, confusion; hypertriglyceridemia, hypercholesterolemia, abnormal fat accumulation, hyperglycemia; nausea, diarrhea, abdominal pain; fever <i>Drug interactions</i> Ketoconazole, zalcitabine, delavirdine, and grapefruit juice increase saquinavir serum concentration. Avoid concomitant use of saquinavir with indinavir, rifampin, rifabutin, phenytoin, carbamazepine, phenobarbital, efavirenz, dexamethasone, nevirapine, and other enzyme inducers. Avoid simvastatin or lovastatin because of rhabdomyolysis; can use pravastatin or fluvastatin	Hard-gel formulation (Invirase, 600 mg po tid within 2 hours of a high-fat meal to increase absorption); not recommended because of poor bioavailability (4%), even when taken with high-fat meal

Table 1. Continued

System, Problem, and Drug Regimen	Duration	Adverse Effects/Drug Interactions	Comments
GENERAL/SYSTEMIC			
Antiretroviral (Anti-HIV) (cont.)			
Ritonavir 400 - 600 mg po bid	Until efficacy wanes or toxicity occurs	See individual agents	Generally well tolerated
plus			Combination therapy provides higher saquinavir levels
Saquinavir soft-gel capsules 400 - 600 mg po bid. Lower dosages (400 mg of each) preferred			
Nelfinavir 750 mg po tid	Until efficacy wanes or toxicity occurs	See individual agents	Generally well tolerated
plus			Combination therapy provides higher saquinavir levels
Saquinavir 800 mg po tid			
Amprenavir (141W94, Agenerase) 1200 mg po bid with or without food	Until efficacy wanes or toxicity occurs	Nausea, diarrhea; hypertriglyceridemia, hypercholesterolemia, abnormal fat accumulation, hyperglycemia; headache; rash, Stevens-Johnson syndrome	Approval expected during 1999
		<i>Drug interactions</i> Metabolized by P-450 system; drug interactions (eg, rifampin, rifabutin) expected. Avoid simvastatin or lovastatin because of rhabdomyolysis; can use pravastatin or fluvastatin	
<i>Nonnucleoside reverse transcriptase inhibitors (NNRTIs)</i>			
Nevirapine (Viramune) 200 mg po qd for 14 days; if no rash develops, increase to 200 mg po bid. Once-daily dosing (400 mg po qd) under investigation.	Until efficacy wanes or toxicity occurs	Maculopapular rash, Stevens-Johnson syndrome; nausea, vomiting, diarrhea; fatigue, fever, headaches; aminotransferase elevations; rare hematologic toxicity	Discontinue drug at any time if rash is severe. Do not increase dosage if any rash is present during first 14-day lead-in period
		<i>Drug interactions</i> P-450 enzyme inducer; avoid concomitant use with saquinavir, rifampin, and rifabutin	Rash from one NNRTI does not predict rash from other NNRTIs
Delavirdine (Rescriptor) 400 mg po tid. Can dissolve in 3 oz water as slurry	Until efficacy wanes or toxicity occurs	Maculopapular rash; nausea; headache; aminotransferase elevations especially when taken with saquinavir; neutropenia when taken with nelfinavir	Delavirdine increases saquinavir and indinavir levels by 50%. Reduce indinavir dosage to 600 mg po q 8 h when used in combination with delavirdine. Separate didanosine or antacid administration from delavirdine administration by at least 1 hour
		<i>Drug interactions</i> P-450 enzyme inhibitor. Avoid concomitant use of astemizole, rifampin, rifabutin, phenytoin, carbamazepine, cisapride, alprazolam, midazolam, triazolam, ergot alkaloids. Ketoconazole, itraconazole, fluconazole, clarithromycin, and fluoxetine can increase delavirdine serum concentrations; dosage reduction might be necessary	Rash from one NNRTI does not predict rash from other NNRTIs
Efavirenz (DMP 266, Sustiva) 600 mg po qhs with or without food; 200 mg po tid if dizziness occurs	Until efficacy wanes or toxicity occurs	Dizziness, anxiety, inability to concentrate, lightheadedness, headache dysphoria, nightmares; nausea; rash (less than other NNRTIs). Avoid in pregnancy	Good central nervous system penetration; resistance might develop more slowly than other NNRTIs.
		<i>Drug interactions</i> Avoid use with saquinavir, astemizole, and cisapride. Increased indinavir dosage to 1 g po q 8 h might be required	Rash from one NNRTI does not predict rash from other NNRTIs

Continued

Table 1. Continued

System, Problem, and Drug Regimen	Duration	Adverse Effects/Drug Interactions	Comments
GENERAL/SYSTEMIC			
Antiretroviral (Anti-HIV) (cont.)			
<i>Other Agents</i>			
Hydroxyurea (Hydrea) 500 mg po bid	Until efficacy wanes or toxicity occurs	Bone marrow suppression including CD4+ count decline during hydroxyurea therapy	Can be used in combination with didanosine (and possibly other antiretroviral drugs) as salvage therapy. Long-term risks and benefits unknown
Adefovir dipivoxil (bis-POM PMEAs, Preveon) 120 mg po qd; lower dosage (60 mg po qd) might be effective		Nausea, vomiting, diarrhea, aminotransferase elevations; malaise; renal insufficiency, Fanconi-like syndrome with hypophosphatemia, glycosuria, renal acidosis, proteinuria, and elevated serum creatinine	Nucleotide analog. Role unclear at this time; might offer benefit in salvage therapy. Carnitine supplementation might be required. Approval expected during 1998. Available through expanded access at 1-800-445-3235
<i>Drug interactions</i> Avoid concurrent use with other nephrotoxins			
<i>Postexposure prophylaxis</i>			
Zidovudine 200 mg po tid plus lamivudine 150 mg po bid with or without nelfinavir 750 mg po tid or indinavir 800 mg po q 8 h	4 weeks	See above adverse effects and drug interactions. Zidovudine and lamivudine appear to be safe in pregnancy	Administration within 1 - 2 hours or as soon as possible after exposure. Can substitute other antiretroviral agents (eg, stavudine plus didanosine) when source patient has received extensive treatment with zidovudine or lamivudine. Add nelfinavir, indinavir, or other PI for high-risk exposures and when source patient suspected to have developed antiretroviral drug resistance. Can call 1-888-HIV-4911 for additional assistance
<i>Pregnancy</i>			
Zidovudine-containing antiretroviral regimen during pregnancy, plus intrapartum zidovudine 2 mg/kg IV for 1 hour, then 1 mg/kg/h until delivery	Until end of pregnancy	See above adverse effects and drug interactions Serious adverse effects on fetus not found in studies to date	Zidovudine therapy, combined with intrapartum intravenous zidovudine plus oral zidovudine 2 mg/kg qid to infants for first 6 weeks of life, decreases transmission to infants Unconfirmed reports of premature deliveries when PI therapy is used
Wasting Syndrome			
Megestrol (Megace) suspension (40 mg/mL) 800 mg po qd	Indefinitely	Nausea, vomiting; edema; adrenal suppression; depression. Progesterone side effects (hyperglycemia, decreased testosterone levels)	Megestrol can increase appetite and cause fat accumulation with weight gain. Uncertain whether this weight gain improves health. Available also as tablets, but large number of tablets required for administration and more expensive
Dronabinol (Tetrahydrocannabinol [THC], Marinol) 2.5 mg po bid 30 minutes to 1 hour before meals. Maximum 20 mg qd	Indefinitely	Restlessness, irritability, insomnia, dizziness, loss of coordination, psychotomimetic effects; fatigue; tachycardia	Increases appetite and can cause weight gain. Uncertain whether this weight gain improves health. Antinauseant. Not recommended for persons sensitive to marijuana effects
Human growth hormone (r-hGH, Serostim) 0.1 mg/kg/d SQ (average dosage 6 mg/d)	Unknown	Arthralgias, joint stiffness, carpal tunnel syndrome; hyperglycemia; hypertriglyceridemia	Can improve exercise endurance and increase weight, characterized by increased lean body mass and decreased fat
Anabolic steroids (eg, testosterone 200 mg IM every 2 weeks or 300 mg IM every 3 weeks, oxandrolone [Oxandrin] 2.5 mg po bid - tid or testosterone patches [Testoderm, Androderm])	Unknown	Edema; cholestatic jaundice, peliosis hepatis, aminotransferase elevations; increased libido, testicular atrophy, priapism; insomnia	Might improve well-being and increase lean body mass. Treatment should be accompanied by exercise

Table 1. Continued

System, Problem, and Drug Regimen	Duration	Adverse Effects/Drug Interactions	Comments
GENERAL/SYSTEMIC			
<i>Mycobacterium avium</i> complex (MAC)			
<i>Prophylaxis</i>			
Clarithromycin (Biaxin) 500 mg po bid	Indefinitely	Clarithromycin and azithromycin side effects include nausea, vomiting, dyspepsia, diarrhea, hearing loss, aminotransferase elevations	Survival benefits shown for clarithromycin. Prophylaxis can be offered to patients with CD4+ cell counts < 50/ μ L
OR			
Azithromycin (Zithromax) 1200 mg po once weekly or 500 mg po qd	Indefinitely	<i>Drug interactions</i> Clarithromycin increases serum levels of rifabutin and can lead to rifabutin toxicity, including severe anterior uveitis. Clarithromycin and azithromycin increase levels of carbamazepine and theophylline. Avoid astemizole in combination with azole antibiotics because of increased risk of torsades de pointes and ventricular tachyarrhythmias	Clarithromycin and rifabutin might provide prophylaxis against <i>Cryptosporidium</i>
OR			
Rifabutin (Mycobutin) 300 mg po qd	Indefinitely	Nausea (can be reduced by administering 150 mg po bid). Rash. Uveitis with dosages > 300 mg po qd and in patients receiving concomitant clarithromycin, fluconazole, or indinavir therapy. Red-orange discoloration of body fluids. Rare neutropenia, thrombocytopenia, anemia; flu-like syndrome; elevated bilirubin and alkaline phosphatase levels, hepatitis <i>Drug interactions</i> Multiple interactions with protease inhibitors (see Antiretroviral drugs, above). Rifabutin increases metabolism of methadone, zidovudine, and clarithromycin; higher dosage of these drugs might be required. Clarithromycin increases rifabutin blood levels and can lead to rifabutin toxicity	Exclude <i>Mycobacterium tuberculosis</i> infection before initiating rifabutin therapy
<i>Acute MAC disease</i> Ethambutol (Myambutol) 15 mg/kg po qd (1 g po qd maximum); dosage reduction in renal failure	Indefinitely, if tolerated (minimum of 12 weeks)	Optic neuritis (if > 25 mg/kg/d); hyperuricemia; nausea, vomiting	Treatment indicated for documented MAC disease and patients with progressive signs, symptoms, and laboratory abnormalities consistent with MAC disease. Clinical improvement might take 2 - 4 weeks. Isolation of MAC in stool or sputum might not indicate systemic disease but is usually treated with ethambutol plus a macrolide antibiotic
plus either			
Clarithromycin 500 mg po bid. Higher dosages associated with higher mortality			
or			
Azithromycin 500 mg po qd			When both <i>M tuberculosis</i> and MAC infections are suspected, add isoniazid, rifampin, and pyrazinamide to ethambutol and clarithromycin pending culture results. See <i>M tuberculosis</i>
<i>For serious illness or failure to respond within 1 month, can add one or two of the following:</i>			
Rifabutin 300 mg po qd	Indefinitely		Rifampin (Rimactane, Rifadin) 450 - 600 mg po qd can be substituted for rifabutin if concerned about <i>M tuberculosis</i>

Continued

Table 1. Continued

System, Problem, and Drug Regimen	Duration	Adverse Effects/Drug Interactions	Comments
GENERAL/SYSTEMIC			
<i>Mycobacterium avium</i> complex (MAC) (cont.)			
Ciprofloxacin (Cipro) 500 - 750 mg po qd - bid	Indefinitely	Nausea, vomiting, diarrhea. Reversible pink to brown-black discoloration of skin, eyes, body secretions; rash. Hyperglycemia. Retinal degeneration <i>Drug interactions</i> Binds to cations, resulting in decreased ciprofloxacin absorption. Administer 2 - 4 hours after antacids, sucralfate, dairy products, and didanosine	
Amikacin (Amikin) 7.5 - 10.0 mg/kg IM/IV qd	2 - 8 weeks	Nephrotoxicity, ototoxicity	Monitor drug levels in patients with renal failure
<i>Mycobacterium tuberculosis</i>			
<i>Prophylaxis</i>			
Isoniazid (INH) 300 mg po qd	6 - 12 months	Aminotransferase elevations and hepatitis; administer with pyridoxine to prevent peripheral neuropathy <i>Drug interactions</i> Increases metabolism of ketoconazole; larger dosages of ketoconazole might be required. Increased phenytoin and carbamazepine toxicity; monitor levels	Prophylaxis for all HIV-infected persons with ≥ 5 -mm intermediate-strength tuberculin skin test induration and those with strong history of tuberculosis exposure regardless of skin test reactivity Isoniazid can be administered concurrently with NRTIs, PIs, NNRTIs
OR			
Rifampin or rifabutin plus Pyrazinamide or isoniazid	2 - 3 months	See individual drug toxicities <i>Drug interactions</i> PIs should not be administered concurrently with rifampin. Rifabutin contraindicated with ritonavir, soft-gel saquinavir, delavirdine, and possibly other antiretroviral drugs	When short-course prophylaxis is administered with or without directly observed therapy (DOT), consultation with tuberculosis experts is recommended. Effective antiretroviral therapy should not be discontinued to permit use of specific antituberculosis drugs
<i>Active tuberculosis</i>			
Combinations of isoniazid, rifampin or rifabutin, pyrazinamide, ethambutol, and streptomycin	Begin with 4 drugs. After 2 months can usually continue 2-drug therapy, depending upon susceptibility testing results. Total treatment: at least 6 months, and 6 months beyond culture conversion	See individual drug adverse effects and drug interactions <i>Drug interactions</i> PIs should not be administered concurrently with rifampin. Rifabutin contraindicated with ritonavir, soft-gel saquinavir, delavirdine, and possibly other antiretroviral drugs	Consultation with tuberculosis experts required. Treatment guidelines available through Centers for Disease Control and Prevention at http://www.cdc.gov
Histoplasmosis			
<i>Acute</i>			
Amphotericin B (Fungizone) 1.0 mg/kg IV qd until 15 mg/kg total dosage has been administered. Decrease to 0.7 - 0.8 mg/kg qd if not tolerated followed by	6 - 8 weeks total acute therapy (amphotericin plus itraconazole)	See CENTRAL NERVOUS SYSTEM, <i>Cryptococcus neoformans</i>	Amphotericin B recommended initially; oral therapy does not appear as effective. Itraconazole 200 mg po bid might be effective
Itraconazole (Sporanox) 200 mg po bid	See above	Nausea, vomiting; hypokalemia; hypertension; aminotransferase elevations; adrenal insufficiency; rhabdomyolysis.	Teratogenic

Table 1. Continued

System, Problem, and Drug Regimen	Duration	Adverse Effects/Drug Interactions	Comments
GENERAL/SYSTEMIC			
Histoplasmosis (cont.)			
Itraconazole (cont.)			
		<i>Drug interactions</i> Potent hepatic enzyme inducers, such as rifampin and phenytoin, increase metabolism of itraconazole; higher itraconazole dosages might be required. Avoid concurrent use with triazolam, alprazolam (Xanax), antacids, H ₂ blockers, and omeprazole	
<i>Maintenance</i> Itraconazole 200 mg po qd	Indefinitely		Fluconazole 400 mg po qd less effective
OR			
Amphotericin B 50 mg IV each week, 2 times a week, or every other week	Indefinitely		Optimum frequency of administration not determined
Coccidioidomycosis			
<i>Acute</i>			
Amphotericin B (as above)	6 - 8 weeks	See CENTRAL NERVOUS SYSTEM <i>Cryptococcus neoformans</i>	Fluconazole penetrates CNS and is preferred initial therapy for CNS coccidioidomycosis
or			
Fluconazole	Indefinitely		
<i>Maintenance</i> Fluconazole	Indefinitely		
Cryptococcosis			
		See CENTRAL NERVOUS SYSTEM, <i>Cryptococcus neoformans</i>	
SKIN/MUCOCUTANEOUS			
Kaposi sarcoma			
Observation	Indefinitely		Treatment not required unless lesions are symptomatic or cosmetically bothersome
OR			
Local treatment (radiation therapy, cryotherapy, excision, or intralesional vinblastine)	Until lesions and symptoms are resolved or controlled	Mucositis in head and neck regions from radiation therapy	Treatment effective for cosmetic purposes, for relief of symptoms, and to help reduce edema caused by lymphatic obstruction
OR			
Systemic chemotherapy with vinblastine and vincristine, vincristine alone, or combination of doxorubicin, bleomycin, and vincristine	Same	Usual chemotherapeutic agent side effects. Liposomal preparations might have some advantages in specific cases	Therapy can help control disease but does not alter prognosis. Consultation by oncologist or AIDS specialist usually required
OR			
Interferon-alpha 3 mU SQ 3 times weekly; increase by 3 mU/d every 2 weeks as tolerated (maximum 27 mU/d)	Indefinitely	Fatigue, myalgia, asthenia; neutropenia, thrombocytopenia; aminotransferase elevations	Hematologic toxicities increased in patients taking zidovudine. Dosages greater than 10 mU/d necessary for efficacy
Seborrheic dermatitis			
Hydrocortisone (HC) cream 2.5% plus ketoconazole cream 2% bid; severe cases can require ketoconazole 200 - 400 mg po qd for 3 - 4 weeks	Until resolved	See ORAL CAVITY, <i>Candida albicans</i> , ketoconazole	Commonly involves face, eyebrows, retroauricular areas, nasolabial folds, and scalp. Addition of antifungal cream enhances therapeutic response and reduces the frequency of steroid application

Continued

Table 1. Continued

System, Problem, and Drug Regimen	Duration	Adverse Effects/Drug Interactions	Comments
SKIN/MUCOCUTANEOUS (cont.)			
Mucocutaneous herpes simplex			
<i>Acute</i>			
Acyclovir (Zovirax) 200 mg po 5 times a day or 400 mg po tid	7 - 10 days	Oral: nausea, vomiting, diarrhea, dizziness	Topical acyclovir ineffective for most episodes
OR			
Valacyclovir (Valtrex) 500 mg - 1 g po bid	10 days	Nausea, vomiting, diarrhea; headache, dizziness, fatigue, insomnia. Hemolytic uremic syndrome (if > 3 g/d)	
OR			
Famciclovir (Famvir) 250 mg po tid	10 days	Nausea, vomiting, diarrhea; headache, dizziness, fatigue, insomnia	
<i>Maintenance</i>			
Acyclovir 200 - 400 mg po 2 - 3 times a day or valacyclovir 500 mg po bid or 1 g po qd or famciclovir 250 mg po bid	Indefinitely		Chronic maintenance therapy might be necessary for repeated episodes
Disseminated, extensive, or persistent herpes simplex			
Acyclovir 5 mg/kg/dose IV q 8 h; dosage reduction in renal failure; maintenance as above	7 - 14 days or until lesions resolve	Intravenous: lethargy, tremors, confusion, hallucinations; phlebitis; increased serum creatinine, reversible crystalline nephropathy	Severe herpes infections (eg, esophagitis, colitis, encephalitis) require intravenous acyclovir. Maintain good urine output and hydration to prevent acyclovir crystallization
OR			
Valacyclovir 1 g po tid	7 - 14 days or until lesions resolve	See above	
Herpes zoster (shingles, disseminated, or persistent zoster)			
Acyclovir 10 mg/kg/dose IV q 8 h; or acyclovir 800 mg po 5 times a day; re- duce dosage of intravenous acyclovir in renal failure	7 - 10 days or until lesions resolve		Alternate drugs are foscarnet, vidarabine, cidofovir, and trifluridine (Viroptic) applied to skin covered with polymyxin B-bacitracin (Polysporin) ointment q 8 h. Keratoconjunctivitis requires more frequent (q 2 h) trifluridine application
OR			
Valacyclovir 1 g po tid	7 - 10 days		
Acyclovir-resistant herpes infections			
Foscarnet 40 mg/kg/dose IV q 8 h; dosage reduction in renal failure	10 - 14 days or until lesions clear	See OPHTHALMOLOGIC, CMV, below	See OPHTHALMOLOGIC, CMV, below
OR			
Trifluridine (Viroptic) 1% solution q 8 h	Same	Rare hypersensitivity reactions	Apply to affected areas and cover with antibiotic ointment such as bacitracin or polymyxin B Keratoconjunctivitis requires more frequent (as often as 2 hours, maximum 9 drops a day) trifluridine application
Cidofovir	Same	See OPHTHALMOLOGIC, CMV, below	Cidofovir might be effective

Table 1. Continued

System, Problem, and Drug Regimen	Duration	Adverse Effects/Drug Interactions	Comments
SKIN/MUCOCUTANEOUS (cont.)			
Bacillary angiomatosis			
Erythromycin 500 mg po qid	2 months	See GENERAL/SYSTEMIC, MAC, clarithromycin, azithromycin. Jarisch-Herxheimer reaction with systemic disease	Skin lesions can resolve in 1 - 3 weeks, but 2 months' treatment needed. Systemic disease (eg, hepatic, splenic, central nervous system, bone) or cutaneous recurrences require treatment for 4 months or indefinitely. Azithromycin 1 g po qd and clarithromycin 500 - 1000 mg po qd can be used as alternatives
OR			
Doxycycline 100 mg po bid	2 months		
Eosinophilic folliculitis			
High-potency fluorinated corticosteroid cream bid	Indefinitely		Itraconazole 200 mg po once daily with food might be effective. If no response in 2 weeks, increase dosage to 200 mg po bid for 2 additional weeks. If no response after 4 weeks, discontinue. Topical metronidazole might be helpful
plus			
Antihistamine (eg, diphenhydramine [Benadryl], hydroxyzine [Atarax, Vistaril], doxepin [Sinequan])	Indefinitely		Avoid astemizole in combination with azole antibiotics because of increased risk of torsades de pointes and ventricular tachyarrhythmias
HEMATOLOGIC			
Thrombocytopenia			
Observation		Discontinue drugs that can cause thrombocytopenia	Treatment not required in absence of bleeding. Consider platelet transfusions prior to invasive procedures. High-dosage zidovudine, corticosteroids (eg, prednisone 60 mg po qd), splenectomy, intravenous gamma globulin, and interferon-alpha can raise platelet count
		Corticosteroids can increase immunodeficiency	
OPHTHALMOLOGIC			
Cytomegalovirus (CMV)			
<i>Prophylaxis</i>			
Ganciclovir (Cytovene) 1 g po tid	Indefinitely	See below	Oral ganciclovir primary prophylaxis is not currently recommended. Efficacy not established
<i>Acute retinitis</i>			
<i>Induction</i>			
Ganciclovir 5 mg/kg/dose IV q 12 h; dosage reduction in renal failure	14 days for acute retinal infection; 14 - 21 days usually required for extraocular infection	Neutropenia, leukopenia, anemia, thrombocytopenia (avoid if platelet count < 20,000/ μ L); aminotransferase elevations; renal failure; phlebitis, rash; nausea. Discontinue zidovudine during induction to minimize additive hematologic toxicity (neutropenia). To avoid hematologic toxicity, substitute didanosine, zalcitabine, or stavudine for zidovudine, or change to foscarnet	Intravitreal ganciclovir by injection or implant appears effective if IV causes unacceptable toxicity. Does not provide systemic therapeutic effect or protection of contralateral eye
OR			Start G-CSF (Filgrastim, Neupogen) 300 μ g SQ qd to 3 times a week for ganciclovir-induced neutropenia (ANC < 500/ μ L) on two consecutive measurements
Foscarnet (Foscavir) 90 mg/kg/dose IV q 12 h as 2-hour infusion, dosage discontinuation or reduction in renal failure	14-day induction	Nephrotoxicity common; tremors, headaches, occasional seizures, muscle spasms; hypocalcemia, hypercalcemia, hypokalemia, hypophosphatemia, hyperphosphatemia; anemia, granulocytopenia; aminotransferase elevations; phlebitis, penile ulcerations	Administered by infusion pump via central line. Infusion of 500 - 1000 mL normal saline before each foscarnet administration can minimize nephrotoxicity. Creatinine clearance should be measured in cachectic patients and in patients with renal insufficiency to ensure proper use of administration nomogram
OR		<i>Drug interactions</i> Avoid concurrent use of nephrotoxic agents when possible	
Ganciclovir plus foscarnet		See individual agents above	Continue maintenance drug, induce with the alternative drug, then continue maintenance therapy with both drugs

Continued

Table 1. Continued

System, Problem, and Drug Regimen	Duration	Adverse Effects/Drug Interactions	Comments
OPHTHALMOLOGIC			
Cytomegalovirus (CMV) (cont.)			
<i>Alternatives to ganciclovir or foscarnet</i>			
Cidofovir (Vistide) 5 mg/kg IV with probenecid (2 g po 3 hours before and 1 g po 2 and 8 hours after infusion) each week for 2 weeks, then every 2 weeks thereafter; contraindicated in renal insufficiency (serum creatinine ≥ 1.5 mg/dL, CrCl ≤ 55 mL/min, 2+ proteinuria)	14-day induction period	Life-threatening nephrotoxicity; fever; nausea, diarrhea; rash; uveitis, iritis, ocular hypotonia; proteinuria, metabolic acidosis; neutropenia. Persons allergic to sulfa compounds can be allergic to probenecid <i>Drug interactions</i> Avoid concomitant administration with any potentially nephrotoxic agent, including nonsteroidal anti-inflammatory drugs	Not known whether cidofovir is as effective as ganciclovir or foscarnet. Indwelling catheter not required Prehydrate with 1 L normal saline. Do not administer within 7 days of other potentially nephrotoxic agents. Patients previously treated with foscarnet are at increased risk for renal failure. Administer G-CSF if ANC consistently $< 500/\mu\text{L}$
OR			
Ganciclovir implant (Vitaset)	Indefinitely	Implant can cause transient visual deterioration, retinal detachment, vitreal hemorrhage, cataracts, and endophthalmitis	Implant ideal for patients unable to take daily intravenous therapy. Combine with oral ganciclovir to provide protection against systemic disease and contralateral eye involvement
plus			
Ganciclovir 1g po tid		Oral ganciclovir: Anemia, neutropenia; nephrotoxicity; neuropathy <i>Drug interactions</i> Oral ganciclovir therapy causes 50% increase in didanosine blood levels; reduce didanosine dosage by 50%	Oral ganciclovir absorption is erratic when diarrhea is present. Administer on empty stomach to improve absorption
<i>Maintenance</i>			
Ganciclovir 5 mg/kg IV qd as 1-hour infusion; dosage reduction in renal failure	Indefinitely		Lifelong suppressive therapy required to prevent recurrence of retinitis. Less frequent (5 - 6 days per week) administration might be acceptable. Administer G-CSF or change to foscarnet if ANC consistently $< 500/\mu\text{L}$
OR			
Foscarnet 90 mg/kg IV qd as 2-hour infusion; discontinuation or dosage reduction in renal failure	Indefinitely		Maintenance with 120 mg/kg/d might be more effective but also more toxic
OR			
Ganciclovir plus foscarnet	Indefinitely	See individual agents above	Combination therapy not routinely recommended as initial therapy. Continue maintenance dosage of current drug; induce alternate drug, followed by maintenance with both drugs. Reinduction with ganciclovir or foscarnet might be helpful for recurrences when alternative drug cannot be administered
OR			
Ganciclovir 1 g po tid	Indefinitely	See above	Oral ganciclovir is not as effective for maintenance therapy as other regimens
OR			
Cidofovir 5 mg/kg as 1-hour infusion every 2 weeks at infusion center	Indefinitely	Life-threatening nephrotoxicity; cannot be given with potentially nephrotoxic drugs	Does not require indwelling catheter; quality of life might be improved
ORAL CAVITY			
<i>Candida albicans</i>			
Clotrimazole (Mycelex) troches 10 mg 5 times a day or vaginal suppositories 100 mg qd - bid. Dissolve troche slowly in mouth	1 - 2 weeks or until resolved; maintenance (with lowest effective dosage) might be required for severe or frequent recurrences	Minimal toxicity. Unpleasant taste, nausea, vomiting; aminotransferase elevations	Troches have high sugar content and often require frequent administration. Suppositories can be more convenient

Table 1. Continued

System, Problem, and Drug Regimen	Duration	Adverse Effects/Drug Interactions	Comments
ORAL CAVITY			
<i>Candida albicans</i> (cont.)			
OR			
Nystatin (Mycostatin) 100,000 U/mL, swish and swallow 5 mL po q 6 h or one 500,000-U tablet dissolved slowly in mouth q 6 h	Same	Large oral doses can produce diarrhea, nausea, vomiting	Generally less effective than ketoconazole, fluconazole, and clotrimazole. Can be effective in fluconazole-resistant candidal infection
OR			
Fluconazole (Diflucan) 100 - 200 mg po qd followed by maintenance therapy 50 - 100 mg po qd; 100 - 200 mg po once weekly is less effective	Same	See CENTRAL NERVOUS SYSTEM, <i>Cryptococcus neoformans</i>	Effective in oral candidiasis unresponsive to above oral agents. Higher dosages might be required. Fluconazole solution or itraconazole 200 mg po qd (or itraconazole solution) might be effective against fluconazole-resistant <i>Candida albicans</i>
OR			
Ketoconazole (Nizoral) 400 mg po qd followed by maintenance therapy 200 mg po qd - bid for 7 consecutive days per month	Same	Nausea, vomiting; aminotransferase elevations; anaphylaxis, urticaria. Higher dosages can suppress testosterone levels; gynecomastia; adrenal suppression <i>Drug interactions</i> Need gastric acidity to be effective; avoid antacids, H ₂ antagonists; administer 2 hours apart from didanosine. Higher dosages might be necessary if taking rifampin. Avoid concurrent use with triazolam or alprazolam	
OR			
Amphotericin B oral suspension 100 mg/mL, swish and swallow 1 - 5 mL qid	Same	Unpalatable; nausea, vomiting, diarrhea; rare urticaria	Not absorbed. No systemic effects. Intravenous amphotericin B might be necessary for severe disease
Periodontal disease			
Hydrogen peroxide gargles for 30 sec bid	Indefinitely		Oral hygiene measures with manual removal of plaque are essential. Severe periodontal disease can require antibiotic therapy with metronidazole 250 mg po tid or 7 - 10 days (alternatives: clindamycin or amoxicillin-clavulanate [Augmentin]). Antiseptic mouthwash (Listerine) gargles can be effective
OR			
Chlorhexidine gluconate (Peridex) oral rinse 15 mL swished in mouth for 30 sec bid	Indefinitely	Staining of teeth	
ESOPHAGEAL			
<i>Candida albicans</i>			
Fluconazole 200 - 400 mg po qd; higher dosages might be required	14 - 21 days; maintenance with lowest effective dosage	See CENTRAL NERVOUS SYSTEM, <i>Cryptococcus neoformans</i>	Empiric treatment for patients with dysphagia or odynophagia who have oral thrush. Endoscopy with biopsy and cultures appropriate for patients who fail to respond within 1 week. Ketoconazole less expensive than fluconazole and effective in most patients
OR			
Ketoconazole 200 mg po bid; see ORAL CAVITY, <i>Candida albicans</i>	Same as above	See ORAL CAVITY, <i>Candida albicans</i>	

Continued

Table 1. Continued

System, Problem, and Drug Regimen	Duration	Adverse Effects/Drug Interactions	Comments
ESOPHAGEAL			
<i>Candida albicans</i> (cont.)			
OR			
Amphotericin B 0.3 - 0.4 mg/kg IV qd	10 days or until resolution		Candidal esophagitis unresponsive to oral agents requires low-dose IV amphotericin B
Cytomegalovirus			
Ganciclovir; foscarnet; see OPHTHALMOLOGIC, CMV	14 - 21 days	See OPHTHALMOLOGIC, CMV	Diagnose by endoscopic appearance plus biopsy showing CMV inclusion bodies and positive culture. Long-term suppressive therapy indicated only after multiple recurrences. Beware of drug resistance
Herpes simplex			
Acyclovir IV or valacyclovir po; see SKIN/MUCOCUTANEOUS, disseminated, extensive, or persistent herpes simplex	10 - 14 days; maintenance required	See SKIN/MUCOCUTANEOUS, disseminated, extensive, or persistent herpes simplex	Diagnose by endoscopic appearance plus positive culture
GASTROINTESTINAL			
Nausea and vomiting			
Prochlorperazine (Compazine) 2.5 - 10.0 mg IV or 5 - 10 mg po, or IM q 6 h, or 25 mg po q 12 h	As needed	Fatigue, drowsiness, dizziness, depression; extrapyramidal reactions; dystonic reactions; aminotransferase elevations; constipation	Combinations of these agents often necessary Haloperidol (Haldol) can also be effective
Metoclopramide (Reglan) 10 mg po qid, or 1 mg/kg IV q 3 h, or 10 mg IM q 4 - 6 h. Dosage reduction in renal failure	As needed	Same as above	Same as above
Lorazepam (Ativan) 0.5 - 2.0 mg po or SL tid - qid	As needed	Similar to benzodiazepines; antegrade amnesia	Effective for anticipatory nausea
Granisetron (Kytril) 1 mg po q 12 h, or 10 µg/kg bid IV, or ondansetron (Zofran) 0.15 mg/kg IV infusion for 15 min q 6 h or 4 - 10 mg po q 6 h	As needed	Constipation, diarrhea, abdominal pain; fever, chills; headache; sedation	Reserved for intractable nausea and vomiting unresponsive to other agents. Ondansetron or granisetron in combination with droperidol helpful for intractable nausea and vomiting
Dronabinol (Marinol) 2.5 - 10.0 mg po q 8 - 12 h	As needed	See GENERAL/SYSTEMIC, wasting syndrome	Effective in drug-induced nausea. Marijuana can be helpful
Droperidol (Inapsine) 2.5 mg IM/IV q 4 - 6 h	As needed	Similar to prochlorperazine	
Diarrhea			
<i>Symptomatic treatment</i>			
Loperamide (Imodium) 4 mg po initially then 2 mg q 6 h around the clock and prn (maximum 16 mg qd)	As needed	Abdominal cramps, nausea, abdominal distention, vomiting; dizziness, drowsiness	Around-the-clock regimen more effective than prn. Treat to 2 - 3 bowel movements per day
Diphenoxylate-atropine (Lomotil) 2.5 - 5.0 mg po 3 - 6 times daily for 24 - 48 hours; then 2.5 - 5.0 mg tid and prn to control diarrhea (maximum 20 mg qd)	As needed	Ileus; nausea, vomiting, abdominal discomfort; anticholinergic side effects secondary to atropine	Same as above. 2.5 mg diphenoxylate-atropine is equivalent to 2 mg morphine sulfate
Paregoric 0.4 mg morphine/mL, 5 - 10 mL qd - qid, or tincture of opium 10 mg morphine/mL, 0.3 - 1.0 mL po qid and prn (maximum 1 mL/dose or 6 mL/d), or equivalent	As needed	Ileus. Altered mental status, hallucinations. Adverse effects common to narcotic analgesics	Same as above. 5 mL paregoric and 0.2 mL tincture of opium are equivalent to 2 mg morphine sulfate

Continued

Table 1. Continued

System, Problem, and Drug Regimen	Duration	Adverse Effects/Drug Interactions	Comments
GASTROINTESTINAL			
Diarrhea (cont.)			
Octreotide (Sandostatin) 100 µg SQ tid, increase by 100 - 200 µg q 1 - 2 wk until maximum of 500 µg SQ tid	Indefinitely	Nausea, steatorrhea; hyperglycemia; pain at injection site	Not approved by FDA. Efficacy not demonstrated. Long-term safety unknown. Octreotide does not improve malabsorption
Cryptosporidium			
Paromomycin (Humatin) 750 mg po tid	10 - 14 days or indefinitely	Nausea, vomiting, diarrhea; rare ototoxicity and nephrotoxicity (similar to other aminoglycosides) only if absorbed through ulcerative bowel lesions	No evidence of efficacy. Addition of azithromycin 600 mg po qod might increase effectiveness
Isospora belli			
Trimethoprim-sulfamethoxazole (TMP-SMX, Septra, Bactrim) 1 DS (double-strength) tablet po qid	21 days	See PULMONARY, PCP	Usually effective
Cytomegalovirus			
Ganciclovir; foscarnet; see OPHTHALMOLOGIC, CMV	14 - 21 days	See OPHTHALMOLOGIC, CMV	Long-term suppressive therapy indicated only after multiple recurrences. Beware of drug resistance
PULMONARY			
Pneumocystis carinii pneumonia (PCP)			
<i>Prophylaxis for patients with AIDS (including CD4+ cell count < 200/µL), unexplained fever, or oral candidiasis</i>			
Trimethoprim-sulfamethoxazole (TMP-SMX) 1 DS tablet po qd or qod or 3 times a week (eg, M-W-F)	Indefinitely	See TMP-SMX below	TMP-SMX considered most effective for prophylaxis. TMP-SMX provides additional prophylaxis against toxoplasmosis and common bacterial infections
<i>Alternatives to TMP-SMX for prophylaxis</i>			
Dapsone 50 mg po bid or 100 mg po qd with or without TMP (Trimprax) 15 mg/kg/d or pyrimethamine (Daraprim) 25 - 75 mg po q wk	Indefinitely	See dapsone plus TMP. Patients allergic to sulfa might tolerate dapsone; some cross-sensitivity	Probably less effective than TMP-SMX; might be less toxic. Check glucose-6 phosphate dehydrogenase (G6PD) before starting dapsone. Lower dosages (eg, 100 mg po 2 times a week) might be effective
OR			
Atovaquone (Mepron) suspension (750 mg/5 mL) 1500 mg po q d or 750 mg po bid, with or without pyrimethamine 25 - 75 mg po q wk	Indefinitely	Headaches; nausea, diarrhea, aminotransferase elevations; rash, drug fever; neutropenia, anemia; transient conjunctivitis; erythema multiforme	Take with food to increase drug absorption. Patients with enteropathy might not absorb a sufficient amount of atovaquone for adequate treatment. Better tolerated than dapsone; efficacy similar
OR			
Inhaled pentamidine (Aeropent) 300 mg q 4 wk using Respigard II nebulizer	Indefinitely	Bronchospasm and coughing are common; pretreatment with inhaled bronchodilator (eg, albuterol) can help. Increased risk of spontaneous pneumothorax. Minimal systemic effects. Rare pancreatitis, hypoglycemia; rare nephrotoxicity	Effective for prophylaxis against primary PCP when CD4+ cell count > 150/µL. Does not prevent extrapulmonary disease. Upper lobe recurrences from poor drug distribution when inhaled in upright position. Do not use in patients with possible <i>M tuberculosis</i> infection because of risk of <i>M tuberculosis</i> spread by aerosolization
OR			
Clindamycin (Cleocin) 450 - 600 mg po bid - tid plus primaquine 15 mg po qd	Indefinitely	See Acute PCP below	Efficacy and proper dosages for PCP prophylaxis unknown

Continued

Table 1. Continued

System, Problem, and Drug Regimen	Duration	Adverse Effects/Drug Interactions	Comments
PULMONARY <i>Pneumocystis carinii</i> pneumonia (PCP) (cont.)			
OR			
Pyrimethamine 25 mg - sulfadoxine 500 mg (Fansidar) 1 po q 2 wk	Indefinitely	Stevens-Johnson syndrome, toxic epidermal necrolysis; bone marrow suppression; gastrointestinal, central nervous system toxicity	No studies clearly show efficacy
<i>Acute PCP</i> TMP-SMX; TMP 15/kg/d given in 3 divided doses either po or as 1- to 2-hour IV infusions; lower dosages (TMP 12 mg/kg/d) can be effective and less toxic	21 days	Adverse effects commonly appear between 7 and 14 days in more than 50% of patients	TMP-SMX is the drug of choice and should be used unless severe reactions (eg, anaphylaxis, Stevens-Johnson syndrome) are of concern. Oral and intravenous routes equally effective
Note: Patients with substantial hypoxemia require concomitant corticosteroids (see below)		Rashes: maculopapular, exfoliative, Stevens-Johnson syndrome	Mild rash does not necessitate stopping or changing treatment: institute antihistamine or rechallenge with lower dosage of TMP-SMX. Desensitization more successful than rechallenge.
		Hematologic: neutropenia, leukopenia, thrombocytopenia, anemia	If ANC < 500/ μ L or if platelet count < 30×10^9 /L and bleeding occurs, consider alternative treatment
		<i>Drug interactions</i> Concurrent leucovorin calcium therapy associated with increased rate of therapeutic failure	
		Gastrointestinal: nausea, vomiting, aminotransferase elevations	Pretreatment with lorazepam, prochlorperazine, metoclopramide, or dronabinol to reduce nausea. Nausea can be less with oral TMP-SMX. Aminotransferase elevations 4 - 5 times normal require treatment change
		Renal: increased blood urea nitrogen (BUN) and creatinine; hyperkalemia secondary to effects of TMP	TMP decreases creatinine tubular secretion and can elevate serum creatinine levels. Discontinue TMP-SMX if serum creatinine > 3.0 mg/dL
		Hyponatremia	Can be caused by large volume of 5% dextrose in water (D5W) needed for IV administration; dilute each 80 mg TMP in 75 mL D5W or change to oral TMP-SMX. For severe hyponatremia ($\text{Na}^+ < 115$ mEq/dL), can dilute in normal saline; administer within 1 hour of preparation to avoid TMP-SMX precipitation
<i>Alternatives to TMP-SMX for acute PCP</i> Pentamidine isethionate (Pentam) 4 mg/kg/d as 1- to 2-hour IV infusion once a day; 3 mg/kg/d might also be effective		Neurologic: tremor, psychosis, aseptic meningitis	Tremors can be confused with seizures
		Drug fever. Sepsis-like syndrome, especially upon rechallenge	Drug fever can herald onset of neutropenia, rash, hepatitis, and bone marrow toxicity
	21 days	Adverse effects commonly appear between 7 and 14 days	
		Orthostatic hypotension can be severe and occur with initial infusion	Slow IV infusion for 2 hours can prevent hypotension. Check blood pressure at end of infusion
		Pancreatitis; early or delayed hypoglycemia (can occur after discontinuation of therapy); hyperglycemia	Hypoglycemia can be profound and prolonged, requiring immediate IV D50W followed by D10W glucose infusions. Permanent diabetes can occur.
		<i>Drug interactions</i> Avoid concomitant pancreatic toxins such as didanosine, zalcitabine, and alcohol	

Continued

Table 1. Continued

System, Problem, and Drug Regimen	Duration	Adverse Effects/Drug Interactions	Comments
PULMONARY <i>Pneumocystis carinii</i> pneumonia (PCP) (cont.)			
Pentamidine isethionate (cont.)		Renal failure; hyperkalemia. Concomitant nephrotoxic agents (eg, nonsteroidal anti-inflammatory agents) and dehydration increase risk of nephrotoxicity	Obtain accurate patient weight every 2 - 3 days to adjust pentamidine dosage. Discontinue pentamidine if creatinine > 3.0 mg/dL. Can resume administration if creatinine < 2 mg/dL
		Rare: neutropenia, thrombocytopenia; hypocalcemia, hypomagnesemia; aminotransferase elevations; cardiac arrhythmias (rare) with prolongation of QT interval and T-wave flattening	
OR			
Clindamycin 600 mg IV or po tid	21 days	Maculopapular rash (day 10 - 12 most common, usually self-limiting), fever; diarrhea, nausea, vomiting, abdominal cramps, <i>Clostridium difficile</i> colitis, aminotransferase elevations	Consider in patients with mild-to-moderate PCP, intolerant of or unresponsive to TMP-SMX
plus			
Primaquine 30-mg base po qd		Methemoglobinemia from primaquine, hemolysis in G6PD-deficient patients; leukopenia	Check G6PD before initiating primaquine therapy. Check methemoglobin levels when clinically indicated (see dapsone). Vitamin C 1 g po tid might prevent methemoglobinemia. Lower dosage of primaquine (15 mg po qd) can be effective
OR			
Dapsone 50 mg po bid plus either TMP 15 mg/kg/d po in 3 - 4 divided doses or pyrimethamine 50 - 75 mg po qd	21 days	See toxicities for TMP-SMX. Patients allergic to sulfa often tolerate dapsone. Methemoglobinemia, dose-related hemolysis, bone marrow suppression; rash; fever; nausea, abdominal pain; hyperkalemia; proteinuria, papillary necrosis	Effective in mild-to-moderate PCP only. Check G6PD before starting dapsone. Check methemoglobin levels if symptomatic or discrepancy between oxygen saturation and simultaneous arterial PaO ₂ . Treat methemoglobinemia > 20% (13% - 20% if anemic or respiratory compromise) with methylene blue 1% solution 2 mg/kg IV once; methylene blue contraindicated in G6PD deficiency. Vitamin C 1 g po tid might prevent methemoglobinemia
		<i>Drug interactions</i> Drug interactions with rifampin and rifabutin can render dapsone ineffective	
OR			
Trimetrexate (Neutrexin) 45 mg/m ² IV qd	21 days	Granulocytopenia, fever, rash; aminotransferase elevations	Can be effective in some patients as salvage therapy
plus			
Dapsone 50 mg po bid	21 days	See above	
plus			
Leucovorin calcium (folinic acid) 20 mg/m ² IV or po q 6 h	24 days		Must be administered for 72 hours after the last dose of trimetrexate. Large IV fluid load with leucovorin administration can result in volume overload
OR			
Atovaquone suspension (750 mg/5 mL) 750 mg po bid with food	21 days	Rash, drug fever; headaches; nausea, diarrhea, aminotransferase elevations; neutropenia, anemia; transient conjunctivitis; erythema multiforme	Higher therapeutic failure rate than TMP-SMX. For patients who fail or are intolerant of TMP-SMX, pentamidine, dapsone-TMP, or clindamycin-primaquine. Take with food to increase drug absorption. Patients with enteropathy might not absorb a sufficient amount of atovaquone for adequate treatment
plus			
Pyrimethamine 50 - 75 mg po qd			
<i>Adjunctive corticosteroid therapy for acute PCP with PaO₂ ≤ 70 mmHg</i>			
Prednisone po or methylprednisolone (Solu-Medrol) IV: 40 mg bid for 5 days followed by 40 mg qd for 5 days, followed by 20 mg qd for 11 days (can be tapered to 0 mg for last 11 days also)	21 days	Hyperglycemia, sodium retention, potassium wasting. Psychiatric syndromes. Exacerbation of Kaposi sarcoma, thrush, herpes infections, and other opportunistic infections	Corticosteroids indicated in conjunction with antipneumocystis therapy in all patients with PaO ₂ ≤ 70 mmHg. Begin corticosteroids concurrently with PCP treatment or if PaO ₂ decreases to ≤ 70 mmHg within 72 hours of initiating PCP treatment.

Continued

Table 1. Continued

System, Problem, and Drug Regimen	Duration	Adverse Effects/Drug Interactions	Comments
CENTRAL NERVOUS SYSTEM			
<i>Toxoplasma gondii</i>			
<i>Prophylaxis</i>			
Most PCP prophylaxis regimens provide some protection against toxoplasmosis	Indefinitely	See PULMONARY, PCP	TMP-SMX, dapsone plus TMP or pyrimethamine, clindamycin plus primaquine, atovaquone plus pyrimethamine, and pyrimethamine-sulfadoxine provide some prophylaxis against toxoplasmosis. Aerosolized pentamidine not effective; adding another agent to provide toxoplasmosis prophylaxis not required. Clarithromycin and azithromycin provide some benefit
<i>Acute</i>			
Pyrimethamine 75 - 100 mg po qd (every other day if bone marrow suppression) plus leucovorin calcium (folinic acid) 10 - 25 mg po qd plus either	6 - 8 weeks for acute therapy	Leukopenia, anemia, thrombocytopenia	Clinical response or regression of lesions on imaging studies is usually noted within 2 weeks. Maintenance required indefinitely to prevent relapse
Sulfadiazine 1.0 - 1.5 g po q 6 h	Same	Rash, drug fever; leukopenia, thrombocytopenia; crystalluria with renal failure	Sulfadiazine probably provides effective prophylaxis against PCP. Ensure adequate fluid intake
or			
Clindamycin 600 - 900 mg po or IV qid	Same	See PULMONARY, PCP	
<i>Alternative when intolerant of sulfadiazine and clindamycin</i>			
Pyrimethamine plus leucovorin as above	Same	See above	
plus one of the following			
Clarithromycin 1 g po bid or azithromycin 1200 - 1500 mg po qd	Same	See GENERAL/SYSTEMIC, MAC	
or			
Atovaquone suspension (750 mg/5 mL) 750 mg po qid with meals	Same	See PULMONARY, PCP	Not proved effective
or			
Doxycycline 100 mg po tid - qid or minocycline 200 mg po bid	Same	Tetracycline side effects	Not proved effective
or			
Dapsone 100 mg po qd	Same	See PULMONARY, PCP	
or			
Pyrimethamine alone 100 - 200 mg po qd	Same	See PULMONARY, PCP	Not as effective as above regimens
OR			
TMP/SMX as for acute PCP	Same	See PULMONARY, PCP	
<i>Maintenance</i>			
Pyrimethamine 25 - 50 mg po qd plus either	Indefinitely		Add leucovorin calcium if evidence of leukopenia
Sulfadiazine 1 g po q 12 h	Indefinitely		
or			
Clindamycin 300 - 450 mg po q 6 - 8 h	Indefinitely		

Continued

Table 1. Continued

System, Problem, and Drug Regimen	Duration	Adverse Effects/Drug Interactions	Comments
CENTRAL NERVOUS SYSTEM			
<i>Cryptococcus neoformans</i>			
<i>Prophylaxis</i> Fluconazole provides limited prophylaxis			Primary prophylaxis not routinely recommended. Can be considered for patients with CD4+ cell counts < 50/ μ L. No long-term survival benefit. Fluconazole resistance reported
<i>Meningitis or disseminated cryptococcosis</i> <i>Acute</i>			
Amphotericin B 0.7 - 1.0 mg/kg/d IV with or without 5-flucytosine (Ancobon) 100 mg/kg po qd in 4 divided doses for first 2 - 4 weeks. If clinically improved after 7.5 mg/kg total amphotericin B administration, can change to fluconazole 400 mg po qd or itraconazole 200 mg po bid	6 - 8 weeks; amphotericin total dosage not to exceed 2 g	Renal failure, hypokalemia, hypomagnesemia. Liposomal amphotericin B might decrease toxicity Fever, chills; anemia, thrombophlebitis Granulocytopenia; nausea, vomiting, diarrhea, aminotransferase elevations; rash from flucytosine Flucytosine toxicities (rash, metallic taste, leukopenia, thrombocytopenia) limit its usefulness	Pretreatment with diphenhydramine, acetaminophen or IV morphine can decrease amphotericin-induced fevers, chills, and rigors. Pretreatment not recommended routinely. Administer for 4 - 6 hours in D5W. Addition of heparin 500 U and hydrocortisone 25 mg to amphotericin IV solution can decrease phlebitis. Infusion of 500 - 1000 mL normal saline before administration of amphotericin B can minimize renal toxicity. 5-Flucytosine not indicated if granulocytopenia or thrombocytopenia is present Markedly increased intracranial pressure (> 240 mm) might require cerebrospinal fluid drainage (20 - 30 mL or more per day by lumbar puncture or continuous lumbar drain), or possibly corticosteroid, mannitol, or acetazolamide (Diamox) therapy
OR			
Fluconazole 400 - 800 mg po qd. Dosage reduction in renal failure. Higher dosages (eg, 800 - 1200 mg po qd) might increase efficacy	8 - 12 weeks	Nausea, vomiting, diarrhea; dizziness; aminotransferase elevations; rare cutaneous reactions, skin pigmentation, alopecia <i>Drug interactions</i> Increased phenytoin (Dilantin) and warfarin (Coumadin) levels; higher fluconazole dosages might be necessary for patients taking rifampin	As effective as amphotericin B against mild or moderate disease; unknown whether equally effective against severe disease. Fluconazole penetrates central nervous system and most body tissues, including prostate. Addition of 5-flucytosine might be of benefit
<i>Maintenance</i>			
Fluconazole 200 - 400 mg po qd	Indefinitely	Same	Higher dosages might be necessary for recurrent disease
OR			
Amphotericin B 0.5 - 0.8 mg/kg/d 3 - 5 times a week	Indefinitely	Same	
Syphilis			
Aqueous crystalline penicillin G 3 - 4 mU IV q 4 h (total 18 - 24 mU/d)	10 - 14 days	Usual penicillin adverse effects, Jarisch-Herxheimer reaction; seizures from high-dosage penicillin in renal failure	Continued serologic and clinical follow-up required to assess adequacy of treatment for neurosyphilis. Persons with ophthalmic auditory, or cranial nerve abnormalities or other syndromes consistent with neurosyphilis should receive daily penicillin therapy for 10 - 14 days. Intravenous penicillin preferred for adequate central nervous system penetration. For penicillin-allergic patients, consultation with an expert advised. Administer additional benzathine penicillin 2.4 mU IM weekly after completion of neurosyphilis treatment to ensure 3 weeks total penicillin therapy
OR			
Procaine penicillin G 2.4 mU IM qd	10 - 14 days	Same. Probenecid rash	
plus			
Probenecid 500 mg po qid	10 - 14 days		<i>Continued</i>

Table 1. Continued

System, Problem, and Drug Regimen	Duration	Adverse Effects/Drug Interactions	Comments
CENTRAL NERVOUS SYSTEM			
Peripheral neuropathy			
Desipramine (Norpramin) or amitriptyline (Elavil) 25 - 150 mg po hs	Indefinitely	Usual tricyclic side effects; drowsiness; orthostatic hypotension; anticholinergic symptoms	Pain relief occurs in 3 - 5 days. Desipramine causes less sedation and fewer anticholinergic effects. Other tricyclic drugs might be equally effective
Phenytoin (diphenylhydantoin, Dilantin) 100 mg po tid	Indefinitely	Usual side effects and drug-drug interactions	Generally ineffective
Carbamazepine (Tegretol) 100 - 300 mg po bid	Indefinitely	Leukopenia, bone marrow suppression, rare agranulocytosis; rash; drowsiness, dizziness; aminotransferase elevations	Less desirable because of bone marrow effects. Need to monitor carbamazepine levels to avoid toxicity
Mexiletine (Mexitil) 150 mg po bid - tid	Indefinitely	Nausea, vomiting, epigastric pain; dizziness, tremor; bradycardia; rare seizures, leukopenia, agranulocytosis	Less desirable because of side effects
Capsaicin (Axsain, Zostrix-HP) 0.075% topical cream qid	Indefinitely	Minor burning sensation, skin irritation, erythema	Pain relief delayed 2 - 4 weeks. No systemic effects
Gabapentin (Neurontin) 300 - 400 mg po tid via dose escalation; dosage reduction in renal failure	Indefinitely	Thrombocytopenia; somnolence, dizziness, ataxia, nystagmus, fatigue, somnolence, headache; nausea, vomiting, diarrhea	Can be helpful when other agents fail

principally in order of efficacy. Where more than one drug is effective for a specific condition, recommendations are usually in an order that favors the least expensive choice among equally effective options. The most common and clinically important adverse effects and drug interactions are listed.

References

A selected bibliography highlights the most important management and therapeutic problems associated with HIV infection and AIDS. References including articles about *P carinii* pneumonia²¹ and other mycobacterial diseases²²; dermatologic,²³⁻²⁵ oropharyngeal,^{26,27} ophthalmologic,²⁸⁻³⁰ and gastrointestinal problems^{31,32}; the AIDS wasting syndrome³³⁻³⁵; and neurologic disease.^{36,37} Additional references are intended to assist primary care clinicians³⁸ with the broad spectrum of problems associated with HIV infection and AIDS,^{39,40} other sexually transmitted diseases,⁴¹ and special treatment considerations for occupational exposures⁴²⁻⁴⁵ and pregnancy.⁴⁶

Other Sources

Information about clinical trials is available through the AIDS Clinical Trials Information Service at 1-800-TRIALS A, and through the AIDS Treatment Information Service (ATIS) at 1-800-HIV-8440. ATIS also has printed and Website guidelines and information about approved therapies and management protocols at <http://hivatis.org>. Our National HIV Telephone Consultation Service (Warmline) at the University of California, San Francisco, Department of Family and Community Medicine at San Francisco General Hospital provides clinical consultation and education for health care providers; the Warmline is in operation on weekdays at 1-800-933-3413. The AIDS Education and Training Centers (AIDS ETCs) of the Health Resources and Services Administration (HRSA) at 1-301-443-6364 offers education, training, and consultation services to health care providers. HRSA also offers special-topic teleconferences. Additional information can be accessed at other Websites, including the San Francisco General Hospital-based site at <http://www.hivinsite.ucsf.edu> and the American Medical Association HIV/AIDS site at <http://www.ama-assn.org>.

We gratefully acknowledge the staff of the HIV Telephone Consultation Service and the faculty, staff, and house staff at San Francisco General Hospital for making this work possible, and Mary A. Hanville for assistance in preparation of this manuscript.

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