

HIGHLIGHTS OF PRESCRIBING INFORMATION

These highlights do not include all the information needed to use ATZANAVIR capsules safely and effectively. See full prescribing information for ATZANAVIR capsules for complete information about ATZANAVIR capsules, for use as initial HIV-1 therapy.

RECENT MAJOR CHANGES

Contraindications (4) 09/2020
Warnings and Precautions 09/2020
Immune Reconstitution Syndrome (5.10)

INDICATIONS AND USAGE

ATZANAVIR capsules are a protease inhibitor indicated for use in combination with other antiretroviral agents for the treatment of HIV-1 infection in adults and in pediatric patients 6 years of age and older weighing at least 15 kg (11).

CONTRAINDICATIONS

ATZANAVIR capsules are contraindicated in patients with previously demonstrated hypersensitivity (eg, Stevens-Johnson syndrome, erythema multiforme, or toxic skin eruptions) to any of the components of this product (4).

ATZANAVIR capsules are contraindicated in patients who are co-administered with rifabutin, quinidine (if atazanavir is co-administered with ritonavir), rifampin, orally administered midazolam, ergot derivatives, ranolazine, prochlorperazine, lurasidone (if atazanavir is co-administered with ritonavir), and toxic skin eruptions, including drug-induced hypersensitivity syndrome (4).

WARNINGS AND PRECAUTIONS

Cardiac conduction abnormalities: Prolongation may occur in some patients. ECG monitoring should be considered in patients with preexisting conduction abnormalities or when administered with other drugs that may prolong the QT interval (5.1).

ADVERSE REACTIONS

Most patients experience asymptomatic increases in indirect bilirubin, which is reversible upon discontinuation. Do not dose reduce. In a concomitant transaminase increase occurs, evaluate for alternative etiologies (5.2).

Hypotension: Patients with hepatitis B or C or infection are at risk of increased transaminase or hepatic decompensation. Monitor hepatic laboratory tests prior to therapy and during treatment (2.8, 5.4, 8.8).

Chronic kidney disease has been reported during postmarketing surveillance in patients with HIV-1 infection treated with atazanavir, with or without ritonavir. Consider alternatives in patients at high risk for renal disease or preexisting renal disease. Monitor renal laboratory tests prior to therapy and during treatment. Consider discontinuation of atazanavir in patients with progressive renal disease (5.5, 8.3).

Methylophilias and cholelithiasis have been reported. Consider temporary interruption or discontinuation (5.6).

The concurrent use of atazanavir with ritonavir and certain other medications may result in potentially significant drug interactions. Consult the full prescribing information prior to and during treatment for potential drug interactions (5.7, 7.3).

Patients receiving atazanavir may develop new onset or exacerbations of diabetes mellitus/hyperglycemia (5.9), immune reconstitution syndrome (6.10), and redistribution/accumulation of body fat (6.11).

Hemophilia: Spontaneous bleeding may occur and additional factor VIII may be required (5.12).

Most common adverse reactions (≥2%) are nausea, jaundice/cholelithiasis, rash, headache, abdominal pain, vomiting, insomnia, peripheral neuropathy, symptoms, dizziness, myalgia, diarrhea, depression, and fever (6.1).

Adverse reactions: ATZANAVIR is not recommended for use in patients with moderate to severe hepatic impairment (2.8, 8.1) or FDA H1-1400-FA-1088 or FDA H1-1400-FA-1089 or FDA H1-1400-FA-1090 or FDA H1-1400-FA-1091 or FDA H1-1400-FA-1092 or FDA H1-1400-FA-1093 or FDA H1-1400-FA-1094 or FDA H1-1400-FA-1095 or FDA H1-1400-FA-1096 or FDA H1-1400-FA-1097 or FDA H1-1400-FA-1098 or FDA H1-1400-FA-1099 or FDA H1-1400-FA-1100 or FDA H1-1400-FA-1101 or FDA H1-1400-FA-1102 or FDA H1-1400-FA-1103 or FDA H1-1400-FA-1104 or FDA H1-1400-FA-1105 or FDA H1-1400-FA-1106 or FDA H1-1400-FA-1107 or FDA H1-1400-FA-1108 or FDA H1-1400-FA-1109 or FDA H1-1400-FA-1110 or FDA H1-1400-FA-1111 or FDA H1-1400-FA-1112 or FDA H1-1400-FA-1113 or FDA H1-1400-FA-1114 or FDA H1-1400-FA-1115 or FDA H1-1400-FA-1116 or FDA H1-1400-FA-1117 or FDA H1-1400-FA-1118 or FDA H1-1400-FA-1119 or FDA H1-1400-FA-1120 or FDA H1-1400-FA-1121 or FDA H1-1400-FA-1122 or FDA H1-1400-FA-1123 or FDA H1-1400-FA-1124 or FDA H1-1400-FA-1125 or 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