Role of atazanavir in the treatment of HIV infection

Pablo Rivas1
Judit Morello2
Carolina Garrido1
Sonia Rodríguez-Nóvoa2
Vincent Soriano1
1Department of Infectious Diseases; 2Pharmacology Unit, Hospital Carlos III, Madrid, Spain

Abstract: Atazanavir (ATV) is one of the latest protease inhibitors (PI) approved for the treatment of HIV infection. The drug has a relatively long-life (∼7 h) and large inhibitory quotient which allows once daily administration. It is generally well tolerated and the main side effect is hyperbilirubinemia, since ATV inhibits the hepatic uridin-glucoronyl-transferase. A signature mutation at the protease gene, I50L, may confer loss of susceptibility to the drug. Interestingly, it produces hypersusceptibility to all other PIs. When ATV is pharmacokinetically boosted with ritonavir (r) 100 mg/day, a greater genetic barrier for resistance is achieved, and generally more than 3 major PI resistance associated mutations are needed to result in ATV resistance. In drug-naïve subjects, regimens based on ATV/r have shown non-inferiority compared to lopinavir (LPV)/r (CASTLE study) or fosamprenavir/r (ALERT trial), generally with improved tolerance (less diarrhea and dyslipidemia). Given its good tolerance and convenience, ATV has been proven to be successful as a simplification strategy in switch studies (ie, SWAN and SLOAT) conducted in patients with complete virological suppression under other PI-based regimens. Finally, ATV/r-based combinations have shown to be equivalent in terms of viral response to other PI/r-containing regimens, including LPV/r, in rescue interventions in patients failing other PI regimens (ie, studies AI424-045 and NADIS).

Keywords: atazanavir, HIV, protease inhibitors, drug resistance

Introduction

The introduction of triple combination therapy following the availability of protease inhibitors (PIs) dramatically changed the natural history of HIV infection in the late 1990s. When the first PI (saquinavir) was marketed in 1995, the recognition of significant gains in CD4 counts and dramatic reductions in the incidence of opportunistic events accompanying by unprecedented declines in virus replication produced a huge enthusiasm in both patients and clinicians. However, this initial joy was soon tempered when patients began to complain of difficulties in pursuing treatment schedules and especially when side effects became clearly manifest. Even worse was the recognition that disturbances in the metabolism of lipids and glucose and disfiguring morphological features due to fat tissue redistribution were a new stigmatizing feature following prolonged PI exposure.

A new generation of PI compounds free of the main limitations of the first-generation PIs has recently entered the HIV armamentarium. ATV, marketed as Reyataz® (Bristol-Myers Squibb), may have an advantage over other PIs because of its favorable effect on lipid profiles, once-daily dosing, low capsule burden and a relatively favorable resistance profile.1 This article reviews the main pharmacologic and clinical features of ATV and updates its role in the treatment of HIV infection.

Mechanism of action

ATV is an azapeptide inhibitor of the HIV-1 protease. The chemical name for ATV sulfate is (3S,8S,9S,12S)-3,12-bis(1,1-dimethylethyl)-8-hydroxy-4,11-dioxo-9-(phenylmethyl)-6-((4-(2-pyridinyl)phenyl)methyl)-2,5,6,10,13-pentazatetradecanedioic acid dimethyl ester sulfate (1:1) (Figure 1). The compound inhibits the virus-specific processing of viral Gag and Gag-Pol poliproteins of HIV-1 group M subtype A, B, C, D, AE, AG,
F, G, and J in infected cells, thus preventing formation of mature virions.\(^1\) As measure of potency, the concentration that inhibits 50% of viral replication (IC50) in the absence of human serum ranged from 0.58 ng/mL to 5.7 ng/mL in a panel of susceptible viruses isolated from 31 PI-naive HIV-infected patients.\(^2,3\) The presence of 40% human serum in cell cultures increased ATV IC50 by 2.7- to 3.6-fold, as noticed for other PIs. The adjusted IC50 for protein binding was estimated to range from 8 to 20 ng/mL against reference viral strains with a conventional cycle cell infection and the PhenoSense\textsuperscript{TM} single assay (ViroLogic Inc., South San Francisco, CA, USA), respectively.\(^2\)

**Drug resistance**

Resistance patterns to ATV differ according to the population exposed to the drug being PI-naive or -experienced, and to ritonavir (r) boosting. The presence of a single major mutation in the protease gene may result in loss of susceptibility to ATV, but in clinical practice ATV resistance generally occurs when several mutations in the protease gene are present. In PI-naive patients, the most frequent mutation at failure under ATV is I50L,\(^4,5\) while in PI-experienced patients mutations I84V and N88S are more commonly selected. Of note, I50L is only selected under ATV pressure and it causes higher susceptibility to other PIs such as amprenavir, darunavir, indinavir, lopinavir (LPV), nelfinavir (NFV) and saquinavir (SQV).\(^6\) The prevalence of I50L in large HIV drug resistance mutation databases is generally very low.\(^7\)

ATV is almost always prescribed boosted with r, but the Food and Drug Administration (FDA) allows also its use unboosted in selected PI-naive patients and in simplification strategies. In contrast, the EMEA has not approved yet the use of ATV without r boosting. When PIs are used without r boosting, drug exposures are significantly lower. In this situation, the resistance barrier may be confined to a single key mutation (eg, I50L).

Interpretation of drug resistance mutations is complicated. First, a qualitative approach is usually performed identifying changes that could affect drug susceptibility. As not all changes in the protease gene have the same influence on drug susceptibility, a quantitative score giving different weight to each mutation and each drug must be used. The International AIDS Society-USA (IAS-USA) Panel describes a list of mutations in the protease gene that are associated with resistance to ATV. This list includes changes at positions 10, 16, 20, 24, 32, 33, 34, 36, 46, 48, 50, 53, 54, 60, 62, 64, 71, 73, 82, 84, 85, 88, 90 and 93, although the only changes considered as major mutations are I50L, I84V and N88S.\(^8\) Table 1 records different algorithms developed to assist in the interpretation of resistance to ATV.

The accumulation of multiple PI resistance mutations has been shown to be a major indicator of ATV loss of susceptibility. In vitro analysis of the genotypic profiles of 943 PI-susceptible and PI-resistant clinical isolates identified a strong correlation between the presence of several amino acid changes at specific residues (10I/V/F, 20R/M/I, 24I, 33I/F/V, 36I/L/V, 46I/L, 48V, 54V/L, 63P, 71V/T/I, 73C/S/T/A, 82A/F/S/T, 84V, and 90M) and decreased susceptibility to ATV.\(^2\) In addition, data from PI-experienced patients revealed that the median number of PI-associated resistance mutations was lower in patients showing virological response than in non-responders.\(^9\)

As with other PIs, failure of ATV is often rather better explained by lack of potency than by the acquisition of primary resistance mutations, especially in patients without previous PI failure.\(^10\) In PI-experienced patients, the genotypic inhibitory quotient (GIQ) has been proven to be a good predictor of virological response to ATV, as it integrates both resistance mutations and drug plasma exposure.\(^9\)

**Cross-resistance**

Although there is no obvious overlapping resistance pattern of ATV with any other particular PI, cross-resistance with ATV is observed in isolates resistant to 4 or more PIs.\(^2\) The NADIS French study proved that ATV/r-based rescue therapy is generally efficient in most PI-experienced patients, except in those who had failed an LPV/r-containing regimen.\(^11\) By contrast, mutation L76V, selected under virologic failure with other PIs, produces hypersusceptibility to ATV.\(^12\)

**Resistance to atazanavir in HIV-1 non B subtypes and in HIV-2**

HIV variants other than subtype B show a high genetic variability within the protease, presenting many polymorphisms at positions associated with PI resistance in subtype B.
Nevertheless, primary resistance mutations do not appear as natural polymorphisms to ATV either in HIV-1 non-B subtypes or in HIV-2 isolates. Indeed, 48-week data from the CASTLE study have confirmed that virologic response to ATV is independent from HIV subtype.\(^{13}\)

**Pharmacokinetics**

ATV is rapidly absorbed with a peak plasma concentration (C\(_{\text{max}}\)) occurring at approximately 2.5 hours and demonstrates non-linear pharmacokinetics. The extent of absorption is highly dependent on gastric pH and increases when taken together with a light meal. ATV is 86% bound to human serum proteins to both alpha-1-acid glycoprotein and albumin to a similar extent. ATV is a substrate for P-glycoprotein (P-gp), an efflux transporter that will act to limit tissue compartment distribution. Like other PIs, ATV is extensively metabolized by hepatic cytochrome P450, primarily the CYP3A4/CYP3A5 isoenzymes. ATV and its metabolites then undergo biliary excretion for 20% and 7% of the administered dose in feces and urine, respectively. Unchanged drug accounts approximately 86% and 86% of an administered dose, respectively. Steady-state is achieved between days 4 and 8, with an accumulation of approximately 2.3-fold. Finally, ATV scarcely enters the cerebrospinal fluid (CSF) and semen, with a CSF/plasma ratio ranging from 0.11 to 4.42.\(^{1}\)

A strategy to enhance PI pharmacokinetic parameters consists of the co-administration of low doses of r (boosting). As r is a potent inhibitor of CYP3A and to a less extent of P-gp, co-administration of ATV and low doses of r (100 mg/day) result in a decrease in ATV gastrointestinal clearance and first-pass hepatic effect. Consequently, ATV normal half-life (\(\sim\)7 h) is increased resulting in higher minimum concentration (C\(_{\text{min}}\)), C\(_{\text{max}}\) and AUC.\(^{1}\) Table 2 depicts the main pharmacokinetic parameters of ATV 400 and ATV/r 300/100 mg/day.\(^{1,14}\)

There is large inter-individual variability in ATV disposition. In a population pharmacokinetic study which included 214 HIV patients, multiple factors that potentially could influence the variability of ATV pharmacokinetics were assessed, including body weight, sex, ethnicity, creatinine clearance, and concomitant medications (r, nevirapine, tenofovir, abacavir, lamivudine and proton pump inhibitors). Only body weight, r and nevirapine appeared to be highly correlated with ATV clearance.\(^{15}\)

**Pharmacokinetics in special populations**

**Pediatric**

ATV has not been fully evaluated in pediatric patients and the optimal dosage has not been established. In the phase I/II

<table>
<thead>
<tr>
<th>Source</th>
<th>Protease inhibitor</th>
<th>Protease mutations</th>
<th>Clinical cut-off</th>
</tr>
</thead>
<tbody>
<tr>
<td>Colonno et al(^{2})</td>
<td>Atazanavir</td>
<td>L10F/I/V, K20I/M/R, L24I, L33F/I/V, M36I/L/V, M46I/L, G48V, I54L/V, L63P, A71I/V/T, G73A/C/S/T, V82A/F/T/S, I84V, L90M, or the presence of I50L alone</td>
<td>&lt;versus ℝ 4</td>
</tr>
<tr>
<td>ANRS 2005(^{22})</td>
<td>Atazanavir–ritonavir</td>
<td>L10F/I/V, K20I/M/R, L24I, L33F/I/V, M46I/L, D60E, I84V, I85V, L90M, or the presence of I50L alone</td>
<td>&lt;versus ℝ 3</td>
</tr>
<tr>
<td>Pellegrin et al(^{26})</td>
<td>Atazanavir–ritonavir</td>
<td>L10F/I/V, K20I/M/R, L24I, M46I/L, I54A/M/V, V82A/F/T/S, I84V, L90M, or the presence of I50L alone</td>
<td>&lt;versus ℝ 5</td>
</tr>
<tr>
<td>Bertoli et al(^{83})</td>
<td>Atazanavir</td>
<td>L10C/I/V, V32I, E34Q, M46I/L, F53L, I54A/M/V, V82A/F/T/S, I84V, L90M, or the presence of I50L alone</td>
<td>&lt;versus ℝ 4</td>
</tr>
<tr>
<td>Bertoli et al(^{83})</td>
<td>Atazanavir–ritonavir</td>
<td>G16V, V32I, K20I/M/R/T/V, L33F/I/V, M46I/L, I54L/V, Q58E, L63P, A71I/V/T, G73A/C/S/T, V82A/F/T/S, I84V, L90M, or the presence of I50L alone</td>
<td>&lt;versus ℝ 3</td>
</tr>
</tbody>
</table>

**Abbreviation:** ANRS, agence nationale de recherche sur le sida et les hépatites Virales.

**Table 1 Different genotypic resistance scores for atazanavir (boosted or unboosted) in relation to clinical responses**

<table>
<thead>
<tr>
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<th>Protease inhibitor</th>
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<th>Clinical cut-off</th>
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<tr>
<td>Colonno et al(^{2})</td>
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<td>L10F/I/V, K20I/M/R, L24I, L33F/I/V, M36I/L/V, M46I/L, G48V, I54L/V, L63P, A71I/V/T, G73A/C/S/T, V82A/F/T/S, I84V, L90M, or the presence of I50L alone</td>
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</tr>
<tr>
<td>ANRS 2005(^{22})</td>
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<td>L10F/I/V, K20I/M/R, L24I, L33F/I/V, M46I/L, D60E, I84V, I85V, L90M, or the presence of I50L alone</td>
<td>&lt;versus ℝ 3</td>
</tr>
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<td>&lt;versus ℝ 5</td>
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</tr>
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**Table 2 Pharmacokinetic parameters at steady state after atazanavir 400 mg once daily and after atazanavir 300 mg with ritonavir 100 mg once daily with a light meal in HIV-infected patients**

<table>
<thead>
<tr>
<th>Bioavailability (%)</th>
<th>ATV 400 mg</th>
<th>ATV/r 300/100 mg</th>
</tr>
</thead>
<tbody>
<tr>
<td>Protein binding (%)</td>
<td>86</td>
<td>86</td>
</tr>
<tr>
<td>Distribution volume</td>
<td>Not available</td>
<td>Not available</td>
</tr>
<tr>
<td>Clh (L/h)</td>
<td>25.2</td>
<td>25.2</td>
</tr>
<tr>
<td>Clr of parent drug (%)</td>
<td>7</td>
<td>Not available</td>
</tr>
<tr>
<td>t(_{1/2}) mean ± SD (h)</td>
<td>6.5 ± 2.6</td>
<td>6.5 ± 2.6</td>
</tr>
<tr>
<td>C(_{\text{max}}) mean ± SD (ng/mL)</td>
<td>3152 ± 2231</td>
<td>5233 ± 3033</td>
</tr>
<tr>
<td>T(_{\text{max}}) median (h)</td>
<td>2</td>
<td>3</td>
</tr>
<tr>
<td>C(_{\text{min}}) mean ± SD (ng/mL)</td>
<td>273 ± 298</td>
<td>862 ± 838</td>
</tr>
<tr>
<td>AUC, mean ± SD (ng/mL/h)</td>
<td>2226 ± 20159</td>
<td>53761 ± 35294</td>
</tr>
</tbody>
</table>

**Abbreviations:** ATV, atazanavir; RTV, ritonavir; Clh, hepatic clearance; Clr, renal clearance; T\(_{1/2}\), half-life; C\(_{\text{max}}\), maximum concentration; T\(_{\text{max}}\), time at C\(_{\text{max}}\); C\(_{\text{min}}\), minimum concentration; AUC, area under the curve.
PATCG 1020A study, ATV was dosed at 310 mg/m² body surface area and adjusted over 24 h in 172 children with a mean age of 1.2 years up to 17 years. AUC presented a wide range between 33,800 ng/mL/h in infants with a mean age of 1.2 years and 67,100 ng/mL/h in children with a mean age of 15 years. The $C_{\text{max}}$ values were 3900 ng/mL in adolescents (mean age, 17 years) and 7400 ng/mL in the youngest children (mean age, 1.2 years).16

**Hepatic impairment**

Although few data have been derived from HIV patients with severe hepatic dysfunction, an increased exposure to the drug in such individuals is anticipated. In non-HIV infected adults with moderate to severe hepatic impairment, $AUC_{(0-\infty)}$ was increased in 42% after a single 400-mg dose compared with healthy volunteers. On the basis on these data, a dose reduction of ATV is advised for patients with moderate hepatic dysfunction.1 In contrast, a study conducted in 58 HCV/HIV-coinfected patients with compensated liver disease, ATV $C_{\text{min}}$ did not differ significantly between patients with and without liver cirrhosis.17

**Renal impairment**

ATV pharmacokinetics have been evaluated in 20 adults with severe renal impairment, including a few on hemodialysis, using multiple doses of 400 mg once daily. The mean ATV $C_{\text{max}}$ was 9% lower, $AUC$ was 19% higher, and $C_{\text{min}}$ was 96% higher in subjects with severe renal impairment not undergoing hemodialysis, than in subjects with normal renal function. When ATV was administered either before or after hemodialysis, the geometric means for $C_{\text{max}}$, $AUC$, and $C_{\text{min}}$ were approximately 25% to 43% lower compared to subjects with normal renal function. No dose adjustment is required in patients who are not managed with hemodialysis and HIV patients with end stage renal disease managed with hemodialysis should receive ATV 300 with r 100 mg daily. ATV should not be administered to antiretroviral-experienced patients with end stage renal disease managed with hemodialysis.1

**Pregnancy**

Although there are no adequate pharmacokinetic studies in pregnant women, data from other PIs suggest a decrease in plasma exposure during the third trimester.18 A pharmacokinetic study was performed in 12 HIV-pregnant women receiving ATV/r 300/100 mg/day. The third trimester $AUC$ and $C_{\text{max}}$ were approximately 40% and 21% lower, respectively, than in adult HIV-infected patients. All subjects reaching delivery achieved plasma HIV-RNA <50 copies/mL and all infants tested were HIV-negative and presented normal bilirubin values through day 14. However, 1 newborn developed grade 3 hyperbilirubinemia at day 15.19 In another study, total bilirubin concentrations were above normal limits at birth and day 3, and 3 newborns had transient neonatal jaundice which did not require phototherapy.20

**Drug interactions**

ATV is a substrate and inhibitor of CYP3A (Ki value of 0.84 to 1.0 μM). ATV also inhibits CYP2C8 (Ki = 2.1 μM) and UGT1A1 ($K_i = 1.9$ μM). It is also a substrate and inhibitor of P-gp with levels comparable to verapamil, a well known P-gp inhibitor.21,22 Therefore, co-administration of ATV and drugs primarily metabolized by CYP3A, CYP2C8, or UGT1A1 and/or substrates of P-gp may result in increased plasma concentrations of the concomitant drug, which potentially could enhance or prolong both their therapeutic and adverse effects. The magnitude of the mediated drug interaction may change when ATV is co-administered with r. On the other hand, drugs that induce CYP3A4 and/or P-gp may decrease ATV plasma concentrations and therefore compromise ATV therapeutic effect. Finally, drugs affecting the gastric pH may alter ATV solubility and consequently its bioavailability.1 ATV and ATV/r drug interactions are summarized in Table 3.

As shown in Figure 2, raltegravir, the first integrase inhibitor approved for the treatment of HIV infection, is metabolized by glucuronidation, but interactions with ATV do not generally result in increased hyperbilirubinemia since different UGT isoenzymes are subject to ATV inhibition and raltegravir metabolism, respectively.

**Therapeutic drug monitoring (TDM)**

Monitoring plasma drug concentrations has proven to be useful for PIs, and may allow tailoring of antiretroviral therapy.23 ATV plasma concentrations demonstrate high inter-individual variability, even in the presence of r, along with a low intra-individual variability, supporting TDM. The definition of a therapeutic range for ATV concentrations has been pursued for a while in an attempt to minimize side effects, mainly hyperbilirubinemia while ensuring maximal efficacy, especially when r unboosted ATV needs to be used.24

**Atazanavir plasma concentrations as predictors of virological response**

Several studies have evaluated the relationship between ATV plasma levels and virological response.9,25,26 A threshold of 0.15 mg/L has been proposed for antiretroviral-naïve patients based on results of a retrospective study performed
on 51 patients. In that study, virological responses according to ATV trough concentrations (C_{trough}) were as follows: 58.3% when <0.15 mg/L, 75% when values between 0.15 and 0.85 mg/L, and 100% when ≥0.85 mg/L.

In PI-experienced patients, the combined use of pharmacokinetics and resistance information has been proposed as a better predictor of virological response than use of these two variables separately. The genotypic inhibitory quotient (GIQ), defined as the ratio of the minimum plasma drug concentration to the number of resistance mutations in the protease gene, has been associated with virological response to ATV in PI-experienced patients. A GIQ > 0.10 mg/L mutation has been found to significantly predict virological response at 24 weeks. Mutations more significant to be included in the GIQ model are the following: L10F/I/V, K20M/R, L24I, D30N, V32I, L33F, I47V/A, G48V, V77I, V82A/F/T, G73S, G77V, L90M.

**Atazanavir plasma levels as predictors of hyperbilirubinemia**

Several studies have found an association between ATV plasma concentrations and serum bilirubin levels. However, at this time there are not enough data available to define threshold for scleral jaundice, which otherwise may vary from one subject to another, although it can generally be recognized when serum bilirubin goes up 2.5 mg/dL.

In HIV-HCV-coinfected patients on stable ATV therapy who begin treatment with pegylated interferon plus ribavirin, hemolytic anemia caused by ribavirin may result in increased levels of bilirubin (Figure 2). In a study performed in 72 HCV/HIV-coinfected patients treated for hepatitis C, up to 45% of those on ATV experienced serum bilirubin increases greater than 1 mg/dL compared with only 3% of individuals under other antiretroviral regimens. Indeed, the proportion of patients on ATV who experienced hyperbilirubinemia grade 3–4 increased 2.5-fold after beginning hepatitis C therapy (from 9% to 45%).

Another population in which TDM for ATV may be particularly useful is the subset of HIV patients treated concomitantly with TDF, the most widely used antiretroviral agent. The co-administration of ATV and TDF results in a 20% to 40% reduction in ATV plasma concentrations. Therefore, the current advice is to give ATV boosted with low doses of r when taken along with TDF. However, this intervention may be regarded as less convenient for drug-naïve subjects because of the need for an additional extra pill of r, which has to be kept in the refrigerator and often produces gastrointestinal disturbances. More importantly, it increases the risk of jaundice and eliminates the lipid-friendly behavior of ATV. In a retrospective study performed in 56 HIV patients with undetectable plasma HIV-RNA under a triple regimen containing ATV/r 300/100 mg once daily (qd), TDM was useful to allow a switch to ATV 400 mg qd. A total of 79% were taking concomitantly TDF. At week 24, therapeutic levels (C_{trough} > 150 ng/mL) were kept in all patients but 4. TDF was part of the antiretroviral regimen in all 4 cases. From a total of 29 (52%) patients on ATV/RTV showing grade 3–4 hyperbilirubinemia, only 7 (12%) still suffered from it upon switching to ATV 400 mg qd. Undetectable viral load was maintained in all patients but one in whom subtherapeutic ATV concentrations were seen. Therefore, TDM can be a useful tool to identify the subset of ATV treated patients in whom removal of r could be attempted without risk of suboptimal plasma ATV exposure and subsequent virological failure.

**Pharmacogenetics**

Hyperbilirubinemia is the most common adverse effect in patients treated with ATV. The underlying mechanism is an inhibition by ATV of the uridine-glucuronosyl-transferase (UGT) 1A1 enzyme, which is involved in bilirubin conjugation (Figure 2). In this context, genetic factors affecting ATV disposition and/or UGT1A1 function could alter bilirubin levels.

Several single nucleotide polymorphisms (SNP) with potential consequences in the expression of P-gp have been identified in the gene that codifies for P-gp, named multidrug resistance gene 1 (MDR1). In patients taking ATV 400 mg qd, the 3435C > T polymorphism has been associated with lower ATP plasma levels and lower serum bilirubin levels. Accordingly, the risk for developing severe hyperbilirubinemia was as high as 24% in subjects with MDR1 wild-type alleles but as low as 0% in those homozygous for the 3435C > T polymorphism. These results have been confirmed thereafter in patients on ATV/RTV, as patients with the CC genotype showed significantly higher median ATV C_{trough} concentrations than those with CT/TT genotypes (939 versus 376 ng/mL, respectively). The C3435T polymorphism is a silent change and thus it is unlikely that it directly influences the expression of the MDR1 gene. Most likely, this polymorphism may be linked to one or more other unidentified changes in MDR1 or distinct metabolizing enzyme genes. The 2677G > T polymorphism has recently been suggested as the possible candidate. Persons homozygous for the MDR1 2677T allele, which is frequently linked to the 3435T allele, show enhanced
<table>
<thead>
<tr>
<th>Family</th>
<th>Drug</th>
<th>Effect on concentration</th>
<th>Recommendation</th>
</tr>
</thead>
<tbody>
<tr>
<td>Antiarrhythmics</td>
<td>amiodarone, bepridil, lidocaine, quinidine</td>
<td>↑ antiarrhythmic</td>
<td>Caution is warranted, TDM recommended.</td>
</tr>
<tr>
<td>Anticoagulants</td>
<td>warfarin</td>
<td>↑ warfarin</td>
<td>Monitoring of INR is recommended.</td>
</tr>
<tr>
<td>Anticonvulsants</td>
<td>carbamazepine, lamotrigine, phenobarbital, phenytoin</td>
<td>↓ trazodone</td>
<td>Use with caution.</td>
</tr>
<tr>
<td>Antidepressants</td>
<td>tricyclic antidepressants trazodone</td>
<td>↑ tricyclic antidepressants trazodone</td>
<td>Use with caution, TDM is recommended. TDM is recommended and lower trazodone doses should be used</td>
</tr>
<tr>
<td>Antifungals</td>
<td>itraconazole, ketoconazole</td>
<td>↑ itraconazole, ketoconazole (ATV 400)</td>
<td>If ATV is used with RTV, itraconazole or ketoconazole doses of &gt;200 mg/day should be used with caution.</td>
</tr>
<tr>
<td>Antihistamines</td>
<td>astemizole, terfenadine</td>
<td>no data available</td>
<td>ATV/r should not be used in combination with drugs that are substrates of the CYP3A4 and have narrow therapeutic windows, such as terfenadine and astemizole.</td>
</tr>
<tr>
<td>Antimicrobial agents</td>
<td>clarithromycin</td>
<td>↑ clarithromycin ↓ I4-OH-clarithromycin ↑ ATV concentrations</td>
<td>Dose reductions by 50% should be considered. Combination with boosted ATV has not yet been studied.</td>
</tr>
<tr>
<td>Antineoplastics</td>
<td>irinotecan</td>
<td>↑ irinotecan</td>
<td>Contraindicated.</td>
</tr>
<tr>
<td>Antimycobacterials</td>
<td>rifabutin</td>
<td>↑ rifabutin</td>
<td>Reduce rifabutin dose to 150 mg every other day or 3 times/week.</td>
</tr>
<tr>
<td></td>
<td>rifampicin</td>
<td>severe ↓ ATV</td>
<td>Contraindicated.</td>
</tr>
<tr>
<td>Antiretroviral agents</td>
<td>NRTIs: didanosine</td>
<td>↓ ATV, ↓ didanosine</td>
<td>Didanosine should be administered (with food) 1 h before or 2 h after ATV/r intake. Avoid combination of tenofovir with r unboosted ATV.</td>
</tr>
<tr>
<td></td>
<td>tenofovir</td>
<td>↓ ATV, ↑ tenofovir</td>
<td>The recommended dose in treatment-naïve patients is ATV/r 300/100 mg/day. No recommendation has been established in treatment-experienced patients.</td>
</tr>
<tr>
<td></td>
<td>NNRTIs: efavirenz</td>
<td>↓ ATV</td>
<td>Co-administration is not recommended. ATV should be boosted with r.</td>
</tr>
<tr>
<td></td>
<td>Nevirapine</td>
<td>expected ↓ ATV</td>
<td>Contraindicated due to synergistic effect on hyperbilirubinemia.</td>
</tr>
<tr>
<td></td>
<td>etravirine</td>
<td>↑ etravirine, ↓ ATV</td>
<td>Appropriate recommendations for this combination have not been established. TDM is recommended.</td>
</tr>
<tr>
<td></td>
<td>IPs: indinavir</td>
<td></td>
<td>ATV and tipranavir should not be co-administered.</td>
</tr>
<tr>
<td></td>
<td>saquinavir (soft gelatin capsules)</td>
<td>↑ saquinavir</td>
<td>The clinical relevance of these data is unknown. No changes in raltegravir design are recommended.</td>
</tr>
<tr>
<td></td>
<td>tipranavir</td>
<td>expected ↓ ATV</td>
<td>Contraindicated.</td>
</tr>
<tr>
<td></td>
<td>INI: raltegravir</td>
<td>↑ raltegravir</td>
<td>Contrainindicated.</td>
</tr>
<tr>
<td>Benzodiazepines</td>
<td>midazolam, triazolam</td>
<td>↑ midazolam, triazolam</td>
<td>Contraindicated.</td>
</tr>
<tr>
<td>Calcium channel blockers</td>
<td>diltiazem</td>
<td>↑ diltiazem and desacetyl-diltiazem</td>
<td>Caution is warranted. 50% dose reduction of diltiazem should be considered.</td>
</tr>
<tr>
<td></td>
<td>felodipine, nifedipine, nicardipine, verapamil</td>
<td>↑ felodipine, nifedipine, nicardipine, verapamil</td>
<td>Caution is warranted and ECG monitoring is recommended. Dose titration should be considered.</td>
</tr>
<tr>
<td>Corticosteroids</td>
<td>fluticasone</td>
<td>↑ fluticasone</td>
<td>Caution is warranted.</td>
</tr>
<tr>
<td>Ergot derivatives</td>
<td>dihydroergotamine, ergotamine, ergonovine, methylergonovine</td>
<td>↑ ergot derivatives</td>
<td>Contraindicated.</td>
</tr>
<tr>
<td>GI motility agent</td>
<td>cisapride</td>
<td>↑ cisapride</td>
<td>Contraindicated.</td>
</tr>
<tr>
<td>Acid suppressive therapy</td>
<td>antaciads</td>
<td>↓ ATV</td>
<td>ATV should be administered 2h before or 1h after intake of antaciads.</td>
</tr>
</tbody>
</table>

(Continued)
Table 3 (Continued)

<table>
<thead>
<tr>
<th>Family</th>
<th>Drug</th>
<th>Effect on concentration</th>
<th>Recommendation</th>
</tr>
</thead>
<tbody>
<tr>
<td>H₂ receptor antagonists</td>
<td>H₂ receptor antagonist</td>
<td>↓ ATV</td>
<td>H₂ Receptor antagonist should not exceed a 40 mg dose equivalent of famotidine twice daily and ATV should be administered with r simultaneously, with and/or at least 10 hours after the dose of the H₂-receptor antagonist. ATV/r is recommended. Proton-pump inhibitor dose should not exceed a 20 mg dose/day, equivalent of omeprazole and must be taken approximately 12 hours prior to ATV/RTV in antiretroviral-naive patients. Proton-pump inhibitors should not be used in treatment-experienced patients.</td>
</tr>
<tr>
<td>proton pump inhibitors</td>
<td>proton pump inhibitors</td>
<td>↓ ATV</td>
<td></td>
</tr>
<tr>
<td>Herbal products</td>
<td>St. John’s wort</td>
<td>expected ↓ ATV</td>
<td>Contraindicated.</td>
</tr>
<tr>
<td>HMG-CoA reductase inhibitors</td>
<td>lovastatin, simvastatin</td>
<td>↑ lovastatin, simvastatin</td>
<td>Contraindicated.</td>
</tr>
<tr>
<td></td>
<td>atorvastatin, rosuvastatin</td>
<td>↑ atorvastatin, rosuvastatin</td>
<td>Use the lowest possible dose of atorvastatin or rosuvastatin with careful monitoring or consider other HMG-CoA reductase inhibitor such as pravastatin or fluvastatin.</td>
</tr>
<tr>
<td>Immuno-suppressants</td>
<td>cyclosporine A, sirolimus</td>
<td>↑ immunosuppressants</td>
<td>TDM is recommended.</td>
</tr>
<tr>
<td>Neureptics</td>
<td>pimozide</td>
<td>↑ pimozide</td>
<td>Contraindicated.</td>
</tr>
<tr>
<td>Oral contraceptives</td>
<td>ethinyl estradiol,</td>
<td>↓↑ oral contraceptives</td>
<td>Due to possible alteration of oral contraceptives concentrations, alternative/additional contraceptive measures should be used when co-administered with ATV or ATV/r.</td>
</tr>
<tr>
<td></td>
<td>norethindrone</td>
<td></td>
<td></td>
</tr>
<tr>
<td>PDE5 inhibitors</td>
<td>sildenafil, tadalafil,</td>
<td>↑ sildenafil, tadalafil,</td>
<td>Do not exceed 25 mg of sildenafil in 48 h, 10 mg of tadalafil in 72 hours or 2.5 mg of vardenafil in 72 h.</td>
</tr>
<tr>
<td></td>
<td>vardenafil</td>
<td>vardenafil</td>
<td></td>
</tr>
</tbody>
</table>

Abbreviations: NRTI, nucleoside reverse transcriptase inhibitor; NNRTI, non-nucleoside reverse transcriptase inhibitor; PI, protease Inhibitor; INI, integrase Inhibitor; ATV, atazanavir; RTV, ritonavir; TDM, therapeutic drug monitoring; INR, international normalized ratio; ECG, electrocardiogram.

Figure 2 Antiretrovirals, bilirubin and UGT inhibition.
constitutive CYP3A4 expression in the liver and gut when compared to subjects homozygous for the 2677G allele.39

Several polymorphisms at the UGT1A1 gene have been described. The wild type allele (UGT1A1*1) contains 6 TA repeats in the promoter (TA6) whereas the most common variant allele (UGTA1A1*28) contains 7 repeats (TA7) and is the main one responsible for the mild form of Gilbert’s syndrome, an inherited unconjugated hyperbilirubinemia disorder.40–42 In a retrospective study conducted in 118 patients on a stable ATV/r regimen, the proportion of patients with grade 3–4 hyperbilirubinemia varied with distinct UGT1A1 genotypes: 80% for 7/7, 29% for 6/7 and 18% for 6/6. In the multivariate analysis, having at least one TA7 allele at UGT1A1 significantly predicted grade 3–4 hyperbilirubinemia.30

Rotger et al quantified the relative contribution of the UGT1A1*28 allele and different antiretroviral combinations on the risk of hyperbilirubinemia in 96 HIV-infected patients. ATV bilirubin levels increased by 0.87 mg/dL, while it was 0.46 mg/dL using indinavir. Patients homozygous for UGT1A1*28 had bilirubin levels increased by 0.3 mg/dL. As a result, 67% of individuals homozygous for UGT1A1*28 and receiving either ATV or indinavir had hyperbilirubinemia in the jaundice range (&gt;2.5 mg/dL), versus only 7% of those with the common allele not receiving any of these drugs. The implementation of UGT1A1*28 genotyping before initiation of antiretroviral therapy would lead to a theoretical 75% reduction in the number of patients experiencing jaundice.42

Other polymorphisms in the genes encoding for the UGT isoenzymes could also be associated with Gilbert’s syndrome.40,41 Lankisch et al examined the effects of different polymorphisms in UGT1A1 (UGT1A1*28), UGT1A3 (–66) and UGT1A7 (–57T &gt; G, W208R, N129K and R131K) genes on the risk of hyperbilirubinemia in patients treated with ATV. These isoenzymes are all inhibited by ATV. Interestingly, the homozygote haplotype for all 4 variants was present in up to 41% of patients with grade 3/4 and in 100% of patients with grade 4 hyperbilirubinemia. Therefore, the combination in homozygosis at *28, UGT1A7–57G, UGT1A3–66C and UGT1A7 was identified as highly predictive of severe hyperbilirubinemia under ATV therapy.41

Toxicity
ATV is generally well tolerated, as shown by the fact that only 5% to 10% of patients discontinued the drug due to adverse events in the main registrational studies with 48 weeks of follow-up. Indirect (unconjugated) bilirubin elevation is the most frequent side effect reported with ATV therapy and is due to inhibition of the UGT, as mentioned above. This unconjugated bilirubin elevation is frequently seen within the first months on ATV therapy and tends to slightly decline thereafter due to metabolic compensation mechanisms. This laboratory abnormality only achieves clinical relevance (grade 3–4) in up to one third of patients across studies. Jaundice is infrequent (&lt;10%) and excessive bilirubin elevations leading to ATV discontinuation are rare (around 1% of treated patients). The risk for hyperbilirubinemia seems to be associated to ATV plasma levels, and is more frequent when high doses of ATV are given (ie, 600 mg qd) or when ATV is boosted with r.9,25,29,30 Of note, hyperbilirubinemia is completely reversible after stopping ATV.

Grade 3–4 elevations in transaminases have been observed in 3% to 14% of patients receiving ATV. Liver enzyme elevations do not correlate with increased serum bilirubin and are more frequently seen in HIV subjects with underlying chronic hepatitis B or C. In the AI424-007 study,41 in which ATV was given in combination with didanosine and stavudine, grade 3–4 elevations in transaminases occurred in 20% of patients with chronic hepatitis B and in 40% of patients with chronic hepatitis C, but in &lt;10% of HIV-monoinfected individuals. In studies AI424-00844 and AI424-034,5 ALT levels &gt;5 times the upper limit of normality were seen respectively in 15%, 14% and 17% of seropositive patients for hepatitis B or C treated with ATV, efavirenz and nelfinavir. In study AI424-045,31 20 patients treated with ATV/r and 18 with LPV/r, all seropositive for hepatitis B and or C, experienced increases in ALT levels &gt;5 times the upper limit of normality in 25% (5/20) and 6% (1/18) of cases, respectively. Therefore, liver function tests should periodically be monitored in patients on ATV with underlying chronic liver disease.

Patients on ATV may occasionally complain of gastrointestinal disturbances, although symptoms are generally mild. In the AI424-007 study,45 grade 3–4 nausea/vomiting, abdominal pain, or diarrhea occurred in only 2% to 3% of patients. Of note, these side effects do not seem to rise when ATV is boosted with r. Indeed, in the AI424-045,31 the incidence of grade 2–4 gastrointestinal symptoms was 3% in the ATV/r arm but was much higher in the LPV/r arm (11%).

Some concern has been raised with respect to the potential cardiac toxicity of ATV, following sporadic reports in the literature. This adversity was not originally reported during the clinical development of the drug and has only rarely been noticed in the post-marketing period.45 ATV has been shown to prolong the PR interval in electrocardiograms performed on healthy volunteers as well as in HIV-infected patients.46
Abnormalities in atrioventricular conduction are generally asymptomatic, concentration-dependent and limited to first-degree atrioventricular block. Anecdotal reports of second-degree atrioventricular block and other conduction abnormalities have been published. A retrospective analysis of patients enrolled in the ATV expanded access program (AI424-900 study) has shown that QRS intervals increased by a median of 5 ms in 56 out of 75 (74.7%) antiretroviral-experienced patients (66.7% male), using either boosted or unboosted ATV. The PR and the QTc intervals did not change significantly. Interestingly, new asymptomatic bundle branch blocks were observed in 4 patients. According to pooled data from the manufacturer’s prescribing information, the incidence of QTc interval prolongation in a total of 1793 patients treated with ATV was comparable to that of patients receiving other PIs, with none of the patients showing a QTc interval >500 ms. Since pre-existing cardiac conduction abnormalities was an exclusion criterion in many early ATV studies, the prevalence of electrocardiogram abnormalities in unselected populations remains unknown. An additive effect of ATV and drugs that prolong the PR interval (eg, beta-blockers, verapamil, digoxin) and the QT interval cannot be excluded; consequently, when possible these combinations should be avoided. Altogether, these observations show that periodic electrocardiogram monitoring is worthwhile in patients treated with ATV, particularly when boosted with r.

Cases of nephrolithiasis in patients on ATV have been reported after marketing of ATV. This complication resembles the well known side effect of indinavir, which was able to produce kidney stones. In some instances, ATV crystals have been demonstrated in urine. However, this complication is very rare. In one retrospective study the prevalence of ATV-associated urolithiasis was 0.97%. Patients with low water intake, high urinary pH, and prior history of urinary stones are at higher risk for ATV-associated urine crystallization.

Unlike other PIs, ATV does not seem to negatively impact the lipid profile. ATV also seems to have less impact on insulin and overall on glucose metabolism than all other PIs. Following this observation a lack of involvement of the drug in morphological lipodystrophy abnormalities has been presumed, which is somewhat unique for PIs, although a longer follow-up is needed to confirm this singularity.

**Clinical experience**

**Atazanavir in treatment-naïve subjects**

ATV has been compared with other potent agents such as nelfinavir, efavirenz, saquinavir and LPV/RTV as part of triple therapy with two nucleoside reverse transcriptase inhibitors (NRTI) in treatment-naïve patients (Table 4). Two studies, AI424-00743 and AI424-008, compared different doses of unboosted ATV with nelfinavir along with didanosine + stavudine or lamivudine + stavudine, respectively. The vire-
logical response to both PIs was similar in terms of mean viral load reduction at 48 weeks and proportion of patients attaining undetectable viremia. Likewise, the increase in CD4+ T lymphocytes was similar for all groups in both studies. ATV and nelfinavir regimens were generally well tolerated, with the best efficacy/safety profile being found when ATV was given as 400 mg qd. Indeed, this was the selected dose for later studies. The metabolic indulge of ATV was one of the most striking findings of the AI424-007 and AI424-008 studies. Increases in total cholesterol at 48 weeks were 25% and 28% for nelfinavir in each trial, but only 5% and 7% for ATV (400 mg qd), respectively (p < 0.05). The same favorable behavior was seen for triglycerides (1.5% and 7% increase in each study) over nelfinavir (42% and 50% increase in each study).

The AI424-034 study compared unboosted ATV with efavirenz (one if not the most preferred third agent in drug-naïve patients) in combination with zidovudine + lamivudine as the nucleoside backbone. Despite the recruitment of many patients with high viral loads (42% with ≥5 log HIV-RNA copies/mL and low CD4 counts (median of 282 cells/µL), the performance was excellent and comparable after 48 weeks: 70% and 64% of patients with ATV and efavirenz, respectively, attained <400 HIV-RNA copies/mL. Some technical problems in the processing of specimens led to low rates of patients with values <50 copies/mL in both groups (32% with ATV and 37% with EFV). The regimens were comparable with respect to the magnitude and rate of CD4 + T cell gains. Interestingly, ATV treatment was associated with a more favorable lipid profile than was efavirenz treatment. In fact, the mean increase in total and low-density lipoprotein (LDL) cholesterol was significantly higher for efavirenz than for ATV (+21% and +18% versus +2% and +1%, respectively). The high-density lipoprotein (HDL) cholesterol fraction rose in both groups, but was more pronounced with efavirenz than with ATV (+24% versus +13%; p < 0.001). Finally, mean triglyceride levels decreased with ATV (−9%) and increased with efavirenz (+23%) (p < 0.001). No significant variations in glucose metabolism were noticed in either group, although patients on efavirenz tended to show slight increases in fasting insulin concentrations.

These initial registrational trials did not compare boosted ATV with other PIs or non-nucleoside reverse transcriptase inhibitors (NNRTI), and boosted ATV was initially disregarded for use in drug-naïve patients, at least in North America. These trials were followed by the AI424-089 study, a randomized, multicenter, 96-week study designed to compare the efficacy and safety of ATV/r to ATV, each given once daily in combination with once-daily lamivudine and extended-release stavudine in drug-naïve patients. In this comparison of boosted and unboosted ATV, the overall efficacy results were comparable. However, while not statistically significant, the rates of response were higher and emergence of PI resistance lower in subjects on ATV/r than in those on unboosted ATV. Based on these results, other studies using boosted ATV in treatment-naïve subjects were designed. The concern about a lower potency of unboosted ATV has been recorded in several antiretroviral treatment guidelines, in which ATV alone is only described as an alternative (but not preferred) PI in drug-naïve patients. Unboosted ATV may be chosen in highly selected circumstances as initial therapy for patients when a once-daily regimen without r is desired, in subjects not candidates for NNRTI-based regimens, and in patients with underlying cardiovascular risk factors in whom hyperlipidemia may be particularly undesirable. Of course, TDM would be desirable in these circumstances.
Atazanavir in HIV therapeutics

Atazanavir in treatment-experienced failing patients

It must be acknowledged that the approval of ATV was accelerated based on the results of another three clinical trials run in patients with virological failure under PI-containing regimens (Table 5). In this adverse setting, the AI424-009 study compared two rescue regimens based on ATV (400 and 600 mg) plus SQV (1200 mg) once a day versus SQV/r (400/400 mg) twice daily either in combination with two NRTI chosen based on phenotypic susceptibility data. The rationale for using ATV + SQV was based on the evidence of a booster effect of the former on SQV by inhibiting the cytochrome P450 3A4 isoenzyme in the liver. More than 85% of patients had prior PI experience and the median duration of prior PI therapy was approximately 2 years. The best virological response at 48 weeks was found in the ATV 400 mg qd group (41%) over the ATV 600 mg qd (29%) or SQV/r groups (35%). There were fewer adverse event discontinuations in the ATV/r groups (9% and 11%, respectively) than in the SQV/r arm (30%). Moreover, none of the ATV groups showed the adverse effects on lipids seen in the SQV/r arm.
In the AI424-043 study all patients had failed with at least one PI and were rescued with unboosted ATV versus LPV/r along with a nucleoside backbone selected according to resistance testing. LPV/RTV resulted in a significantly greater reduction in plasma HIV-RNA than unboosted ATV (−2.02 versus −1.59 log_{10} copies/mL, \( p < 0.001 \)) at week 48. Moreover, CD4 gains were higher with LPV/r than with ATV (169 versus 112 cells/μL). However, in the subset of patients with lack of NRTI resistance mutations at baseline, both regimens demonstrated comparable virologic suppression. The metabolic indulgence of ATV was especially manifest in this study. While LPV/r increased lipid levels and induced insulin resistance, these parameters remained stable in the ATV arm.

The third study (AI424-045) compared boosted ATV with LPV/r. The trial expected to find comparable efficacy and less toxicity with ATV/r. Patients who had failed two or more HAART regimens were randomized to ATV/r (300/100 mg qd), ATV/SQV (400/1200 mg qd) or LPV/r (400/100 mg bid), all along with TDF once daily and another NRTI. Approximately 40% of patients harbored ≥4 NRTI resistance mutations and more than one third harbored ≥4 PI resistance mutations. The virologic and immunologic results were worse for patients treated with ATV/SQV as compared with the other two groups (LPV/r and ATV/r), and therefore this arm was stopped at 24 weeks. In the intention-to-treat analysis at 96 weeks, similar virological efficacy was demonstrated for the remaining ATV/r and LPV/r arms, with 44% versus 46% of patients achieving <400 HIV-RNA copies/mL, respectively. CD4 gains at that time were also comparable (+160 and +142 cells/μL, respectively). Although response rates were similar when fewer than 4 PI-resistance associated mutations were present at baseline, LPV/r was superior to ATV/r in patients with more PI resistance mutations. This result suggested that ATV/r is of similar efficacy to LPV/r in antiretroviral-experienced patients with a limited extent of PI resistance, but it was inferior when a greater PI resistance was present. The tolerability of the three regimens was good and comparable, although only the ATV-containing groups benefited from less frequent gastrointestinal disturbances and significant reductions in total cholesterol, LDL cholesterol, and triglyceride concentrations. Moreover, in contrast with LPV/r, patients on ATV/r showed a significantly lower use of anti-diarrheal and lipid-lowering agents. The only drawback was grade 3–4 hyperbilirubinemia, which developed in 53% on ATV/r, a rate greater than expected and for which there is no clear explanation so far.

Despite the good performance of boosted ATV-based regimens in short- and long-term treatment of antiretroviral-experienced patients, its efficacy may be impaired in heavily pretreated patients. In terms of virological potency, many studies have concluded that as salvage therapy in antiretroviral-experienced patients, ATV/r could be a valuable option, unless a high number of PI resistance mutations are present. Moving a further step, the results of the French prospective NADIS cohort have concluded that ATV/r-containing regimes may be useful in heavily antiretroviral-experienced patients as long as failure has not occurred on LPV/r.

Studies comparing the antiretroviral efficacy, safety and effect of serum lipids of ATV/r versus LPV/r in HIV subjects who have experienced their first virologic failure while receiving a NNRTI-containing HAART regimen are currently ongoing (eg, AI424-103 trial).

Finally, boosted double PI regimens (r plus 2 additional PIs) gained interest a few years ago as a treatment option for multiple treatment-experienced patients with little or no options. The advent of new antiretrovirals, such as enfuvirtide, raltegravir, darunavir, maraviroc and etravirine, with the ability to achieve long-term viral load reductions in heavily antiretroviral-experienced patients with relatively few, manageable toxicities, and low drug interactions has clearly halted the research on double boosted PIs.

**Atazanavir in switch studies**

The long-term adverse effects and the difficulty of continuous almost perfect adherence to antiretroviral regimens have led to assessment of the impact of strategies aimed to simplify therapy. Obviously, a requirement is preservation of viral suppression and reduced risk of toxicities. As a simplification strategy, ATV-based regimens may offer several therapeutic advantages, including a convenient once-daily dosing schedule, low pill burden and lack of metabolic undesirable side effects compared to efavirenz or other PI-based regimens (Table 6).

In the AI424-044 study, patients completing ≥48 weeks in another trial (AI424-008) with a plasma HIV-RNA <10,000 copies/mL were randomized to continue on ATV (400 or 600 mg qd) or switch from nelfinavir to ATV (400 mg qd). After 24 weeks of ATV therapy, 83%, 85% and 87% of subjects in the ATV 400 mg, ATV 600 mg and switch groups, respectively, had plasma HIV-RNA <400 copies/mL. These figures were 76%, 76% and 63%, respectively, at week 48 in the prior AI424-008 study. Patients switched from nelfinavir to ATV showed significant mean declines in total cholesterol

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Atazanavir in HIV therapeutics

(−16%), fasting LDL cholesterol (−21%), and fasting triglycerides (−28%) (p < 0.0001) by week 12 of ATV therapy, approaching baseline values.

Further studies have confirmed the benign lipid profile of ATV. The AI424-067 study was designed as a switch trial with the aim of managing hyperlipidemia in patients with well-controlled HIV replication. Preliminary 12-week data were reported a few years ago from 246 participants on stable HAART with undetectable viral load and elevated LDL cholesterol. They were randomly assigned to either continue their current PI-based regimen or switch to unboosted ATV while remaining on the same NRTIs. After 12 weeks, patients in the ATV arm maintained virological suppression while experiencing a 35% decrease in triglycerides, a 15% reduction in LDL cholesterol, an 18% drop in total cholesterol, and 14% increase in HDL cholesterol. In addition, the study comprised an immediate switch group in which the PI was replaced by ATV 400 mg qd and a delayed switch group in which subjects continued to receive their prior PI-containing regimen for the first 24 weeks and then switched to ATV 400 mg qd. The results at 48 weeks confirmed that the switch from any PI (r-boosted or not) to ATV 400 mg qd resulted in clinically relevant reductions in fasting LDL cholesterol. However, rates of viral rebound were 15% and 10% in the immediate and delayed switch groups, respectively. It should be noted that most of these patients had been exposed to suboptimal dual NRTI therapy in the past, and in this situation boosted instead of unboosted ATV would have been preferred by most clinicians to ensure enough drug exposure.

The Switch to Another Protease Inhibitor Study (SWAN or AI424-097 study) was a 48-week, open-label, prospective trial involving HIV patients with virologic suppression who were receiving stable PI-based regimens (with or without r). Patients were randomized 2:1 to switch to ATV 400 mg qd or to continue on their prior PI. After 48 weeks, patients who switched to ATV showed significantly fewer total cholesterol, fasting triglyceride, and non-HDL cholesterol.

### Table 6 Main switch studies with atazanavir

<table>
<thead>
<tr>
<th>Study</th>
<th>Phase</th>
<th>n</th>
<th>Backbone</th>
<th>Previous regimen</th>
<th>Intervention</th>
<th>Follow-up (weeks)</th>
<th>Main conclusion</th>
</tr>
</thead>
<tbody>
<tr>
<td>AI424-044</td>
<td>III</td>
<td>346</td>
<td>d4T + 3TC</td>
<td>ATV 400 mg qd</td>
<td>Continue on ATV (400 or 600 mg) or switch from NFV to ATV 400 mg qd</td>
<td>24</td>
<td>Maintenance of virologic suppression and significant improvement in lipid profile in patients switched to ATV</td>
</tr>
<tr>
<td>AI424-067</td>
<td>III</td>
<td>246</td>
<td>2 NRTI</td>
<td>Pl with or without RTV</td>
<td>Immediate switch to ATV 400 mg qd or delayed switch after 24 weeks</td>
<td>48</td>
<td>Virologic rebounds &gt;50 copies/mL, 15% in immediate switch and 10% in delayed switch group. Improvement in lipid profile in hyperlipidemic patients</td>
</tr>
<tr>
<td>AI424-097</td>
<td>III</td>
<td>419</td>
<td>2 NRTI</td>
<td>Pl with or without RTV</td>
<td>Continue existing PI or switch to ATV (400 mg qd) or to ATV/RTV (300/100 mg qd) if receiving TDF</td>
<td>48</td>
<td>Maintenance of virologic suppression and improvement in lipid profile in patients switched to ATV</td>
</tr>
<tr>
<td>SLOAT</td>
<td>IV</td>
<td>189</td>
<td>2 NRTI</td>
<td>LPV/RTV</td>
<td>Continue on LPV/RTV or switch to ATV (400 mg qd) or to ATV/RTV (300/100 mg qd) if receiving TDF</td>
<td>24</td>
<td>Maintenance of virologic suppression and improvement in lipid profile in patients switched to ATV</td>
</tr>
<tr>
<td>ATAZIP</td>
<td>IV</td>
<td>265</td>
<td>2 NRTI</td>
<td>LPV/RTV</td>
<td>Continue on LPV/RTV or switch to ATV RTV (300/100 mg qd)</td>
<td>48</td>
<td>Maintenance of virologic suppression and improvement in lipid profile but no significant change in body composition in patients switched to ATV</td>
</tr>
<tr>
<td>ReAL</td>
<td>IV</td>
<td>200</td>
<td>2 NRTI</td>
<td>Pl/ritonavir (72% LPV/RTV)</td>
<td>Continue existing PI or switch to ATV/RTV (300/100 mg qd) in patients with lipoatrophy</td>
<td>48</td>
<td>Maintenance of virologic suppression and improvement in lipid profile but no significant change in body composition in patients switched to ATV</td>
</tr>
</tbody>
</table>

Abbreviations: NRTI, nucleoside reverse transcriptase inhibitor; PI, protease inhibitor; ITT, intent-to-treat; VL, viral load; 3TC, lamivudine; d4T, stavudine; LPV, lopinavir; NFV, nevirapine; RTV, ritonavir; ATV, atazanavir; bid, twice daily; qd, once daily.

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elevations than did patients in the comparator PI group (p < 0.001). For patients with prior exposure to boosted PI regimens, the 2 treatment groups had comparable rates of virologic rebound (8% for the ATV group versus 11% for the comparator PI group). In contrast, significantly lower rebound rates were recognized for ATV versus comparator PI groups in the subset of patients with prior exposure to unboosted PIs (5% versus 22%; p < 0.001). A limitation of the study was that nearly half of the study participants entered the trial taking an obsolete unboosted PI modality and almost everyone switched to ATV did not take r, which is not the current standard of care.

The AI424-100 study tried to compare the change from baseline in fasting non-HDL cholesterol between hyperlipidemic subjects who were switched to an ATV/RTV-regimen and those who continued on a LPV/r regimen. Unfortunately, the study was prematurely discontinued due to poor enrolment and no efficacy analyses were performed. This issue was finally addressed by two Spanish phase IV studies, the SLOAT and the ATAZIP trials.

In the Simplification Lopinavir to Atazanavir (SLOAT) study, patients receiving LPV/r-based regimens (with or without hyperlipidemia) and having undetectable plasma HIV-RNA for longer than 24 weeks were randomized to continue on the same therapy or switch to ATV (400 mg qd) or to ATV/r (300/100 mg qd) if receiving TDF. The rate of virological failures at 48 weeks did not differ between groups. A significant reduction (p < 0.001) was seen in median total cholesterol (−19 mg/dL) and triglycerides (−80 mg/dL) in the ATV switch group, whereas no significant changes occurred in the control LPV/r arm. Greater reductions in total cholesterol and triglycerides were seen in patients switched to ATV without r boosting.

The ATAZIP study evaluated 248 individuals on stable LPV/RTV-based therapy (plasma HIV-RNA <200 copies/mL for longer than 6 months) who were randomized to either continue on the capsule formulation of LPV/r twice daily or switch to ATV/r (300/100 mg) qd. All individuals with more than 4 PI-associated resistance mutations and/or who had failed more than 2 PI-containing regimens were excluded from the study. Although HDL cholesterol did not change in either group, significantly more favorable shifts in total cholesterol, LDL cholesterol and triglycerides were observed in the ATV/r arm in comparison to the LPV/r arm. Virologic failure was uncommon and occurred in only 5% of ATV/r recipients and 6% of LPV/r patients. Virologic failure was typically associated with a prior history of PI failure and/or presence of baseline major PI resistance mutations.

In addition to a better lipid profile, several studies have attempted to evaluate the effects on glucose tolerance following switch to ATV from PI-based regimens. The rationale was derived from prior information showing that ATV seems to have less effect on glucose metabolism than other PIs. In a study which included 21 highly pretreated HIV-infected subjects with metabolic alterations, the switch to an unboosted ATV-containing regimen revealed a significant improvement in glucose tolerance. Likewise, in 9 HIV-infected men with dyslipidemia under a stable PI-containing regimen, the switch to ATV/RTV improved significantly insulin resistance, measured by the gold-standard euglycemic clamp method.

ATV is less likely to induce lipodystrophy than other PIs. Based on data at week 48 from a substudy of the AI424-034 trial, it would appear that ATV has little effect on changes in fat redistribution. This substudy focused on subjects with CD4 counts >100 cells/μL, whose initial antiretroviral therapy consisted of ZDV + 3TC plus either EFV or ATV. There was no evidence of lipoatrophy at 48 weeks in any treatment arm; although there was a small increase in both abdominal subcutaneous and limb fat levels, which was not statistically significant. Moreover, there was an increase in the visceral fat by computed tomography with either drug combination. This study could be criticized in that the 48-week follow-up might be insufficient to detect subtle or delayed deleterious effects. However, in the AI424-007 trial, lipodystrophy continued to be infrequent and not significant compared to nelfinavir after 48 weeks on unboosted ATV.

It is still unclear whether a switch from other PIs to ATV could ameliorate or even improve lipodystrophic effects of prior antiretroviral regimens. In a case series, 3 patients experienced a regression of dorso-cervical and abdominal fat accumulation after switching the existing PI to ATV. However, switching from PIs to ATV/r did not significantly reduce visceral fat accumulation in the abdomen at 48 weeks in the ReAL study. This trial evaluated the impact on body composition of switching from any twice-daily r-boosted PI to a once-daily ATV/r regimen in patients with lipohypertrophy (fat accumulation). Participants had been taking a boosted PI for a median of 24 months (LPV/r in 72%). No significant differences in visceral fat were seen comparing the two groups after 1 year, suggesting no meaningful effect of treatment switch on the primary study outcome. However, patients who switched to ATV experienced a significant reduction in all fasting lipids, while antiviral efficacy was preserved. This study is still ongoing and results at week 96 will soon be available. Nevertheless, lipid improvements seen with ATV most likely will not translate into significant body

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fat changes within 2 years, since fat abnormalities generally take longer to recover. Long-term studies will be needed to assess the true effect of ATV on lipodystrophy.

Although ATV has proved to be a more metabolic-friendly drug that other antiretrovirals, and may help to manage metabolic side effects, it remains unclear whether switching to ATV may be a better approach than adding lipid-lowering medications to prior medications. However, any improvement in the lipid profile on ATV is likely to be beneficial in terms of reducing cardiovascular risk, and may decrease the required dose of lipid-lowering medications. A recent retrospective study has shown that a switch to ATV in HIV-infected antiretroviral-experienced patients caused significant reductions in plasma lipids and a modest but significant reduction in the normalized-for-age cardiovascular risk score, mainly in patients with higher baseline total cholesterol and triglycerides. Nevertheless, another study revealed that the switch from PIs to ATV in antiretroviral-experienced patients did not result in recognizable improvements in endothelial function despite benefits in the serum lipids.

**Atazanavir/ritonavir monotherapy**

In an attempt to simplify antiretroviral regimens as much as possible, recent reports have claimed that PI/r monotherapy could be an alternative option in some patients, especially as a maintenance strategy once virological control has been achieved. The proof-of-concept studies for this strategy were originally conducted with LPV/r,

following which 3 uncontrolled studies with ATV/RTV monotherapy have been performed, resulting in conflicting results.

ATV/r was tested in a prospective pilot study of 34 HIV adults with virological suppression for ≥48 weeks receiving their first PI regimen. All participants switched to ATV/r at entry and discontinued the NRTI backbone after 6 weeks. Three participants (9%) experienced virologic failure at weeks 12, 14 and 20 after simplification, with plasma HIV-RNA levels of 4730, 1285 and 28397 copies/mL, respectively. Intriguingly, resistance testing at failure did not identify any PI resistance mutation. Plasma ATV concentrations at failure were low or below detection in 2 out of 3 subjects. The authors concluded that ATV/r maintenance monotherapy could be a valuable option in a subset of HIV patients. However, predicting those who would fail was unclear.

In another pilot trial using ATV/r as maintenance therapy, premature discontinuation was recommended when 15 of the planned 30 patients had been recruited, because 5 cases of virological failure were documented. In failing patients, viral rebound occurred between weeks 12 and 16. Plasma ATV concentrations did not predict the outcome. Again, no PI resistance mutations were found in failing patients.

The investigators concluded that ATV/r as maintenance monotherapy in HIV-1 infection should not be considered as potent as conventional triple antiretroviral therapy.

The third study, named ATARITMO, tried to determine the feasibility of ATV/r maintenance monotherapy along with its effects on viral replication in compartments other than plasma, such as the CSF and seminal plasma. At week 24, 3/20 patients had detectable viremia (HIV-RNA > 100 copies/mL) in CSF and 2/15 in semen, despite viral suppression in plasma (<50 HIV-RNA copies/mL). Interestingly, samples with elevated HIV-RNA (>500 copies/mL) in CSF were all wild type in CSF.

Altogether, the general view is that ATV/r monotherapy, as with other PIs, must not be considered as an acceptable optional strategy when any modality of triple antiretroviral regimens can be afforded.

**Conclusion**

ATV has opened a new PI era with safer and more convenient compounds. While it retains the potency of the oldest robust PIs, it offers the advantage of simplicity and minimal toxicity both in treatment-naïve and in treatment-experienced patients. Once the possibility of controlling HIV infection has already been achieved by choosing combinations of the 25 drugs currently available within the HIV treatment armamentarium, the availability of this kind of friendly drug may enable an improvement in the quality of life, a variable of growing relevance as the life expectancy of HIV-infected persons is approaching that of uninfected individuals.

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