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SUMMARY OF PRODUCT CHARACTERISTICS

1. NAME OF THE MEDICINAL PRODUCT

Atazanavir Sandoz 150 mg, harde capsules Atazanavir Sandoz 200 mg, harde capsules Atazanavir Sandoz 300 mg, harde capsules

2. QUALITATIVE AND QUANTITATIVE COMPOSITION

100 mg hard capsules

Each hard capsule contains 100 mg of atazanavir (as sulfate).

Excipient with known effect

Each hard capsule contains 43.7 mg of lactose (as monohydrate).

150 mg hard capsules

Each hard capsule contains 150 mg of atazanavir (as sulfate).

Excipient with known effect

Each hard capsule contains 65.6 mg of lactose (as monohydrate).

200 mg hard capsules

Each hard capsule contains 200 mg of atazanavir (as sulfate).

Excipient with known effect

Each hard capsule contains 87.4 mg of lactose (as monohydrate).

300 mg hard capsules

Each hard capsule contains 300 mg of atazanavir (as sulfate).

Excipient with known effect

Each hard capsule contains 131.1 mg of lactose (as monohydrate).

For the full list of excipients, see section 6.1.

3. PHARMACEUTICAL FORM

Hard capsule.

100 mg hard capsules:

Opaque, blue and white capsule of size 2 printed with white ink, with "100 mg" on the cap.

150 mg hard capsules:

Opaque, blue and powder blue capsule of size 1 printed with white ink, with "150 mg" on the cap.

200 mg hard capsules:

Opaque, blue capsule of size 0 printed with white ink, with "200 mg" on the cap.

300 mg hard capsules:

Opaque red and blue capsule of size 00 printed with white ink, with "300 mg" on the cap.

4. CLINICAL PARTICULARS

4.1 Therapeutic indications

[Nationally completed name] capsules, co-administered with low dose ritonavir, are indicated for the treatment of HIV-1-infected adults and paediatric patients 6 years of age and older in combination with other antiretroviral medicinal products (see section 4.2).

Based on available virological and clinical data from adult patients, no benefit is expected in patients with strains resistant to multiple protease inhibitors (≥ 4 PI mutations).

The choice of [Nationally completed name] in treatment-experienced adult and paediatric patients should be based on individual viral resistance testing and the patient's treatment history (see sections 4.4 and 5.1).

4.2 Posology and method of administration

Therapy should be initiated by a physician experienced in the management of HIV infection.

<u>Posology</u>

Adults

The recommended dose of [Nationally completed name] capsules is 300 mg once daily taken with ritonavir 100 mg once daily and with food. Ritonavir is used as a booster of atazanavir pharmacokinetics (see sections 4.5 and 5.1). (See also section 4.4 Withdrawal of ritonavir only under restrictive conditions).

Paediatric patients (6 years to less than 18 years of age and weighing at least 15 kg)
The dose of atazanavir capsules for paediatric patients is based on body weight as shown in Table 1 and should not exceed the recommended adult dose. [Nationally completed name] capsules must be taken with ritonavir and have to be taken with food.

Table 1: Dose for paediatric patients (6 years to less than 18 years of age and weighing at least 15 kg) for [Nationally completed name] capsules with ritonavir

0 0		
Body Weight (kg)	[Nationally completed name] once daily dose	ritonavir once
		daily dose ^a
15 to less than 35	200 mg	100 mg
at least 35	300 mg	100 mg

^a Ritonavir capsules, tablets or oral solution.

Paediatric patients (at least 3 months of age and weighing at least 5 kg): Other formulations of atazanavir may be available for paediatric patients at least 3 months of age and weighing at least 5 kg (see relevant Summary of Product Characteristics for alternative forms). Switching to capsules from other formulations is encouraged as soon as patients are able to consistently swallow capsules.

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When transitioning between formulations, a change in dose may be needed. Consult the dosing table for the specific formulation (see relevant Summary of Product Characteristics).

Special populations

Renal impairment

No dosage adjustment is needed. [Nationally completed name] with ritonavir is not recommended in patients undergoing haemodialysis (see sections 4.4 and 5.2).

Hepatic impairment

Atazanavir with ritonavir has not been studied in patients with hepatic impairment. [Nationally completed name] with ritonavir should be used with caution in patients with mild hepatic impairment. [Nationally completed name] with ritonavir must not be used in patients with moderate to severe hepatic impairment (see sections 4.3, 4.4 and 5.2).

In case of withdrawal of ritonavir from the initial recommended ritonavir-boosted regimen (see section 4.4), unboosted [Nationally completed name] could be maintained in patients with mild hepatic impairment at a dose of 400 mg, and in patients with moderate hepatic impairment with a reduced dose of 300 mg once daily with food (see section 5.2). Unboosted [Nationally completed name] must not be used in patients with severe hepatic impairment.

Pregnancy and Postpartum

During the second and third trimesters of pregnancy:

[Nationally completed name] 300 mg with ritonavir 100 mg may not provide sufficient exposure to atazanavir, especially when the activity of atazanavir or the whole regimen may be compromised due to drug resistance. Since there are limited data available and due to interpatient variability during pregnancy, Therapeutic Drug Monitoring (TDM) may be considered to ensure adequate exposure.

The risk of a further decrease in atazanavir exposure is expected when atazanavir is given with medicinal products known to reduce its exposure (e.g., tenofovir disoproxil or H2-receptor antagonists).

- If tenofovir disoproxil or an H2-receptor antagonist is needed, a dose increase to [Nationally completed name] 400 mg with ritonavir 100 mg with TDM may be considered (see sections 4.6 and 5.2).
- It is not recommended to use [Nationally completed name] with ritonavir for pregnant patients who are receiving both tenofovir disoproxil and an H2-receptor antagonist.

(See section 4.4 Withdrawal of ritonavir only under restrictive conditions).

During postpartum:

Following a possible decrease in atazanavir exposure during the second and third trimester, atazanavir exposures might increase during the first two months after delivery (see section 5.2). Therefore, postpartum patients should be closely monitored for adverse reactions.

• During this time, postpartum patients should follow the same dose recommendation as for non-pregnant patients, including those for co-administration of medicinal products known to affect atazanavir exposure (see section 4.5).

Paediatric patients (less than 3 months of age)

[Nationally completed name] should not be used in children less than 3 months because of safety concerns especially taking into account the potential risk of kernicterus.

Method of administration:

For oral use. The capsules should be swallowed whole.

4.3 Contraindications

Hypersensitivity to the active substance or to any of the excipients listed in section 6.1.

Atazanavir is contraindicated in patients with severe hepatic insufficiency (see sections 4.2, 4.4 and 5.2). Atazanavir with ritonavir is contraindicated in patients with moderate hepatic insufficiency (see sections 4.2, 4.4 and 5.2).

Co-administration with simvastatin or lovastatin (see section 4.5).

Combination of rifampicin (see section 4.5).

Combination of the PDE5 inhibitor sildenafil when used for the treatment of pulmonary arterial hypertension (PAH) only (see section 4.5). For co-administration of sildenafil for the treatment of erectile dysfunction see sections 4.4 and 4.5.

Co-administration with medicinal products that are substrates of the CYP3A4 isoform of cytochrome P450 and have narrow therapeutic windows (e.g., quetiapine, lurasidone, alfuzosin, astemizole, terfenadine, cisapride, pimozide, quinidine, bepridil, triazolam, midazolam administered orally (for caution on parenterally administered midazolam, see section 4.5), lomitapide, and ergot alkaloids, particularly, ergotamine, dihydroergotamine, ergonovine, methylergonovine) (see section 4.5).

Co-administration with grazoprevir-containing products, including elbasvir/grazoprevir fixed-dose combination (see section 4.5).

Co-administration with glecaprevir/pibrentasvir fixed-dose combination (see section 4.5).

Co-administration with products containing St. John's wort (*Hypericum perforatum*) (see section 4.5).

Co-administration with apalutamide, encorafenib and ivosidenib (see section 4.5).

Co-administration with carbamazepine, phenobarbital, and phenytoin (see section 4.5).

4.4 Special warnings and precautions for use

Co-administration of atazanavir with ritonavir at doses greater than 100 mg once daily has not been clinically evaluated. The use of higher ritonavir doses may alter the safety profile of atazanavir (cardiac effects, hyperbilirubinaemia) and therefore is not recommended. Only when atazanavir with ritonavir is co-administered with efavirenz, a dose increase of ritonavir to 200 mg once daily could be considered. In this instance, close clinical monitoring is warranted (see Interaction with other Medicinal Products below).

Patients with coexisting conditions

Hepatic impairment: Atazanavir is primarily hepatically metabolised and increased plasma concentrations were observed in patients with hepatic impairment (see sections 4.2 and 4.3). The safety and efficacy of atazanavir has not been established in patients with significant underlying liver disorders. Patients with chronic hepatitis B or C and treated with combination antiretroviral therapy are at an increased risk for severe and potentially fatal hepatic adverse

reactions. In case of concomitant antiviral therapy for hepatitis B or C, please refer also to the relevant Summary of Product Characteristics for these medicinal products (see section 4.8).

Patients with pre-existing liver dysfunction, including chronic active hepatitis, have an increased frequency of liver function abnormalities during combination antiretroviral therapy and should be monitored according to standard practice. If there is evidence of worsening liver disease in such patients, interruption or discontinuation of treatment must be considered.

Renal impairment: No dosage adjustment is needed in patients with renal impairment. However, [Nationally completed name] is not recommended in patients undergoing haemodialysis (see sections 4.2 and 5.2).

QT prolongation: Dose-related asymptomatic prolongations in PR interval with atazanavir have been observed in clinical studies. Caution should be used with medicinal products known to induce PR prolongations. In patients with pre-existing conduction problems (second degree or higher atrioventricular or complex bundle-branch block), [Nationally completed name] should be used with caution and only if the benefits exceed the risk (see section 5.1). Particular caution should be used when prescribing [Nationally completed name] in association with medicinal products which have the potential to increase the QT interval and/or in patients with pre-existing risk factors (bradycardia, long congenital QT, electrolyte imbalances (see sections 4.8 and 5.3).

Haemophiliac patients: There have been reports of increased bleeding, including spontaneous skin haematomas and haemarthroses, in type A and B haemophiliac patients treated with protease inhibitors. In some patients additional factor VIII was given. In more than half of the reported cases, treatment with protease inhibitors was continued or reintroduced if treatment had been discontinued. A causal relationship has been suggested, although the mechanism of action has not been elucidated. Haemophiliac patients should therefore be made aware of the possibility of increased bleeding.

Weight and metabolic parameters

An increase in weight and in levels of blood lipids and glucose may occur during antiretroviral therapy. Such changes may in part be linked to the disease control and life style. For lipids, there is in some cases evidence for a treatment effect, while for weight gain there is no strong evidence relating this to any particular treatment. For monitoring of blood lipids and glucose reference is made to established HIV treatment guidelines. Lipid disorders should be managed as clinically appropriate.

In clinical studies, atazanavir (with or without ritonavir) has been shown to induce dyslipidaemia to a lesser extent than comparators.

<u>Hyperbilirubinaemia</u>

Reversible elevations in indirect (unconjugated) bilirubin related to inhibition of UDP-glucuronosyl transferase (UGT) have occurred in patients receiving atazanavir (see section 4.8). Hepatic transaminase elevations that occur with elevated bilirubin in patients receiving [Nationally completed name] should be evaluated for alternative aetiologies. Alternative antiretroviral therapy to [Nationally completed name] may be considered if jaundice or scleral icterus is unacceptable to a patient. Dose reduction of [Nationally completed name] is not recommended because it may result in a loss of therapeutic effect and development of resistance.

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Indinavir is also associated with indirect (unconjugated) hyperbilirubinaemia due to inhibition of UGT. Combinations of atazanavir and indinavir have not been studied and co-administration of these medicinal products is not recommended (see section 4.5).

Withdrawal of ritonavir only under restrictive conditions

The recommended standard treatment is [Nationally completed name] boosted with ritonavir, ensuring optimal pharmacokinetic parameters and level of virologic suppression.

The withdrawal of ritonavir from the boosted regimen of [Nationally completed name] is not recommended, but may be considered in adults patients at the dose of 400 mg once daily with food only under the following combined restrictive conditions:

- absence of prior virologic failure
- undetectable viral load during the last 6 months under current regimen
- viral strains not harbouring HIV resistance associated mutations (RAMs) to current regimen.

[Nationally completed name] given without ritonavir should not be considered in patients treated with a backbone regimen containing tenofovir disoproxil and with other concomitant medications that reduce atazanavir bioavailability (see section 4.5 In case of withdrawal of ritonavir from the recommended atazanavir boosted regimen) or in case of perceived challenging compliance.

[Nationally completed name] given without ritonavir should not be used in pregnant patients given that it could result in suboptimal exposure of particular concern for the mother infection and vertical transmission.

Cholelithiasis

Cholelithiasis has been reported in patients receiving atazanavir (see section 4.8). Some patients required hospitalization for additional management and some had complications. If signs or symptoms of cholelithiasis occur, temporary interruption or discontinuation of treatment may be considered.

Chronic kidney disease

Chronic kidney disease in HIV-infected patients treated with atazanavir, with or without ritonavir, has been reported during postmarketing surveillance. A large prospective observational study has shown an association between an increased incidence of chronic kidney disease and cumulative exposure to atazanavir/ritonavir-containing regimen in HIV-infected patients with an initially normal eGFR. This association was observed independently of exposure to tenofovir disoproxil. Regular monitoring of the renal function of patients should be maintained throughout the treatment duration (see section 4.8).

Nephrolithiasis

Nephrolithiasis has been reported in patients receiving atazanavir (see section 4.8). Some patients required hospitalization for additional management and some had complications. In some cases, nephrolithiasis has been associated with acute renal failure or renal insufficiency. If signs or symptoms of nephrolithiasis occur, temporary interruption or discontinuation of treatment may be considered.

Immune reactivation syndrome

In HIV-infected patients with severe immune deficiency at the time of institution of combination antiretroviral therapy (CART), an inflammatory reaction to asymptomatic or residual opportunistic pathogens may arise and cause serious clinical conditions, or aggravation of symptoms. Typically, such reactions have been observed within the first few weeks or months of initiation of CART. Relevant examples are cytomegalovirus retinitis,

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generalised and/or focal mycobacterial infections, and *Pneumocystis jirovecii* pneumonia. Any inflammatory symptoms should be evaluated and treatment instituted when necessary. Autoimmune disorders (such as Graves' disease and autoimmune hepatitis) have also been reported to occur in the setting of immune reactivation; however, the reported time to onset is more variable and these events can occurs many months after initiation of treatment.

Osteonecrosis

Although the aetiology is considered to be multifactorial (including corticosteroid use, alcohol consumption, severe immunosuppression, higher body mass index), cases of osteonecrosis have been reported particularly in patients with advanced HIV disease and/or long-term exposure to combination antiretroviral therapy (CART). Patients should be advised to seek medical advice if they experience joint aches and pain, joint stiffness or difficulty in movement.

Rash and associated syndromes

Rashes are usually mild-to-moderate maculopapular skin eruptions that occur within the first 3 weeks of starting therapy with atazanavir.

Stevens-Johnson syndrome (SJS), erythema multiforme, toxic skin eruptions and drug rash with eosinophilia and systemic symptoms (DRESS) syndrome have been reported in patients receiving atazanavir. Patients should be advised of the signs and symptoms and monitored closely for skin reactions. [Nationally completed name] should be discontinued if severe rash develops.

The best results in managing these events come from early diagnosis and immediate interruption of any suspect medicines. If the patient has developed SJS or DRESS associated with the use of atazanavir, [Nationally completed name] may not be restarted.

Interactions with other medicinal products

The combination of atazanavir with atorvastatin is not recommended (see section 4.5).

Co-administration of atazanavir with nevirapine or efavirenz is not recommended (see section 4.5). If the co-administration of atazanavir with an NNRTI is required, an increase in the dose of both [Nationally completed name] and ritonavir to 400 mg and 200 mg, respectively, in combination with efavirenz could be considered with close clinical monitoring.

Atazanavir is metabolised principally by CYP3A4. Co-administration of [Nationally completed name] and medicinal products that induce CYP3A4 is not recommended (see sections 4.3 and 4.5).

PDE5 inhibitors used for the treatment of erectile dysfunction: particular caution should be used when prescribing PDE5 inhibitors (sildenafil, tadalafil, or vardenafil) for the treatment of erectile dysfunction in patients receiving atazanavir. Co-administration of [Nationally completed name] with these medicinal products is expected to substantially increase their concentrations and may result in PDE5-associated adverse reactions such as hypotension, visual changes and priapism (see section 4.5).

Co-administration of voriconazole and atazanavir with ritonavir is not recommended, unless an assessment of the benefit/risk justifies the use of voriconazole.

In the majority of patients, a reduction in both voriconazole and atazanavir exposures are expected. In a small number of patients without a functional CYP2C19 allele, significantly increased voriconazole exposures are expected (see section 4.5).

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Concomitant use of atazanavir/ritonavir and fluticasone or other glucocorticoids that are metabolised by CYP3A4 is not recommended unless the potential benefit of treatment outweighs the risk of systemic corticosteroid effects, including Cushing's syndrome and adrenal suppression (see section 4.5).

Concomitant use of salmeterol and atazanavir may result in increased cardiovascular adverse events associated with salmeterol. Co-administration of salmeterol and [Nationally completed name] is not recommended (see section 4.5).

The absorption of atazanavir may be reduced in situations where gastric pH is increased irrespective of cause.

Co-administration of atazanavir with proton pump inhibitors is not recommended (see section 4.5). If the combination of atazanavir with a proton pump inhibitor is judged unavoidable, close clinical monitoring is recommended in combination with an increase in the dose of atazanavir to 400 mg with 100 mg of ritonavir; doses of proton pump inhibitors comparable to omeprazole 20 mg should not be exceeded.

Co-administration of atazanavir with other hormonal contraceptives or oral contraceptives containing progestogens other than norgestimate or norethindrone has not been studied, and therefore should be avoided (see section 4.5).

Paediatric population

Safety

Asymptomatic PR interval prolongation was more frequent in paediatric patients than adults. Asymptomatic first- and second-degree AV block was reported in paediatric patients (see section 4.8). Caution should be used with medicinal products known to induce PR prolongations. In paediatric patients with pre-existing conduction problems (second degree or higher atrioventricular or complex bundle-branch block), [Nationally completed name] should be used with caution and only if the benefits exceed the risk. Cardiac monitoring is recommended based on the presence of clinical findings (e.g., bradycardia).

Efficacy

Atazanavir/ritonavir is not effective in viral strains harbouring multiple mutations of resistance.

[Nationally completed name] contains sodium and lactose

This medicinal product contains less than 1 mmol sodium (23 mg) per hard capsule, that is to say essentially 'sodium-free'.

Patients with rare hereditary problems of galactose intolerance, total lactase deficiency or glucose-galactose malabsorption should not take this medicinal product.

4.5 Interaction with other medicinal products and other forms of interaction

When atazanavir and ritonavir are co-administered, the metabolic drug interaction profile for ritonavir may predominate because ritonavir is a more potent CYP3A4 inhibitor than atazanavir. The Summary of Product Characteristics for ritonavir must be consulted before initiation of therapy with [Nationally completed name] and ritonavir.

Atazanavir is metabolised in the liver through CYP3A4. It inhibits CYP3A4. Therefore, atazanavir is contraindicated with medicinal products that are substrates of CYP3A4 and have a narrow therapeutic index: quetiapine, lurasidone, alfuzosin, astemizole, terfenadine, cisapride,

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pimozide, quinidine, bepridil, triazolam, orally administered midazolam, lomitapide, and ergot alkaloids, particularly ergotamine and dihydroergotamine (see section 4.3). Co-administration of atazanavir with grazoprevir-containing products, including elbasvir/grazoprevir fixed-dose combination is contraindicated because of the increase in grazoprevir and elbasvir plasma concentrations and potential for the increase in risk of ALT elevations associated with increased grazoprevir concentrations (see section 4.3). Co-administration of atazanavir with glecaprevir/pibrentasvir fixed-dose combination is contraindicated because of the potential increase in the risk of ALT elevations due to a significant increase in glecapreir and pibrentasvir plasma concentrations (see section 4.3).

Other interactions

Interactions between atazanavir and other medicinal products are listed in the table below (increase is indicated as " \uparrow ", decrease as " \downarrow ", no change as " \leftrightarrow "). If available, 90% confidence intervals (CI) are shown in parentheses. The studies presented in Table 2 were conducted in healthy subjects unless otherwise noted. Of importance, many studies were conducted with unboosted atazanavir, which is not the recommended regimen of atazanavir (see section 4.4).

If withdrawal of ritonavir is medically warranted under restrictive conditions (see section 4.4), special attention should be given to atazanavir interactions that may differ in the absence of ritonavir (see information below Table 2).

Table 2: Interactions between atazanavir and other medicinal products

Medicinal products by	Interaction	Recommendations
therapeutic area		concerning co-
		administration
ANTI-HCV AGENTS		
Grazoprevir 200 mg once daily (atazanavir 300 mg / ritonavir 100 mg once daily)	Atazanavir AUC: ↑43% (↑30% ↑57%) Atazanavir C _{max} : ↑12% (↑1% ↑24%) Atazanavir C _{min} : ↑23% (↑13% ↑134%) Grazoprevir AUC: ↑958% (↑678% ↑1339%) Grazoprevir C _{max} : ↑524% (↑342% ↑781%) Grazoprevir C _{min} : ↑1064% (↑696% ↑1602%)	Co-administration of atazanavir and elbasvir/grazoprevir is contraindicated because of a significant increase in grazoprevir plasma concentrations and an associated potential increase in the risk of ALT elevations (see section 4.3).
Elbasvir 50 mg once daily (atazanavir 300 mg / ritonavir 100 mg once daily)	Grazoprevir concentrations were greatly increased when coadministered with atazanavir/ritonavir. Atazanavir AUC: ↑7% (↓2% ↑17%) Atazanavir C _{max} : ↑2% (↓4% ↑8%) Atazanavir C _{min} : ↑15% (↑2% ↑29%) Elbasvir AUC: ↑376% (↑307% ↑456%)	

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1.3.1.1	Samenvalling	van de	Productkenmerken

Medicinal products by	Interaction	Recommendations
therapeutic area		concerning co-
	Elbasvir C _{max} : †315% (†246%	administration
	1397%)	
	Elbasvir C _{min} : ↑545% (↑451%	
	↑654%)	
	Elbasvir concentrations were	
	increased when co-administered	
Sofoshuvin 400 mg/	with atazanavir/ritonavir. Sofosbuvir AUC: \\$\dagger40\% (\\$\dagger25\%)	Co-administration of
Sofosbuvir 400 mg / velpatasvir 100 mg	\$500Sbuvii ACC. 40% (25% 157%)	atazanavir with
/voxilaprevir 100 mg single	Sofosbuvir C _{max} : †29% (†9%	voxilaprevir containing
dose*	↑52%)	products is expected to
(atazanavir 300 mg / ritonavir		increase the
100 mg once daily)	Velpatasvir AUC: ↑93% (↑58%	concentration of
	↑136%)	voxilaprevir. Co-
	Velpatasvir C_{max} : $\uparrow 29\%$ ($\uparrow 7\%$	administration of atazanavir with
	† 56%)	voxilaprevir-containing
	Voxilaprevir AUC: ↑331%	regimens is not
	(†276% †393%)	recommended.
	Voxilaprevir C _{max} : †342% (†265%	
	↑435%)	
	*Lack of pharmacokinetics	
	interaction bounds 70-143%	
	Effect on atazanavir and ritonavir	
	exposure has not been studied.	
	Expected:	
	← Ritonavir	
	The mechanism of int	
	The mechanism of interaction between atazanavir/ritonavir and	
	sofosbuvir/velpatasvir/voxilaprevir	
	is inhibition of OATP1B, P-gp,	
	and CYP3A.	
Glecaprevir 300 mg/	Glecaprevir AUC: ↑553% (↑424%	Co-administration of
pibrentasvir 120 mg once	↑714%)	atazanavir with
daily	Glecaprevir C_{max} : $\uparrow 306\%$ ($\uparrow 215\%$	glecaprevir/pibrentasvir
(atazanavir 300 mg / ritonavir 100 mg once daily*)	↑423%) Glecaprevir C _{min} : ↑1330% (↑885%	is contraindicated because of the potential
100 ling office daily)	1970%)	increase in the risk of
		ALT elevations due to
	Pibrentasvir AUC: ↑64% (↑48%	a significant increase in
	↑82%)	glecaprevir and
	Pibrentasvir C _{max} : ↑29% (↑15%	pibrentasvir plasma
	1 †45%)	concentrations (see
	Pibrentasvir C_{min} : $\uparrow 129\%$ ($\uparrow 95\%$	section 4.3).
	<u>↑168%)</u>	

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Medicinal products by therapeutic area	Interaction	Recommendations concerning co- administration
	*Effect of atazanavir and ritonavir on the first dose of glecaprevir and pibrentasvir is reported.	
ANTIPLATELETS	I m	
Ticagrelor	The mechanism of the interaction is CYP3A4 inhibition by atazanavir and/or ritonavir.	Co-administration of atazanavir with ticagrelor is not recommended due to potential increase in the antiplatelet activity of ticagrelor.
Clopidogrel	The mechanism of the interaction is CYP3A4 inhibition by atazanavir and or/ritonavir.	Co-administration with clopidogrel is not recommended due to potential reduction of the antiplatelet activity of clopidogrel.
Prasugrel	The mechanism of the interaction is CYP3A4 inhibition by atazanavir and or/ritonavir.	No dose adjustment is needed when prasugrel is co-administered with atazanavir (with or without ritonavir).
ANTI-RETROVIRALS		
inhibitors has not been studie	o-administration of atazanavir/ritonad but would be expected to increase exadministration is not recommended.	
Ritonavir 100 mg once	Atazanavir AUC: †250% (†144%	Ritonavir 100 mg once
daily (atazanavir 300 mg once daily) Studies conducted in HIV-	↑403%)* Atazanavir C _{max} : ↑120% (↑56% ↑211%)* Atazanavir C _{min} : ↑713% (↑359% ↑1339%)*	daily is used as a booster of atazanavir pharmacokinetics.
infected patients.	*In a combined analysis, atazanavir 300 mg and ritonavir 100 mg (n=33) was compared to atazanavir 400 mg without ritonavir (n=28). The mechanism of interaction between atazanavir and ritonavir is	

Nucleoside/nucleotide reverse transcriptase inhibitors (NRTIs)

Lamivudine 150 mg twice No significant effect on lamivudine Based on these data and

inhibition of UGT.

CYP3A4 inhibition.

Indinavir is associated with indirect unconjugated

hyperbilirubinaemia due to

Co-administration of

atazanavir and

indinavir is not

recommended (see section 4.4).

<u>Indinavir</u>

1.5.1.1 Gamenvatung van de i Toddotkenmerken			
Medicinal products by	Interaction	Recommendations	
therapeutic area		concerning co-	
		administration	
daily + zidovudine 300 mg	and zidovudine concentrations was	because ritonavir is not	
twice daily	observed.	expected to have a	
(atazanavir 400 mg once		significant impact on	
daily)		the pharmacokinetics	
		of NRTIs, the co-	
		administration of these	
		medicinal products and	
		atazanavir is not	
		expected to	
		significantly alter the	
		exposure of the co-	
		administered medicinal	
		products.	
Abacavir	The co-administration of abacavir		
	and atazanavir is not expected to		
	significantly alter the exposure of		
	abacavir.		
Didanosine (buffered	Atazanavir, simultaneous	Didanosine should be	
tablets) 200 mg/stavudine	administration with ddI+d4T	taken at the fasted state	
40 mg, both single dose	(fasted)	2 hours after atazanavir	
(atazanavir 400 mg single	Atazanavir AUC: \$100 (\$100)	taken with food. The	
dose)	↓79%) 	co-administration of	
	Atazanavir C_{max} : $\downarrow 89\%$ ($\downarrow 94\%$	stavudine with	
	\\ \dagger{1}{2}\\ \dagger{1}{2}\\ \dagger{1}{2}\\ \dagger{1}\\ \dagge	atazanavir is not	
	Atazanavir C_{min} : $\downarrow 84\%$ ($\downarrow 90\%$	expected to	
	↓73%)	significantly alter the	
	Atazanavir, dosed 1 hr after	exposure of stavudine.	
	ddI+d4T (fasted)		
	Atazanavir AUC: ↔3% (↓36%		
	↑67%)		
	Atazanavir C_{max} : $\uparrow 12\%$ ($\downarrow 33\%$		
	118%)		
	Atazanavir C_{min} : $\leftrightarrow 3\%$ ($\downarrow 39\%$		
	173%)		
	Atazanavir concentrations were		
	greatly decreased when co-		
	administered with didanosine		
	(buffered tablets) and stavudine.		
	The mechanism of interaction is a		
	reduced solubility of atazanavir		
	with increasing pH related to the		
	presence of anti-acid agent in		
	didanosine buffered tablets.		
	No significant effect on didanosine		
	and stavudine concentrations was		
D.I.	observed.		
Didanosine (enteric coated	Didanosine (with food)		
capsules) 400 mg single dose	Didanosine AUC: ↓34% (↓41%		

Medicinal products by Interaction Recommendation		
therapeutic area	The action	concerning co-
(atazanavir 300 mg once daily with ritonavir 100 mg once daily) Tenofovir disoproxil fumarate 300 mg once daily (atazanavir 300 mg once	↓27%) Didanosine C _{max} : ↓38% (↓48% ↓26%) Didanosine C _{min} : ↑25% (↓8% ↑69%) No significant effect on atazanavir concentrations was observed when administered with enteric-coated didanosine, but administration with food decreased didanosine concentrations. Atazanavir AUC: ↓22% (↓35% ↓6%)* Atazanavir C _{max} : ↓16% (↓30%	When co-administered with tenofovir disoproxil fumarate, it
daily with ritonavir 100 mg once daily) 300 mg tenofovir disoproxil fumarate is equivalent to 245 mg tenofovir disoproxil. Studies conducted in HIV-infected patients	Atazanavir C _{min} : ↓23% (↓43% ↑2%)* *In a combined analysis from several clinical studies, atazanavir/ritonavir 300/100 mg co-administered with tenofovir disoproxil fumarate 300 mg (n=39) was compared to atazanavir/ritonavir 300/100 mg (n=33). The efficacy of atazanavir/ritonavir in combination with tenofovir disoproxil fumarate in treatment-experienced patients has been demonstrated in clinical study 045 and in treatment naive patients in clinical study 138 (see sections 4.8 and 5.1). The mechanism of interaction between atazanavir and tenofovir disoproxil fumarate is unknown.	is recommended that atazanavir 300 mg be given with ritonavir 100 mg and tenofovir disoproxil fumarate 300 mg (all as a single dose with food).
Tenofovir disoproxil fumarate 300 mg once daily (atazanavir 300 mg once daily with ritonavir 100 mg once daily) 300 mg tenofovir disoproxil fumarate is equivalent to 245 mg tenofovir disoproxil.	Tenofovir disoproxil fumarate AUC: ↑37% (↑30% ↑45%) Tenofovir disoproxil fumarate C _{max} : ↑34% (↑20% ↑51%) Tenofovir disoproxil fumarate C _{min} : ↑29% (↑21% ↑36%)	Patients should be closely monitored for tenofovir disoproxil fumarate-associated adverse reactions, including renal disorders.

Medicinal products by therapeutic area	Interaction	Recommendations concerning co-
therapeutic area		administration
Non-nucleoside reverse transci	riptase inhibitors (NNRTIs)	
Efavirenz 600 mg once daily (atazanavir 400 mg once daily with ritonavir 100 mg once daily)	Atazanavir (pm): all administered with food Atazanavir AUC: $\leftrightarrow 0\%$ ($\downarrow 9\%$ $\uparrow 10\%$)* Atazanavir C_{max} : $\uparrow 17\%$ ($\uparrow 8\%$ $\uparrow 27\%$)* Atazanavir C_{min} : $\downarrow 42\%$ ($\downarrow 51\%$ $\downarrow 31\%$)*	Co-administration of efavirenz and atazanavir is not recommended (see section 4.4).
Efavirenz 600 mg once daily (atazanavir 400 mg once daily with ritonavir 200 mg once daily)	Atazanavir (pm): all administered with food Atazanavir AUC: ←6% (↓10% ↑26%)*/** Atazanavir C _{max} : ←9% (↓5% ↑26%)*/** Atazanavir C _{min} : ←12% (↓16% ↑49%)*/** *When compared to atazanavir 300 mg/ritonavir 100 mg once daily in the evening without efavirenz. This decrease in atazanavir C _{min} might negatively impact the efficacy of atazanavir. The mechanism of efavirenz/atazanavir interaction is CYP3A4 induction. **Based on historical comparison.	
Nevirapine 200 mg twice daily (atazanavir 400 mg once daily with ritonavir 100 mg once daily) Study conducted in HIV-infected patients	Nevirapine AUC: $\uparrow 26\%$ ($\uparrow 17\%$) $\uparrow 36\%$) Nevirapine C_{max} : $\uparrow 21\%$ ($\uparrow 11\%$) $\uparrow 32\%$) Nevirapine C_{min} : $\uparrow 35\%$ ($\uparrow 25\%$) $\uparrow 47\%$) Atazanavir AUC: $\downarrow 19\%$ ($\downarrow 35\%$) $\uparrow 22\%$)* Atazanavir C_{max} : $\leftrightarrow 2\%$ ($\downarrow 15\%$) $\uparrow 24\%$)* Atazanavir C_{min} : $\downarrow 59\%$ ($\downarrow 73\%$) $\downarrow 40\%$)* *When compared to atazanavir 300 mg and ritonavir 100 mg without nevirapine. This decrease in atazanavir C_{min} might negatively impact the efficacy of atazanavir. The mechanism of nevirapine/atazanavir interaction is CYP3A4 induction.	Co-administration of nevirapine and atazanavir is not recommended (see section 4.4).
Integrase Inhibitors		
Raltegravir 400 mg twice	Raltegravir AUC: ↑41%	No dose adjustment

Medicinal products by therapeutic area	Interaction	Recommendations
therapeutic area		concerning co- administration
daily	Raltegravir C _{max} : ↑24%	required for raltegravir.
(atazanavir/ritonavir)	Raltegravir C_{12hr} : $\uparrow 77\%$	required for fameBrushis
	5 12.11 1	
	The mechanism is UGT1A1	
	inhibition.	
ANTIBIOTICS		
Clarithromycin 500 mg	Clarithromycin AUC: †94%	No recommendation
twice daily (atazanavir 400 mg once	$(\uparrow 75\% \uparrow 116\%)$ Clarithromycin C _{max} : $\uparrow 50\% (\uparrow 32\%)$	regarding dose reduction can be made; therefore,
daily)	↑71%)	caution should be
(daily)	Clarithromycin C _{min} : ↑160%	exercised if atazanavir is
	(†135% †188%)	co-administered with
	(1-00.1)	clarithromycin.
	14-OH clarithromycin	·
	14-OH clarithromycin AUC: ↓70%	
	(↓74% ↓66%)	
	14-OH clarithromycin C _{max} : ↓72%	
	(\psi 76% \psi 67%)	
	14-OH clarithromycin C _{min} : ↓62%	
	(↓66% ↓58%)	
	Atazanavir AUC: †28% (†16%	
	†43%)	
	Atazanavir C_{max} : \leftrightarrow 6% (\downarrow 7%	
	†20%)	
	Atazanavir C_{min} : $\uparrow 91\%$ ($\uparrow 66\%$	
	↑121%)	
	A dose reduction of clarithromycin	
	may result in subtherapeutic	
	concentrations of 14-OH	
	clarithromycin. The mechanism of	
	the clarithromycin/atazanavir	
	interaction is CYP3A4 inhibition.	
ANTIFUNGALS	NT	V.4
Ketoconazole 200 mg once	No significant effect on atazanavir concentrations was observed.	Ketoconazole and itraconazole should be
daily (atazanavir 400 mg once daily)	concentrations was observed.	used cautiously with
Itraconazole	Itraconazole, like ketoconazole, is a	atazanavir/ritonavir, high
	potent inhibitor as well as a	doses of ketoconazole
	substrate of CYP3A4.	and itraconazole (>200
	Based on data obtained with other	mg/day) are not
	boosted PIs and ketoconazole,	recommended.
	where ketoconazole AUC showed a	
	3-fold increase, atazanavir/ritonavir	
	is expected to increase ketoconazole	
Varianazala 200 ma turias	or itraconazole concentrations.	Co administration of
Voriconazole 200 mg twice daily	Voriconazole AUC: ↓33% (↓42% ↓22%)	Co-administration of voriconazole and
(atazanavir 300 mg/ritonavir	Voriconazole C_{max} : $\downarrow 10\%$ ($\downarrow 22\%$	voi iconazore and
(atazanavn 300 mg/monavn	VOLICOHAZOIC C _{max} . ↓10/0 (↓22/0	

Medicinal products by	Interaction	Recommendations
therapeutic area		concerning co-
		administration
100 mg once daily)	↓4%)	atazanavir with ritonavir
	Voriconazole C _{min} : ↓39% (↓49%	is not recommended
Subjects with at least one	↓28%)	unless an assessment of
functional CYP2C19 allele.		the benefit/risk to the
	1.175	patient justifies the use
	Atazanavir AUC: ↓12% (↓18%	of voriconazole (see
	\$\frac{15\%}{120\chi}\$	section 4.4).
	Atazanavir C_{max} : $\downarrow 13\%$ ($\downarrow 20\%$	
	14%)	At the time voriconazole
	Atazanavir C_{min} : $\downarrow 20 \% (\downarrow 28 \%)$	treatment is required, a
	↓10%)	patient's CYP2C19
		genotype should be performed if feasible.
	D:tamaria AIIC. 120/ (170/	performed if leasible.
	Ritonavir AUC: ↓12% (↓17%	Therefore, if the
	$\begin{array}{c} \downarrow 7\%) \\ \text{Ritonavir C}_{\text{max}} \colon \downarrow 9\% \ (\downarrow 17\% \leftrightarrow 0\%) \end{array}$	combination is
	Ritonavir C_{max} : $\downarrow 9\% (\downarrow 1/\% \leftrightarrow 0\%)$ Ritonavir C_{min} : $\downarrow 25\% (\downarrow 35\%)$	unavoidable, the
	$\downarrow 14\%)$	following
	\ \frac{1470}{}	recommendations are
	In the majority of patients with at	made according to the
	least one functional CYP2C19	CYP2C19 status:
	allele, a reduction in both	
	voriconazole and atazanavir	- in patients with at least
	exposures are expected.	one functional CYP2C19
Voriconazole 50 mg twice	Voriconazole AUC: \^561\%	allele, close clinical
daily	(†451% †699%)	monitoring for a loss of
(atazanavir 300 mg/ritonavir	Voriconazole C _{max} : \dagger 438%	both voriconazole
100 mg once daily)	(†355% †539%)	(clinical signs) and
	Voriconazole C_{min} : $\uparrow 765\%$	atazanavir (virologic
Subjects without a functional	(†571% †1,020%)	response) efficacy is
CYP2C19 allele.		recommended.
	Atazanavir AUC: ↓20% (↓35%	
	↓3%)	- in patients without a
	Atazanavir C_{max} : $\downarrow 19\%$ ($\downarrow 34\%$	functional CYP2C19
	↔0.2%)	allele, close clinical and
	Atazanavir C_{min} : $\downarrow 31\%$ ($\downarrow 46\%$	laboratory monitoring of voriconazole-associated
	↓13%)	adverse events is
	Ritonavir AUC: \$\frac{11\%}{20\%}\$	recommended.
	$\downarrow 1\%$	1000iiiiiioiidod.
	Ritonavir C_{max} : $\downarrow 11\%$ ($\downarrow 24\% \uparrow 4\%$)	If genotyping is not
	Ritonavir C_{max} : $\downarrow 17\%$ ($\downarrow 24\%$ $\uparrow 47\%$) Ritonavir C_{min} : $\downarrow 19\%$ ($\downarrow 35\%$ $\uparrow 1\%$)	feasible, full monitoring
	Tatomarii Oniii. \$1770 (\$5570 170)	of safety and efficacy
	In a small number of patients	should be performed.
	without a functional CYP2C19	•
	allele, significantly increased	
	voriconazole exposures are	
	expected.	
Fluconazole 200 mg once	Atazanavir and fluconazole	No dosage adjustments
daily	concentrations were not	are needed for

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Medicinal products by	Interaction	Recommendations
therapeutic area		concerning co-
		administration
(atazanavir 300 mg and	significantly modified when	fluconazole and
ritonavir 100 mg once daily)	atazanavir/ritonavir was co-	atazanavir.
	administered with fluconazole.	
ANTIMYCOBACTERIAL	T	
Rifabutin 150 mg twice	Rifabutin AUC: ↑48% (↑19%	When given with
weekly	↑84%)**	atazanavir, the
(atazanavir 300 mg and	Rifabutin C_{max} : $\uparrow 149\%$ ($\uparrow 103\%$	recommended dose of
ritonavir 100 mg once daily)	1 \(\frac{1}{206} \) \(\) \(\text{**} \) \(\text{Pifobution } C \) \(\text{*} \(\text{*} \) \(\text{*} \(\text{*} \) \(\text{*} \)	rifabutin is 150 mg 3
	Rifabutin C _{min} : †40% (†5% †87%)**	times per week on set days (for example
	[6770]	Monday-Wednesday-
	25-O-desacetyl-rifabutin AUC:	Friday). Increased
	†990% (†714% †1361%)**	monitoring for rifabutin-
	25-O-desacetyl-rifabutin C _{max} :	associated adverse
	↑677% (↑513% ↑883%)**	reactions including
	25-O-desacetyl-rifabutin C _{min} :	neutropenia and uveitis
	1045% (†715% †1510%)**	is warranted due to an
		expected increase in
	**When compared to rifabutin 150	exposure to rifabutin.
	mg once daily alone. Total rifabutin	Further dosage reduction
	and 25-O-desacetyl-rifabutin	of rifabutin to 150 mg
	AUC: †119% (†78% †169%).	twice weekly on set days is recommended for
	In previous studies, the	patients in whom the
	pharmacokinetics of atazanavir was	150 mg dose 3 times per
	not altered by rifabutin.	week is not tolerated. It
		should be kept in mind
		that the twice weekly
		dosage of 150 mg may
		not provide an optimal
		exposure to rifabutin
		thus leading to a risk of
		rifamycin resistance and
		a treatment failure. No
		dose adjustment is
Diforminin	Diferenciais is a stress CVD2 A 4	needed for atazanavir. The combination of
Rifampicin	Rifampicin is a strong CYP3A4 inducer and has been shown to	rifampicin and
	cause a 72% decrease in atazanavir	atazanavir is
	AUC which can result in virological	
	failure and resistance development.	section 4.3).
	During attempts to overcome the	
	decreased exposure by increasing	
	the dose of atazanavir or other	
	protease inhibitors with ritonavir, a	
	high frequency of liver reactions	
	was seen.	
ANTIPSYCHOTICS	I =	~
Quetiapine	Due to CYP3A4 inhibition by	Co-administration of
	atazanavir, concentrations of	quetiapine with

1.3.1.1 Samenvatting van de Productkenmerken

Medicinal products by	Interaction	Recommendations
therapeutic area	Interaction	concerning co-
therapeutic area		administration
	quetiapine are expected to increase	atazanavir is
	quettapme are expected to mercuse	contraindicated as
		atazanavir may increase
		quetiapine-related
		toxicity. Increased
		plasma concentrations
		of quetiapine may lead
		to coma (see section
		4.3)
Lurasidone	Atazanavir is expected to increase	Co-administration of
	plasma levels of lurasidone due to	lurasidone with
	CYP3A4 inhibition.	atazanavir is
		contraindicated as this
		may increase
		lurasidone-related
		toxicity (see section 4.3).
ACID REDUCING AGENTS		4.3).
H2-Receptor antagonists	,	
Without Tenofovir		
In HIV-infected patients with a	tazanavir/ritonavir at the	For patients not taking
recommended dose 300/100 m		tenofovir, if atazanavir
Famotidine 20 mg twice	Atazanavir AUC: ↓18% (↓25%	300 mg/ritonavir 100 mg
daily	↑1%)	and H ₂ -receptor
	Atazanavir C_{max} : $\downarrow 20\% (\downarrow 32\% \downarrow 7\%)$	antagonists are co-
	Atazanavir C_{min} : $\leftrightarrow 1\%$ ($\downarrow 16\%$	administered, a dose
	↑18%)	equivalent to famotidine
Famotidine 40 mg twice	Atazanavir AUC: ↓23% (↓32%	20 mg twice daily should
daily	↓14%)	not be exceeded. If a
	Atazanavir C_{max} : $\downarrow 23\%$ ($\downarrow 33\%$	higher dose of an H ₂ -receptor antagonist is
	12%)	required (e.g.,
	Atazanavir C_{min} : $\downarrow 20\%$ ($\downarrow 31\%$	famotidine 40 mg twice
In healthy walnut and with star	1 \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \	daily or equivalent) an
of 400/100 mg once daily	anavir/ritonavir at an increased dose	increase of the
Famotidine 40 mg twice	Atazanavir AUC: ↔3% (↓14%	atazanavir/ritonavir dose
daily	↑22%)	from 300/100 mg to
•	Atazanavir C_{max} : $\leftrightarrow 2\%$ ($\downarrow 13\%$	400/100 mg can be
	↑8%)	considered.
	Atazanavir C _{min} : ↓14% (↓32%	
	↑8%)	
With Tenofovir disoproxil fun disoproxil)	arate 300 mg once daily (equivalent to	245 mg tenofovir
In HIV-infected patients with a	tazanavir/ritonavir at the	For patients who are
recommended dose of 300/100		taking tenofovir
Famotidine 20 mg twice	Atazanavir AUC: \21% (\34%	disoproxil fumarate, if
daily	↓4%)*	atazanavir/ritonavir with
	Atazanavir C _{max} : ↓21% (↓36%	both tenofovir disoproxil
	↓4%)*	fumarate and an H ₂ -
	Atazanavir C _{min} : ↓19% (↓37%	receptor antagonist are

Medicinal products by	Interaction	Recommendations
therapeutic area		concerning co- administration
	<u> </u>	co-administered, a dose
Famotidine 40 mg twice daily	Atazanavir AUC: ↓24% (↓36% ↓11%)* Atazanavir C _{max} : ↓23% (↓36% ↓8%)* Atazanavir C _{min} : ↓25% (↓47% ↑7%)*	increase of atazanavir to 400 mg with 100 mg of ritonavir is recommended. A dose equivalent to famotidine 40 mg twice daily
In HIV-infected patients with a	tazanavir/ritonavir at an increased	should not be exceeded.
dose of 400/100 mg once daily		
Famotidine 20 mg twice daily	Atazanavir AUC: †18% (†6.5% †30%)*	
	Atazanavir C _{max} : ↑18% (↑6.7% ↑31%)*	
	Atazanavir C _{min} : ↑24 % (↑10% ↑39%)*	
Famotidine 40 mg twice daily	Atazanavir AUC: ↔2.3% (↓13% ↑10%)*	
	Atazanavir C_{max} : \leftrightarrow 5% (\downarrow 17% \uparrow 8.4%)*	
	Atazanavir C_{min} : $\leftrightarrow 1.3\%$ ($\downarrow 10\%$)*	
	*When compared to atazanavir 300 mg once daily with ritonavir 100 mg once daily and tenofovir disoproxil fumarate 300 mg all as a single dose with food. When compared to atazanavir 300 mg with ritonavir 100 mg without tenofovir disoproxil fumarate, atazanavir concentrations are expected to be additionally decreased by about 20%. The mechanism of interaction is decreased solubility of atazanavir as intra-gastric pH increases with H ₂ -blockers.	
Proton pump inhibitors		
Omeprazole 40 mg once daily (atazanavir 400 mg once daily with ritonavir 100 mg once daily)	Atazanavir (am): 2 hr after omeprazole Atazanavir AUC: ↓61% (↓65% ↓55%) Atazanavir C _{max} : ↓66% (↓62% ↓49%) Atazanavir C _{min} : ↓65% (↓71% ↓59%)	Co-administration of atazanavir with ritonavir and proton pump inhibitors is not recommended. If the combination is judged unavoidable, close clinical monitoring is
Omeprazole 20 mg once daily (atazanavir 400 mg once daily with ritonavir 100 mg	Atazanavir (am): 1 hr after omeprazole Atazanavir AUC: ↓30% (↓43% ↓14%)*	recommended in combination with an increase in the dose of atazanavir to 400 mg

Medicinal products by	Interaction	Recommendations
therapeutic area		concerning co-
*		administration
once daily)	Atazanavir C _{max} : ↓31% (↓42%	with 100 mg of
	↓17%)*	ritonavir; doses of
	Atazanavir C _{min} : \J31% (\J46%	proton pump inhibitors
	12%)*	comparable to
	\ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \	omeprazole 20 mg
	*When compared to atazanavir 300	should not be exceeded
	mg once daily with ritonavir 100	(see section 4.4).
	mg once daily.	(see section 1.1).
	The decrease in AUC, C_{max} , and	
	C _{min} was not mitigated when an	
	increased dose of	
	atazanavir/ritonavir (400/100 mg	
	once daily) was temporally	
	separated from omeprazole by 12	
	hours. Although not studied, similar	
	results are expected with other	
	proton pump inhibitors. This	
	decrease in atazanavir exposure	
	might negatively impact the	
	efficacy of atazanavir. The	
	mechanism of interaction is	
	decreased solubility of atazanavir as	
	intra-gastric pH increases with	
Autorita	proton pump inhibitors.	
Antacids	D. 1 1 . 1	A 4 1 1 1 1
Antacids and medicinal	Reduced plasma concentrations of	Atazanavir should be
products containing buffers	atazanavir may be the consequence	administered 2 hours before or 1 hour after
	of increased gastric pH if antacids,	
	including buffered medicinal	antacids or buffered
	products, are administered with	medicinal products.
ALDUA 1 ADDENODECED	atazanavir.	
ALPHA 1-ADRENORECEP		C 1 : :
Alfuzosin	Potential for increased alfuzosin	Co-administration of
	concentrations which can result in	alfuzosin with
	hypotension. The mechanism of	atazanavir is
	interaction is CYP3A4 inhibition by	contraindicated (see
ANTERCOACTE ANTEC	atazanavir and/or ritonavir.	section 4.3).
ANTICOAGULANTS	(0.040)	
Direct-acting oral anticoagula		0 1
Apixaban	Potential for increased apixaban	Co-administration of
Rivaroxaban	and rivaroxaban concentrations	apixaban or
	which can result in a higher risk of	rivaroxaban and
	bleeding.	atazanavir with
	The mechanism of interaction is	ritonavir is not
	inhibition of CYP3A4 / and P-gp	recommended.
	by atazanavir/ritonavir.	
	Ritonavir is a strong inhibitor of	
	both CYP3A4 and P-gp.	

Medicinal products by	Interaction	Recommendations
therapeutic area		concerning co- administration
	Atazanavir is an inhibitor of	uummst utton
	CYP3A4. The potential inhibition	
	of P-gp by atazanavir is unknown and cannot be excluded.	
Dabigatran	Potential for increased dabigatran	Co-administration of
	concentrations which can result in	dabigatran and
	a higher risk of bleeding. The mechanism of interaction is P-gp	atazanavir with ritonavir is not
	inhibition.	recommended.
	Ritonavir is a strong P-gp inhibitor.	
	Potential P-gp inhibition by	
	atazanavir is unknown and cannot	
D. I	be excluded.	D :
Edoxaban	Potential for increased edoxaban concentrations which can result in	Exercise caution when edoxaban is used with
	a higher risk of bleeding. The	atazanavir.
	mechanism of interaction is P-gp	
	inhibition by atazanavir/ritonavir.	Please refer to the
	Ritonavir is a strong P-gp	edoxaban SmPC sections 4.2 and 4.5 for
	inhibitor.	appropriate edoxaban
		dosage
	Potential P-gp inhibition by	recommendations for
	atazanavir is unknown and cannot be excluded.	co-administration with P-gp inhibitors.
Vitamin K antagonists	be exeruded.	1 -gp mmotors.
Warfarin	Co-administration with atazanavir	It is recommended that
	has the potential to increase or	the International
	decrease warfarin concentrations.	Normalised Ratio (INR) be monitored
		carefully during
		treatment with
		atazanavir, especially
		when commencing therapy.
ANTIEPILEPTICS		
Carbamazepine	Atazanavir may increase plasma	Carbamazepine should
	levels of carbamazepine due to CYP3A4 inhibition.	be used with caution in combination with
	Due to carbamazepine inducing	atazanavir. If necessary,
	effect, a reduction in atazanavir	monitor carbamazepine
	exposure cannot be ruled out.	serum concentrations
		and adjust the dose
		accordingly. Close monitoring of the
		patient's virologic
		response should be

Medicinal products by	Interaction	Recommendations
therapeutic area		concerning co- administration
Phenytoin, phenobarbital	Ritonavir may decrease plasma levels of phenytoin and/or phenobarbital due to CYP2C9 and CYP2C19 induction. Due to phenytoin/phenobarbital inducing effect, a reduction in atazanavir exposure cannot be ruled out.	excercised. Phenobarbital and phenytoin should be used with caution in combination with atazanavir/ritonavir. When atazanavir/ritonavir is co-administered with either phenytoin or phenobarbital, a dose adjustment of phenytoin or phenobarbital may be required.
		Close monitoring of patient's virologic response should be exercised.
Lamotrigine	Co-administration of lamotrigine and atazanavir/ritonavir may decrease lamotrigine plasma concentrations due to UGT1A4 induction.	Lamotrigine should be used with caution in combination with atazanavir/ritonavir. If necessary, monitor lamotrigine
		concentrations and adjust the dose accordingly.
ANTINEOPLASTICS AND I	MMUNOSUPRESSANTS	
Antineoplastics		
Apalutamide	The mechanism of interaction is CYP3A4 induction by apalutamide and CYP3A4 inhibition by atazanavir/ritonavir.	Co-administration with atazanavir (with or without ritonavir) is contraindicated due to the potential for decreased atazanavir and ritonavir plasma concentration with subsequent loss of virologic response and possible resistance to the class of protease inhibitors (see section 4.3). In addition, serum concentrations of apalutamide may be increased when coadministered with

1211	Companyatting	van da	Draduatkanmarkan
1.3.1.1	Samenvalling	van de	Productkenmerken

Medicinal products by therapeutic area	Interaction	Recommendations concerning co-
T. T		administration
		atazanavir/ritonavir, resulting in the potential for serious adverse events including seizure.
Encorafenib	The mechanism of interaction is CYP3A4 inhibition by atazanavir and/or ritonavir.	Co-administration of encorafenib with atazanavir (with or without ritonavir) is contraindicated due to the potential for loss of virologic response, development of resistance, increase in encorafenib plasma concentration and subsequent risk of serious adverse events such as QT interval prolongation (see section 4.3).
Ivosidenib	The mechanism of interaction is CYP3A4 inhibition by atazanavir and/or ritonavir.	Co-administration of ivosidenib with atazanavir (with or without ritonavir) is contraindicated due to potential for loss of virologic response, development of resistance, increase in ivosidenib plasma concentration and subsequent risk of serious adverse events such as QT interval prolongation (see section 4.3).
Irinotecan	Atazanavir inhibits UGT and may interfere with the metabolism of irinotecan, resulting in increased irinotecan toxicities.	If atazanavir is co- administered with irinotecan, patients should be closely monitored for adverse events related to irinotecan.
Immunosuppressants		
Cyclosporin	Concentrations of these	More frequent
Tacrolimus	immunosuppressants may be	therapeutic
Sirolimus	increased when co-administered with atazanavir due to CYP3A4 inhibition.	concentration monitoring of these medicinal products is recommended until

Medicinal products by	Interaction	Recommendations
therapeutic area	The action	concerning co-
		administration
		plasma levels have been
		stabilised.
CARDIOVASCULARAGEN	ITS	
Antiarrhythmics		
Amiodarone, Systemic	Concentrations of these	Caution is warranted
lidocaine, Quinidine	antiarrhythmics may be increased	and therapeutic
	when co-administered with	concentration
	atazanavir. The mechanism of	monitoring is
	amiodarone or systemic lidocaine/atazanavir interaction is	recommended when available. The
	CYP3A inhibition. Quinidine has a	concomitant use of
	narrow therapeutic window and is	quinidine is
	contraindicated due to potential	contraindicated (see
	inhibition of CYP3A by atazanavir.	section 4.3).
Calcium channel blockers	, amzanavii.	<u>, , , , , , , , , , , , , , , , , , , </u>
Bepridil	Atazanavir should not be used in	Co-administration with
	combination with medicinal	bepridil is contraindicated
	products that are substrates of	(see section 4.3).
	CYP3A4 and have a narrow	
700	therapeutic index.	
Diltiazem 180 mg once	Diltiazem AUC: †125% (†109%	An initial dose reduction
daily	141%)	of diltiazem by 50% is
(atazanavir 400 mg once daily)	Diltiazem C _{max} : ↑98% (↑78%	recommended, with subsequent titration as
daily)	Diltiazem C _{min} : †142% (†114%	needed and ECG
	173%)	monitoring.
		8
	Desacetyl-diltiazem AUC: ↑165%	
	(†145% †187%)	
	Desacetyl-diltiazem C _{max} : ↑172%	
	(†144% †203%)	
	Desacetyl-diltiazem C _{min} : †121%	
	(†102% †142%)	
	No significant effect on atazanavir	
	concentrations was observed.	
	There was an increase in the	
	maximum PR interval compared to	
	atazanavir alone. Co-	
	administration of diltiazem and	
	atazanavir/ritonavir has not been	
	studied. The mechanism of	
	diltiazem/atazanavir interaction is	
\$7	CYP3A4 inhibition.	C4:1 1.11
Verapamil	Serum concentrations of verapamil	Caution should be
	may be increased by atazanavir due to CYP3A4 inhibition.	exercised when
	to C 1 F 3A4 IIIII OIII OII.	verapamil is co- administered with
		atazanavir.
CORTICOSTEROIDS	1	

Medicinal products by therapeutic area	Interaction	Recommendations concerning co-
Dexamethasone and other corticosteroids (all routes of	Co-administration with dexamethasone or other	administration Co-administration with corticosteroids (all
administration)	corticosteroids that induce CYP3A may result in loss of therapeutic effect of atazanavir and development of resistance to atazanavir and/or ritonavir. Alternative corticosteroids should be considered. The mechanism of interaction is CYP3A4 induction by dexamethasone and CYP3A4 inhibition by atazanavir and/or ritonavir.	routes of administration) that are metabolized by CYP3A, particularly for long-term use, may increase the risk for development of systemic corticosteroid effects including Cushing's syndrome and adrenal suppression. The potential benefit of treatment versus the risk of systemic corticosteroid effects should be considered. For co-administration of cutaneously administered corticosteroids sensitive to CYP3A inhibition, consult the Summary of
		Product Characteristics of the corticosteroid for condition or uses that augment its systemic absorption.
Fluticasone propionate intranasal 50 µg 4 times daily for 7 days (ritonavir 100 mg capsules twice daily)	The fluticasone propionate plasma levels increased significantly, whereas the intrinsic cortisol levels decreased by approximately 86% (90% confidence interval 82%-89%). Greater effects may be	Co-administration of atazanavir/ritonavir and these glucocorticoids metabolised by CYP3A4 is not recommended unless the potential benefit of
And Inhaled/Nasal	expected when fluticasone propionate is inhaled. Systemic	treatment outweighs the risk of systemic corticosteroid effects
Corticosteroids	corticosteroid effects including Cushing's syndrome and adrenal suppression have been reported in patients receiving ritonavir and inhaled or intranasally administered fluticasone propionate; this could also occur with other corticosteroids metabolised via the P450 3A pathway, e.g., budesonide. The effects of high fluticasone systemic exposure on ritonavir plasma levels are yet unknown. The mechanism of interaction is	(see section 4.4). A dose reduction of the glucocorticoid should be considered with close monitoring of local and systemic effects or a switch to a glucocorticoid, which is not a substrate for CYP3A4 (e.g., beclomethasone). Moreover, in case of

Samenvatting van de Productkenmerken Januari 2025

Medicinal products by	Interaction	Recommendations
herapeutic area		concerning co-
		administration
	CYP3A4 inhibition.	withdrawal of
	Community and arrange of advances in	glucocorticoids,
	Concomitant use of atazanavir	progressive dose
	(with or without ritonavir) and other Inhaled/Nasal Corticosteroids is	3
	expected to produce the same	performed over a longer period.
	effects.	period.
	chects.	Concomitant use of
		Inhaled/Nasal
		Corticosteroids and
		atazanavir (with or
		without ritonavir) may
		increase plasma
		concentrations of
		Inhaled/Nasal
		corticosteroids. Use with
		caution. Consider
		alternatives to Inhaled/Nasal
		Corticosteroids,
		particularly for long-
		term use.
ERECTILE DYSFUNCTION	1	term ase.
PDE5 inhibitors		
Sildenafil, tadalafil,	Sildenafil, tadalafil and vardenafil	Patients should be
ardenafil	are metabolised by CYP3A4. Co-	warned about these
	administration with atazanavir may	possible side effects
	result in increased concentrations of	when using PDE5
	the PDE5 inhibitor and an increase	inhibitors for erectile
	in PDE5-associated adverse events,	dysfunction with atazanavir (see section
	including hypotension, visual changes, and priapism. The	4.4).
	mechanism of this interaction is	Also see PULMONARY
	CYP3A4 inhibition.	ARTERIAL
		HYPERTENSION in
		this table for further
		information regarding
		co-administration of
		atazanavir with
		sildenafil.
	SING HORMONE (GnRH) RECEP	
Elagolix	The mechanism of interaction is	Concomitant use of
	anticipated increase in elagolix exposure in the presence of	elagolix 200 mg twice daily with atazanavir
	CYP3A4 inhibition by atazanavir	(with or without
	and/or ritonavir.	ritonavir) for more than
		1 month is not
		1 month is not recommended due to the

Madiainal maduata ha	Internation	Recommendations
Medicinal products by	Interaction	
therapeutic area		concerning co-
		administration
		and hepatic
		transaminase elevations.
		Limit concomitant use
		of elagolix 150 mg once
		daily with atazanavir
		(with or without
		ritonavir) to 6 months.
KINASE INHIBITORS		
Fostamatinib	The mechanism of interaction is	Concomitant use of
	CYP3A4 inhibition by atazanavir	fostamatinib with
	and/or ritonavir.	atazanavir (with or
		without ritonavir) may
		increase the plasma
		concentration of R406,
		the active metabolite of
		fostamatinib. Monitor
		for toxicities of R406
		exposure resulting in
		dose-related adverse
		events such as
		hepatotoxicity and
		neutropenia.
		Fostamatinib dose
		reduction may be
HEDDAL DDODUCTS		required.
HERBAL PRODUCTS	Concomitant use of St. John's wort	Co-administration of
St. John's wort (Hypericum		
perforatum)	with atazanavir may be expected to	atazanavir with products
	result in significant reduction in	containing St. John's
	plasma levels of atazanavir. This	wort is contraindicated.
	effect may be due to an induction of	
	CYP3A4. There is a risk of loss of	
	therapeutic effect and development	
HODNOVAL COMPAGE	of resistance (see section 4.3).	
HORMONALCONTRACE		TC 1
Ethinyloestradiol 25 μg +	Ethinyloestradiol AUC: ↓19%	If an oral contraceptive
norgestimate	(↓25% ↓13%)	is administered with
(atazanavir 300 mg once	Ethinyloestradiol C _{max} : ↓16% (↓26%	atazanavir/ritonavir, it is
daily with ritonavir 100 mg	↓5%)	recommended that the
once daily)	Ethinyloestradiol C_{min} : $\downarrow 37\%$ ($\downarrow 45\%$	oral contraceptive
	↓29%)	contain at least 30 µg of
		ethinyloestradiol and
	Norgestimate AUC: ↑85% (↑67%	that the patient be
	↑105%)	reminded of strict
	Norgestimate C _{max} : ↑68% (↑51%	compliance with this
	188%)	contraceptive dosing
	Norgestimate C _{min} : $\uparrow 102\%$ ($\uparrow 77\%$	regimen. Co-
	↑131%)	administration of
	'	atazanavir/ritonavir with
	While the concentration of	other hormonal

1.3.1.1 Samenvatung van de Productkenmerken			
Medicinal products by therapeutic area	Interaction	Recommendations concerning co-administration	
	ethinyloestradiol was increased with atazanavir given alone, due to both UGT and CYP3A4 inhibition by atazanavir, the net effect of atazanavir/ritonavir is a decrease in ethinyloestradiol levels because of the inducing effect of ritonavir. The increase in progestin exposure may lead to related side-effects	contraceptives or oral contraceptives containing progestogens other than norgestimate has not been studied, and therefore should be avoided. An alternate reliable method of contraception is recommended.	
	(e.g., insulin resistance, dyslipidemia, acne and spotting), thus possibly affecting the compliance.		
Ethinyloestradiol 35 µg + norethindrone (atazanavir 400 mg once daily)	Ethinyloestradiol AUC: \uparrow 48% (\uparrow 31% \uparrow 68%) Ethinyloestradiol C _{max} : \uparrow 15% (\downarrow 1% \uparrow 32%) Ethinyloestradiol C _{min} : \uparrow 91% (\uparrow 57% \uparrow 133%)		
	Norethindrone AUC: $\uparrow 110\%$ ($\uparrow 68\%$) $\uparrow 162\%$) Norethindrone C_{max} : $\uparrow 67\%$ ($\uparrow 42\%$) $\uparrow 196\%$) Norethindrone C_{min} : $\uparrow 262\%$ ($\uparrow 157\%$) $\uparrow 409\%$)		
	The increase in progestin exposure may lead to related side-effects (e.g., insulin resistance, dyslipidemia, acne and spotting), thus possibly affecting the compliance.		
LIPID MODIFYING AGEN			
HMG-CoA reductase inhibitor Simvastatin Lovastatin		Co-administration of	
	Simvastatin and lovastatin are highly dependent on CYP3A4 for their metabolism and co-administration with atazanavir may result in increased concentrations.	simvastatin or lovastatin with atazanavir is contraindicated due to an increased risk of myopathy including rhabdomyolysis (see section 4.3).	
Atorvastatin	The risk of myopathy including rhabdomyolysis may also be increased with atorvastatin, which is also metabolised by CYP3A4.	Co-administration of atorvastatin with atazanavir is not recommended. If the use of atorvastatin is considered strictly necessary, the lowest	

Medicinal products by	Interaction	Recommendations
therapeutic area		concerning co-
		administration
		possible dose of atorvastatin should be administered with careful safety monitoring (see section 4.4).
Pravastatin	Although not studied, there is a	Caution should be
Fluvastatin	potential for an increase in pravastatin or fluvastatin exposure when coadministered with protease inhibitors. Pravastatin is not metabolised by CYP3A4. Fluvastatin is partially metabolised by CYP2C9.	exercised.
Other lipid-modifying agents		
Lomitapide	Lomitapide is highly dependent on CYP3A4 for metabolism and coadministration with atazanavir with ritonavir may result in increased concentrations.	Co-administration of lomitapide and atazanavir with ritonavir is contraindicated due to a potential risk of markedly increased transaminase levels and hepatotoxicity (see section 4.3).
INHALED BETA AGONIS		
Salmeterol	Co-administration with atazanavir may result in increased concentrations of salmeterol and an increase in salmeterol-associated adverse events. The mechanism of interaction is CYP3A4 inhibition by atazanavir and/or ritonavir.	Co-administration of salmeterol with atazanavir is not recommended (see section 4.4).
OPIOIDS	and/or ritonavir.	
Buprenorphine, once daily, stable maintenance dose (atazanavir 300 mg once daily with ritonavir 100 mg once daily)	Buprenorphine AUC: ↑67% Buprenorphine C _{max} : ↑37% Buprenorphine C _{min} : ↑69% Norbuprenorphine AUC: ↑105% Norbuprenorphine C _{max} : ↑61% Norbuprenorphine C _{min} : ↑101% The mechanism of interaction is CYP3A4 and UGT1A1 inhibition. Concentrations of atazanavir (when given with ritonavir) were not significantly affected.	Co-administration with atazanavir with ritonavir warrants clinical monitoring for sedation and cognitive effects. A dose reduction of buprenorphine may be considered.
Methadone, stable	No significant effect on methadone	No dosage adjustment is
maintenance dose	concentrations was observed. Given	necessary if methadone

1.3.1.1 Samenvatting van de Pro	Interaction	Dogommondotions
Medicinal products by therapeutic area	interaction	Recommendations concerning co-administration
(atazanavir 400 mg once daily)	that low dose ritonavir (100 mg twice daily) has been shown to have no significant effect on methadone concentrations, no interaction is expected if methadone is coadministered with atazanavir, based on these data.	is co-administered with atazanavir.
PULMONARY ARTERIAI		
PDE5 inhibitors		
Sildenafil SEDATIVES	Co-administration with atazanavir may result in increased concentrations of the PDE5 inhibitor and an increase in PDE5-inhibitor-associated adverse events. The mechanism of interaction is CYP3A4 inhibition by atazanavir and/or ritonavir.	A safe and effective dose in combination with atazanavir has not been established for sildenafil when used to treat pulmonary arterial hypertension. Sildenafil, when used for the treatment of pulmonary arterial hypertension, is contraindicated (see section 4.3).
Midazolam Triazolam	Midazolam and triazolam are extensively metabolised by CYP3A4. Co-administration with atazanavir may cause a large increase in the concentration of these benzodiazepines. No drug interaction study has been performed for the co-administration of atazanavir with benzodiazepines. Based on data for other CYP3A4 inhibitors, plasma concentrations of midazolam are expected to be significantly higher when midazolam is given orally. Data from concomitant use of parenteral midazolam with other protease inhibitors suggest a possible 3-4-fold increase in midazolam plasma levels.	Co-administration of atazanavir with triazolam or orally administered midazolam is contraindicated (see section 4.3), whereas caution should be used with co-administration of atazanavir and parenteral midazolam. If atazanavir is co-administered with parenteral midazolam, it should be done in an intensive care unit (ICU) or similar setting which ensures close clinical monitoring and appropriate medical management in case of respiratory depression and/or prolonged sedation. Dosage adjustment for midazolam should be considered, especially if more than a single dose

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Medicinal products by therapeutic area	Interaction	Recommendations concerning co-administration
		of midazolam is administered.

In case of withdrawal of ritonavir from the recommended atazanavir-boosted regimen (see section 4.4)

The same recommendations for drug interactions would apply except:

- that co-administration is not recommended with tenofovir, carbamazepine, phenytoin, phenobarbital, proton pump inhibitors, and buprenorphine.
- that co-administration with famotidine is not recommended but if required, atazanavir without ritonavir should be administered either 2 hours after famotidine or 12 hours before. No single dose of famotidine should exceed 20 mg, and the total daily dose of famotidine should not exceed 40 mg.
- the need to consider that
 - co-administration of apixaban, dabigatran, or rivaroxaban and atazanavir without ritonavir may affect apixaban, dabigatran, or rivaroxaban concentrations
 - co-administration of voriconazole and atazanavir without ritonavir may affect atazanavir concentrations
 - co-administration of fluticasone and atazanavir without ritonavir may increase fluticasone concentrations relative to fluticasone given alone
 - if an oral contraceptive is administered with atazanavir without ritonavir, it is recommended that the oral contraceptive contain no more than 30 μg of ethinyloestradiol
 - no dose adjustment of lamotrigine is required

Paediatric population

Interaction studies have only been performed in adults.

4.6 Fertility, pregnancy and lactation

Pregnancy

A moderate amount of data in pregnant women (between 300-1000 pregnancy outcomes) indicate no malformative toxicity of atazanavir. Animal studies do not indicate reproductive toxicity (see section 5.3). The use of [Nationally completed name] with ritonavir may be considered during pregnancy only if the potential benefit justifies the potential risk.

In clinical trial AI424-182 atazanavir/ritonavir (300/100 mg or 400/100 mg) in combination with zidovudine/lamivudine was administered to 41 pregnant women during the second or third trimester. Six of 20 (30%) women on atazanavir/ritonavir 300/100 mg and 13 of 21 (62%) women on atazanavir/ritonavir 400/100 mg experienced grades 3 to 4 hyperbilirubinaemia. There were no cases of lactic acidosis observed in the clinical trial AI424-182.

The study assessed 40 infants who received antiretroviral prophylactic treatment (which did not include atazanavir) and were negative for HIV-1 DNA at the time of delivery and/or during the first 6 months postpartum. Three of 20 infants (15%) born to women treated with atazanavir/ritonavir 300/100 mg and four of 20 infants (20%) born to women treated with atazanavir/ritonavir 400/100 mg experienced grade 3-4 bilirubin. There was no evidence of pathologic jaundice and six of 40 infants in this study received phototherapy for a maximum of 4 days. There were no reported cases of kernicterus in neonates.

For dosing recommendations see section 4.2 and for pharmacokinetic data see section 5.2.

It is not known whether atazanavir with ritonavir administered to the mother during pregnancy will exacerbate physiological hyperbilirubinaemia and lead to kernicterus in neonates and infants. In the prepartum period, additional monitoring should be considered.

Breast-feeding

Atazanavir has been detected in human milk. In order to avoid transmission of HIV to the infant it is recommended that women living with HIV do not breast-feed their infants.

Fertility

In a nonclinical fertility and early embryonic development study in rats, atazanavir altered oestrus cycling with no effects on mating or fertility (see section 5.3).

4.7 Effects on ability to drive and use machines

Patients should be informed that dizziness has been reported during treatment with regimens containing [Nationally completed name] (see section 4.8).

4.8 Undesirable effects

Summary of the safety profile

Atazanavir has been evaluated for safety in combination therapy with other antiretroviral medicinal products in controlled clinical trials in 1,806 adult patients receiving atazanavir 400 mg once daily (1,151 patients, 52 weeks median duration and 152 weeks maximum duration) or atazanavir 300 mg with ritonavir 100 mg once daily (655 patients, 96 weeks median duration and 108 weeks maximum duration).

Adverse reactions were consistent between patients who received atazanavir 400 mg once daily and patients who received atazanavir 300 mg with ritonavir 100 mg once daily, except that jaundice and elevated total bilirubin levels were reported more frequently with atazanavir plus ritonavir.

Among patients who received atazanavir 400 mg once daily or atazanavir 300 mg with ritonavir 100 mg once daily, the only adverse reactions of any severity reported very commonly with at least a possible relationship to regimens containing atazanavir and one or more NRTIs were nausea (20%), diarrhoea (10%), and jaundice (13%). Among patients receiving atazanavir 300 mg with ritonavir 100 mg, the frequency of jaundice was 19%. In the majority of cases, jaundice was reported within a few days to a few months after the initiation of treatment (see section 4.4).

Chronic kidney disease in HIV-infected patients treated with atazanavir, with or without ritonavir, has been reported during postmarketing surveillance. A large prospective observational study has shown an association between an increased incidence of chronic kidney disease and cumulative exposure to atazanavir/ritonavir-containing regimen in HIV-infected patients with an initially normal eGFR. This association was observed independently of exposure to tenofovir disoproxil. Regular monitoring of the renal function of patients should be maintained throughout the treatment duration (see section 4.4).

Tabulated list of adverse reactions

Assessment of adverse reactions for atazanavir is based on safety data from clinical studies and post-marketing experience. Frequency is defined using the following convention: very common ($\geq 1/10$), common ($\geq 1/100$) to < 1/100), uncommon ($\geq 1/100$), rare ($\geq 1/10,000$ to <

1/1,000), very rare (< 1/10,000). Within each frequency grouping, undesirable effects are presented in order of decreasing seriousness.

Immune system disorders:	uncommon: hypersensitivity
Metabolism and nutrition disorders:	uncommon: weight decreased, weight gain,
	anorexia, appetite increased
Psychiatric disorders:	uncommon: depression, disorientation,
	anxiety, insomnia, sleep disorder, abnormal
	dream
Nervous system disorders:	common: headache;
	uncommon: peripheral neuropathy, syncope,
	amnesia, dizziness, somnolence, dysgeusia
Eye disorders:	common: ocular icterus
Cardiac disorders:	uncommon: torsades de pointes ^a
	rare: QTc prolongation ^a , oedema, palpitation
Vascular disorders:	uncommon: hypertension
Respiratory, thoracic and mediastinal	uncommon: dyspnoea
disorders:	
Gastrointestinal disorders:	common: vomiting, diarrhoea, abdominal
	pain, nausea, dyspepsia;
	uncommon: pancreatitis, gastritis, abdominal
	distension, stomatitis aphthous, flatulence,
	dry mouth
Hepatobiliary disorders:	common: jaundice;
	uncommon: hepatitis, cholelithiasis ^a ,
	cholestasis ^a ;
	rare: hepatosplenomegaly, cholecystitis ^a
Skin and subcutaneous tissue disorders:	common: rash;
	uncommon: erythemia multiforme ^{a,b} , toxic
	skin eruptions ^{a,b} , drug rash with eosinophilia
	and systemic symptoms (DRESS)
	syndrome ^{a,b} , angioedema ^a , urticaria, alopecia,
	pruritus;
	rare: Stevens-Johnson syndrome ^{a,b} ,
	vesiculobullous rash, eczema, vasodilatation
Musculoskeletal and connective tissue	uncommon: muscle atrophy, arthralgia,
disorders:	myalgia;
	rare: myopathy
Renal and urinary disorders:	uncommon: nephrolithiasis ^a , haematuria,
	proteinuria, pollakiuria, interstitial nephritis,
	chronic kidney disease ^a ;
	rare: kidney pain
Reproductive system and breast disorders:	uncommon: gynaecomastia
General disorders and administration site	common: fatigue;
conditions:	uncommon: chest pain, malaise, pyrexia,
	asthenia;
	rare: gait disturbance

^a These adverse reactions were identified through post-marketing surveillance, however, the frequencies were estimated from a statistical calculation based on the total number of patients exposed to atazanavir in randomised controlled and other available clinical trials (n = 2321).

^b See description of selected adverse reactions for more details.

Description of selected adverse reactions

1.3.1.1 Samenvatting van de Productkenmerken

In HIV-infected patients with severe immune deficiency at the time of initiation of combination antiretroviral therapy (CART), an inflammatory reaction to asymptomatic or residual opportunistic infections may arise. Autoimmune disorders (such as Graves' disease and autoimmune hepatitis) have also been reported; however, the reported time to onset is more variable and these events can occur many months after initiation of treatment (see section 4.4).

Cases of osteonecrosis have been reported, particularly in patients with generally acknowledged risk factors, advanced HIV disease or long-term exposure to combination antiretroviral therapy (CART). The frequency of this is unknown (see section 4.4).

Metabolic parameters

Weight and levels of blood lipids and glucose may increase during antiretroviral therapy (see section 4.4).

Rash and associated syndromes

Rashes are usually mild-to-moderate maculopapular skin eruptions that occur within the first 3 weeks of starting therapy with atazanavir.

Stevens-Johnson syndrome (SJS), erythema multiforme, toxic skin eruptions and drug rash with eosinophilia and systemic symptoms (DRESS) syndrome have been reported with the use of atazanavir (see section 4.4).

Laboratory abnormalities

The most frequently reported laboratory abnormality in patients receiving regimens containing atazanavir and one or more NRTIs was elevated total bilirubin reported predominantly as elevated indirect [unconjugated] bilirubin (87% Grade 1, 2, 3, or 4). Grade 3 or 4 elevation of total bilirubin was noted in 37% (6% Grade 4). Among experienced patients treated with atazanavir 300 mg once daily with 100 mg ritonavir once daily for a median duration of 95 weeks, 53% had Grade 3-4 total bilirubin elevations. Among naive patients treated with atazanavir 300 mg once daily with 100 mg ritonavir once daily for a median duration of 96 weeks, 48% had Grade 3-4 total bilirubin elevations (see section 4.4).

Other marked clinical laboratory abnormalities (Grade 3 or 4) reported in $\geq 2\%$ of patients receiving regimens containing atazanavir and one or more NRTIs included: elevated creatine kinase (7%), elevated alanine aminotransferase/serum glutamic-pyruvic transaminase (ALT/SGPT) (5%), low neutrophils (5%), elevated aspartate aminotransferase/serum glutamic-oxaloacetic transaminase (AST/SGOT) (3%), and elevated lipase (3%).

Two percent of patients treated with atazanavir experienced concurrent Grade 3-4 ALT/AST and Grade 3-4 total bilirubin elevations.

Paediatric population

In a clinical study AI424-020, paediatric patients 3 months to less than 18 years of age who received either the oral powder or capsule formulation had a mean duration of treatment with atazanavir of 115 weeks. The safety profile in this study was overall comparable to that seen in adults. Both asymptomatic first-degree (23%) and second-degree (1%) atrioventricular block were reported in paediatric patients. The most frequently reported laboratory abnormality in paediatric patients receiving atazanavir was elevation of total bilirubin (≥ 2.6 times ULN, Grade 3-4) which occurred in 45% of patients.

In clinical studies AI424-397 and AI424-451, paediatric patients 3 months to less than 11 years of age had a mean duration of treatment with atazanavir oral powder of 80 weeks. No deaths

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were reported. The safety profile in these studies was overall comparable to that seen in previous paediatric and adult studies. The most frequently reported laboratory abnormalities in paediatric patients receiving atazanavir oral powder was elevation of total bilirubin (≥ 2.6 times ULN, Grade 3-4; 16%) and increased amylase (Grade 3-4; 33%), generally of nonpancreatic origin. Elevation in ALT levels were more frequently reported in paediatric patients in these studies than in adults.

Other special populations

Patients co-infected with hepatitis B and/or hepatitis C virus

Among 1,151 patients receiving atazanavir 400 mg once daily, 177 patients were co-infected with chronic hepatitis B or C, and among 655 patients receiving atazanavir 300 mg once daily with ritonavir 100 mg once daily, 97 patients were co-infected with chronic hepatitis B or C. Co-infected patients were more likely to have baseline hepatic transaminase elevations than those without chronic viral hepatitis. No differences in frequency of bilirubin elevations were observed between these patients and those without viral hepatitis. The frequency of treatment emergent hepatitis or transaminase elevations in co-infected patients was comparable between atazanavir and comparator regimens (see section 4.4).

Reporting of suspected adverse reactions

Reporting suspected adverse reactions after authorisation of the medicinal product is important. It allows continued monitoring of the benefit/risk balance of the medicinal product. Healthcare professionals are asked to report any suspected adverse reactions via the national reporting system listed in Appendix V.

4.9 Overdose

Human experience of acute overdose with atazanavir is limited. Single doses up to 1,200 mg have been taken by healthy volunteers without symptomatic untoward effects. At high doses that lead to high drug exposures, jaundice due to indirect (unconjugated) hyperbilirubinaemia (without associated liver function test changes) or PR interval prolongations may be observed (see sections 4.4 and 4.8).

Treatment of overdose with atazanavir should consist of general supportive measures, including monitoring of vital signs and electrocardiogram (ECG), and observations of the patient's clinical status. If indicated, elimination of unabsorbed atazanavir should be achieved by emesis or gastric lavage. Administration of activated charcoal may also be used to aid removal of unabsorbed drug. There is no specific antidote for overdose with [Nationally completed name]. Since atazanavir is extensively metabolised by the liver and is highly protein bound, dialysis is unlikely to be beneficial in significant removal of this medicinal product.

5. PHARMACOLOGICAL PROPERTIES

5.1 Pharmacodynamic properties

Pharmacotherapeutic group: antivirals for systemic use, protease inhibitors, ATC code: J05AE08

Mechanism of action

Atazanavir is an azapeptide HIV-1 protease inhibitor (PI). The compound selectively inhibits the virus-specific processing of viral Gag-Pol proteins in HIV-1 infected cells, thus preventing formation of mature virions and infection of other cells.

Antiviral activity in vitro: atazanavir exhibits anti-HIV-1 (including all clades tested) and anti-HIV-2 activity in cell culture.

Resistance

Antiretroviral treatment naive adult patients

In clinical trials of antiretroviral treatment-naive patients treated with unboosted atazanavir, the I50L substitution, sometimes in combination with an A71V change, is the signature resistance substitution for atazanavir. Resistance levels to atazanavir ranged from 3.5- to 29-fold without evidence of phenotypic cross resistance to other PIs. In clinical trials of antiretroviral treatment-naive patients treated with boosted atazanavir, the I50L substitution did not emerge in any patient without baseline PI substitutions. The N88S substitution has been rarely observed in patients with virologic failure on atazanavir (with or without ritonavir). While it may contribute to decreased susceptibility to atazanavir when it occurs with other protease substitutions, in clinical studies N88S by itself does not always lead to phenotypic resistance to atazanavir or have a consistent impact on clinical efficacy.

Table 3: De novo substitutions in treatment-naive patients failing therapy with atazanavir + ritonavir (Study 138, 96 weeks)

Frequency	de novo PI substitution (n=26) ^a		
>20%	none		
10-20%	none		

^a Number of patients with paired genotypes classified as virological failures (HIV RNA \geq 400 copies/mL).

The M184I/V substitution emerged in 5/26 atazanavir/ritonavir and 7/26 lopinavir/ritonavir virologic failure patients, respectively.

Antiretroviral treatment experienced adult patients

In antiretroviral treatment experienced patients from Studies 009, 043, and 045, 100 isolates from patients designated as virological failures on therapy that included either atazanavir, atazanavir + ritonavir, or atazanavir + saquinavir were determined to have developed resistance to atazanavir. Of the 60 isolates from patients treated with either atazanavir or atazanavir + ritonavir, 18 (30%) displayed the I50L phenotype previously described in naive patients.

Table 4: De novo substitutions in treatment experienced patients failing therapy with atazanavir + ritonavir (Study 045, 48 weeks)

Frequency	de novo PI substitution (n=35) ^{a,b}
>20%	M36, M46, I54, A71, V82
10-20%	L10, I15, K20, V32, E35, S37, F53, I62, G73, I84, L90

^a Number of patients with paired genotypes classified as virological failures (HIV RNA \geq 400 copies/mL).

None of the de novo substitutions (see Table 4) are specific to atazanavir and may reflect reemergence of archived resistance on atazanavir + ritonavir in Study 045 treatment-experienced population.

The resistance in antiretroviral treatment experienced patients mainly occurs by accumulation of the major and minor resistance substitutions described previously to be involved in protease inhibitor resistance.

^b Ten patients had baseline phenotypic resistance to atazanavir + ritonavir (fold change [FC]>5.2). FC susceptibility in cell culture relative to the wild-type reference was assayed using PhenoSenseTM (Monogram Biosciences, South San Francisco, California, USA)

Clinical results

In antiretroviral naive adult patients

Study 138 is an international randomised, open-label, multicenter, prospective trial of treatment-naïve patients comparing atazanavir/ritonavir (300 mg/100 mg once daily) to lopinavir/ritonavir (400 mg/100 mg twice daily), each in combination with fixed-dose tenofovir disoproxil fumarate/emtricitabine (300 mg/200 mg tablets once daily). The atazanavir/ritonavir arm showed similar (non-inferior) antiviral efficacy compared to the lopinavir/ritonavir arm, as assessed by the proportion of patients with HIV RNA < 50 copies/mL at Week 48 (Table 5). Analyses of data through 96 weeks of treatment demonstrated durability of antiviral activity (Table 5).

Table 5: Efficacy Outcomes in Study 138 a

Sable 5:Efficacy Out	comes in Study 13			
Parameter	Atazanavir/ritonavir ^b (300 mg/100 mg once daily) n=440		Lopinavir/ritonavir ^c (400 mg/100 mg twice daily) n=443	
	Week 48	Week 96	Week 48	Week 96
HIV RNA <50 copies/mL,	%			
All patients ^d	78	74	76	68
Difference estimate [95% CI] ^d		Week 48: 1.7% Week 96: 6.1%		
Per protocol analysis ^e	86	91	89	89
	$(n=392^{f})$	(n=352)	(n=372)	(n=331)
Difference estimate ^e		Week 48: -3%	[-7.6%, 1.5%]	,
[95% CI]		Week 96: 2.2%	[-2.3%, 6.7%]	
HIV RNA <50 copies/mL,	% by Baseline Ch	aracteristic ^d		
HIV RNA				
<100,000 copies/mL	82 (n=217)	75 (n=217)	81 (n=218)	70 (n=218)
≥100,000 copies/mL	74 (n=223)	74 (n=223)	72 (n=225)	66 (n=225)
CD4 count <50 cells/mm ³	78 (n=58)	78 (n=58)	63 (n=48)	58 (n=48)
50 to <100 cells/mm ³	76 (n=45)	71 (n=45)	69 (n=29)	69 (n=29)
100 to <200 cells/mm ³	75 (n=106)	71 (n=106)	78 (n=134)	70 (n=134)
\geq 200 cells/mm ³	80 (n=222)	76 (n=222)	80 (n=228)	69 (n=228)
HIV RNA Mean Change f		\ /	· / 1	, ,
All patients	-3.09 (n=397)	-3.21 (n=360)	-3.13 (n=379)	-3.19 (n=340)
CD4 Mean Change from I	Baseline, cells/mm ³			, ,
All patients	203 (n=370)	268 (n=336)	219 (n=363)	290 (n=317)
CD4 Mean Change from Ba	aseline, cells/mm ³ b	y Baseline Charact	eristic	
HIV RNA	179 (n=183)	243 (n=163)	194 (n=183)	267 (n=152)
<100,000 copies/mL		·	·	
≥100,000 copies/mL	227 (n=187)	291 (n=173)	245 (n=180)	310 (n=165)

a Mean baseline CD4 cell count was 214 cells/mm³ (range 2 to 810 cells/mm³) and mean baseline plasma HIV-1 RNA was 4.94 log₁₀ copies/mL (range 2.6 to 5.88 log₁₀ copies/mL)

Data on withdrawal of ritonavir from atazanavir boosted regimen (see also section 4.4)

^b Atazanavir/RTV with tenofovir disoproxil fumarate/emtricitabine (fixed-dose 300 mg/200 mg tablets once daily).

^c Lopinavir/RTV with tenofovir disoproxil fumarate/emtricitabine (fixed-dose 300 mg/200 mg tablets once daily).

d Intent-to-treat analysis, with missing values considered as failures.

^e Per protocol analysis: Excluding non-completers and patients with major protocol deviations.

f Number of patients evaluable.

Study 136 (INDUMA)

In an open-label, randomised, comparative study following a 26- to 30-week induction phase with atazanavir 300 mg + ritonavir 100 mg once daily and two NRTIs, unboosted atazanavir 400 mg once daily and two NRTIs administered during a 48-week maintenance phase (n=87) had similar antiviral efficacy compared with atazanavir + ritonavir and two NRTIs (n=85) in HIV-infected subjects with fully suppressed HIV replication, as assessed by the proportion of subjects with HIV RNA < 50 copies/mL: 78% of subjects on unboosted atazanavir and two NRTIs compared with 75% on atazanavir + ritonavir and two NRTIs.

Eleven subjects (13%) in the unboosted atazanavir group and 6 (7%) in the atazanavir + ritonavir group, had virologic rebound. Four subjects in the unboosted atazanavir group and 2 in the atazanavir + ritonavir group had HIV RNA > 500 copies/mL during the maintenance phase. No subject in either group showed emergence of protease inhibitor resistance. The M184V substitution in reverse transcriptase, which confers resistance to lamivudine and emtricitabine, was detected in 2 subjects in the unboosted atazanavir and 1 subject in the atazanavir + ritonavir group.

There were fewer treatment discontinuations in the unboosted atazanavir group (1 vs. 4 subjects in the atazanavir + ritonavir group). There was less hyperbilirubinaemia and jaundice in the unboosted atazanavir group compared with the atazanavir + ritonavir group (18 and 28 subjects, respectively).

In antiretroviral experienced adult patients

<u>Study 045</u> is a randomised, multicenter trial comparing atazanavir /ritonavir (300/100 mg once daily) and atazanavir/saquinavir (400/1,200 mg once daily), to lopinavir + ritonavir (400/100 mg fixed-dose combination twice daily), each in combination with tenofovir disoproxil fumarate (see sections 4.5 and 4.8) and one NRTI, in patients with virologic failure on two or more prior regimens containing at least one PI, NRTI, and NNRTI. For randomised patients, the mean time of prior antiretroviral exposure was 138 weeks for PIs, 281 weeks for NRTIs, and 85 weeks for NNRTIs. At baseline, 34% of patients were receiving a PI and 60% were receiving an NNRTI. Fifteen of 120 (13%) patients in the atazanavir + ritonavir treatment arm and 17 of 123 (14%) patients in the lopinavir + ritonavir arm had four or more of the PI substitutions L10, M46, I54, V82, I84, and L90. Thirty-two percent of patients in the study had a viral strain with fewer than two NRTI substitutions.

The primary endpoint was the time-averaged difference in change from baseline in HIV RNA through 48 weeks (Table 6).

Table 6: Efficacy Outcomes at Week 48^a and at Week 96 (Study 045)

Parameter	mg/ 100	ATV/RTV ^b (300 ng/ 100 mg once daily) n=120		LPV/RTV ^c (400 mg/ 100 mg twice daily) n=123		Time-averaged difference ATV/RTV-LPV/RTV [97.5% CI ^d]	
	Week 48	Week 96	Week 48	Week 96	Week 48	Week 96	
HIV RNA Me	an Change froi	m Baseline, log	g ₁₀ copies/mL				
A 11	-1.93	-2.29	-1.87	-2.08	0.13	0.14	
All patients	(n=90	(n=64)	(n=99)	(n=65)	[-0.12, 0.39]	[-0.13, 0.41]	
HIV RNA <50 copies/mL, % ¹ (responder/evaluable)							
All patients	36 (43/120)	32 (38/120)	42 (52/123)	35 (41/118)	NA	NA	
HIV RNA <50 copies/mL by select baseline PI substitutions, 1, g % (responder/evaluable)							
0-2	44 (28/63)	41 (26/63)	56 (32/57)	48 (26/54)	NA	NA	
3	18 (2/11)	9 (1/11)	38 (6/16)	33 (5/15)	NA	NA	
≥4	27 (12/45)	24 (11/45)	28 (14/50)	20 (10/49)	NA	NA	

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CD4 Mean Change from Baseline, cells/mm ³						
All patients	110 (n=83)	122 (n=60)	121 (n=94)	154 (n=60)	NA	NA

^a The mean baseline CD4 cell count was 337 cells/mm³ (range: 14 to 1,543 cells/mm³) and the mean baseline plasma HIV-1 RNA level was 4.4 log₁₀ copies/mL (range: 2.6 to 5.88 log₁₀ copies/mL).

g Select substitutions include any change at positions L10, K20, L24, V32, L33, M36, M46, G48, I50, I54, L63, A71, G73, V82, I84, and L90 (0-2, 3, 4 or more) at baseline. NA = not applicable.

Through 48 weeks of treatment, the mean changes from baseline in HIV RNA levels for atazanavir + ritonavir and lopinavir + ritonavir were similar (non-inferior). Consistent results were obtained with the last observation carried forward method of analysis (time-averaged difference of 0.11, 97.5% confidence interval [-0.15, 0.36]). By as-treated analysis, excluding missing values, the proportions of patients with HIV RNA < 400 copies/mL (< 50 copies/mL) in the atazanavir + ritonavir arm and the lopinavir + ritonavir arm were 55% (40%) and 56% (46%), respectively.

Through 96 weeks of treatment, mean HIV RNA changes from baseline for atazanavir + ritonavir and lopinavir + ritonavir met criteria for non-inferiority based on observed cases. Consistent results were obtained with the last observation carried forward method of analysis. By as-treated analysis, excluding missing values, the proportions of patients with HIV RNA <400 copies/mL (<50 copies/mL) for atazanavir + ritonavir were 84% (72%) and for lopinavir + ritonavir were 82% (72%). It is important to note that at time of the 96-week analysis, 48 % of patients overall remained on study. Atazanavir + saquinavir was shown to be inferior to lopinavir + ritonavir.

Paediatric population

Assessment of the pharmacokinetics, safety, tolerability, and efficacy of atazanavir is based on data from the open-label, multicenter clinical trial AI424-020 conducted in patients from 3 months to 21 years of age. Overall in this study, 182 paediatric patients (81 antiretroviral-naive and 101 antiretroviral-experienced) received once daily atazanavir (capsule or powder formulation), with or without ritonavir, in combination with two NRTIs.

The clinical data derived from this study are inadequate to support the use of atazanavir (with or without ritonavir) in children below 6 years of age.

Efficacy data observed in the 41 paediatric patients aged 6 years to less than 18 years that received atazanavir capsules with ritonavir are presented in Table 7. For treatment-naive paediatric patients, the mean baseline CD4 cell count was 344 cells/mm³ (range: 2 to 800 cells/ mm³) and mean baseline plasma HIV-1 RNA was 4.67 log₁₀ copies/mL (range: 3.70 to 5.00 log₁₀ copies/mL). For treatment- experienced paediatric patients, the mean baseline CD4 cell count was 522 cells/mm³ (range: 100 to 1157 cells/ mm³) and mean baseline plasma HIV-1 RNA was 4.09 log₁₀ copies/mL (range: 3.28 to 5.00 log₁₀ copies/mL).

Efficacy Outcomes (paediatric patients 6 years to less than 18 years of Table 7: age) at Week 48 (Study AI424-020)

^b ATV/RTV with tenofovir disoproxil fumarate/emtricitabine (fixed-dose 300 mg/200 mg tablets once daily).

^c LPV/RTV with tenofovir disoproxil fumarate/emtricitabine (fixed-dose 300 mg/200 mg tablets once daily).

d Confidence interval.

^e Number of patients evaluable.

f Intent-to-treat analysis, with missing values considered as failures. Responders on LPV/RTV who completed treatment before Week 96 are excluded from Week 96 analysis. The proportion of patients with HIV RNA < 400 copies/mL were 53% and 43% for ATV/RTV and 54% and 46% for LPV/RTV at Weeks 48 and 96 respectively.

Parameter	Treatment-Naive atazanavir Capsules/ritonavir (300 mg/100 mg once daily) n=16	Treatment- Experienced atazanavir Capsules/ritonavir (300 mg/100 mg once daily) n=25
HIV RNA <50 copies/mL, %) a	
All patients	81 (13/16)	24 (6/25)
HIV RNA <400 copies/mL,	2/o ^a	
All patients	88 (14/16)	32 (8/25)
CD4 Mean Change from Bas	seline, cells/mm ³	
All patients	293 (n=14 ^b)	229 (n=14 ^b)
HIV RNA <50 copies/mL by	select baseline PI substitutions, 6 % (responder/evaluable ^d)
0-2	NA	27 (4/15)
3	NA	-
≥4	NA	0 (0/3)

^a Intent-to-treat analysis, with missing values considered as failures.

5.2 Pharmacokinetic properties

The pharmacokinetics of atazanavir were evaluated in healthy adult volunteers and in HIV-infected patients; significant differences were observed between the two groups. The pharmacokinetics of atazanavir exhibit a non-linear disposition.

Absorption: in HIV-infected patients (n=33, combined studies), multiple dosing of atazanavir 300 mg once daily with ritonavir 100 mg once daily with food produced a geometric mean (CV%) for atazanavir, C_{max} of 4466 (42%) ng/mL, with time to C_{max} of approximately 2.5 hours. The geometric mean (CV%) for atazanavir C_{min} and AUC was 654 (76%) ng/ml and 44185 (51%) ng•h/mL, respectively. In HIV-infected patients (n=13), multiple dosing of atazanavir 400 mg (without ritonavir) once daily with food produced a geometric mean (CV%) for atazanavir C_{max} of 2298 (71) ng/mL, with time to C_{max} of approximately 2.0 hours. The geometric mean (CV%) for atazanavir C_{min} and AUC were 120 (109) ng/ml and 14874 (91) ng•h/mL, respectively.

<u>Food effect:</u> co-administration of atazanavir and ritonavir with food optimises the bioavailability of atazanavir. Co-administration of a single 300-mg dose of atazanavir and 100-mg dose of ritonavir with a light meal resulted in a 33% increase in the AUC and a 40% increase in both the C_{max} and the 24 hour concentration of atazanavir relative to the fasting state. Co-administration with a high-fat meal did not affect the AUC of atazanavir relative to fasting conditions and the C_{max} was within 11% of fasting values. The 24-hour concentration following a high-fat meal was increased by approximately 33% due to delayed absorption; the median T_{max} increased from 2.0 to 5.0 hours. Administration of atazanavir with ritonavir with either a light or a high-fat meal decreased the coefficient of variation of AUC and C_{max} by approximately 25% compared to the fasting state. To enhance bioavailability and minimise variability, atazanavir is to be taken with food.

b Number of patients evaluable.

^c PI major L24I, D30N, V32I, L33F, M46IL, I47AV, G48V, I50LV, F53LY, I54ALMSTV, L76V, V82AFLST, I84V, N88DS, L90M; PI minor: L10CFIRV, V11I, E35G, K43T, Q58E, A71ILTV, G73ACST, T74P, N83D, L89V.

d Includes patients with baseline resistance data.
 NA = not applicable.

Distribution: atazanavir was approximately 86% bound to human serum proteins over a concentration range of 100 to 10,000 ng/mL. Atazanavir binds to both alpha-1-acid glycoprotein (AAG) and albumin to a similar extent (89% and 86%, respectively, at 1,000 ng/mL). In a multiple-dose study in HIV- infected patients dosed with 400-mg of atazanavir once daily with a light meal for 12 weeks, atazanavir was detected in the cerebrospinal fluid and semen.

Metabolism: studies in humans and in vitro studies using human liver microsomes have demonstrated that atazanavir is principally metabolised by CYP3A4 isozyme to oxygenated metabolites. Metabolites are then excreted in the bile as either free or glucuronidated metabolites. Additional minor metabolic pathways consist of N-dealkylation and hydrolysis. Two minor metabolites of atazanavir in plasma have been characterised. Neither metabolite demonstrated in vitro antiviral activity.

Elimination: following a single 400 mg dose of ¹⁴C-atazanavir, 79% and 13% of the total radioactivity was recovered in the faeces and urine, respectively. Unchanged drug accounted for approximately 20% and 7% of the administered dose in the faeces and urine, respectively. Mean urinary excretion of unchanged drug was 7% following 2 weeks of dosing at 800 mg once daily. In HIV-infected adult patients (n=33, combined studies) the mean half-life within a dosing interval for atazanavir was 12 hours at steady state following a dose of 300 mg daily with ritonavir 100 mg once daily with a light meal.

Special populations

Renal impairment: in healthy subjects, the renal elimination of unchanged atazanavir was approximately 7% of the administered dose. There are no pharmacokinetic data available for atazanavir with ritonavir in patients with renal insufficiency. atazanavir (without ritonavir) has been studied in adult patients with severe renal impairment (n=20), including those on haemodialysis, at multiple doses of 400 mg once daily. Although this study presented some limitations (i.e., unbound drug concentrations not studied), results suggested that the atazanavir pharmacokinetic parameters were decreased by 30% to 50% in patients undergoing haemodialysis compared to patients with normal renal function. The mechanism of this decrease is unknown. (See sections 4.2 and 4.4.)

Hepatic impairment: atazanavir is metabolised and eliminated primarily by the liver. atazanavir (without ritonavir) has been studied in adult subjects with moderate-to-severe hepatic impairment (14 Child-Pugh Class B and 2 Child-Pugh Class C subjects) after a single 400-mg dose. The mean AUC $_{(0-\infty)}$ was 42% greater in subjects with impaired hepatic function than in healthy subjects. The mean half-life of atazanavir in hepatically impaired subjects was 12.1 hours compared to 6.4 hours in healthy subjects. The effects of hepatic impairment on the pharmacokinetics of atazanavir after a 300 mg dose with ritonavir have not been studied. Concentrations of atazanavir with or without ritonavir are expected to be increased in patients with moderately or severely impaired hepatic function (see sections 4.2, 4.3, and 4.4).

Age/Gender: a study of the pharmacokinetics of atazanavir was performed in 59 healthy male and female subjects (29 young, 30 elderly). There were no clinically important pharmacokinetic differences based on age or gender.

Race: a population pharmacokinetic analysis of samples from Phase II clinical trials indicated no effect of race on the pharmacokinetics of atazanavir.

Pregnancy: The pharmacokinetic data from HIV-infected pregnant women receiving atazanavir capsules with ritonavir are presented in Table 8.

Table 8: Steady-State Pharmacokinetics of Atazanavir with ritonavir in HIV-Infected Pregnant Women in the Fed State

	atazanavir 300 mg with ritonavir 100 mg			
Pharmacokinetic Parameter	2nd Trimester	3rd Trimester	postpartum ^a	
	(n=9)	(n=20)	(n=36)	
C _{max} ng/mL	3729.09	3291.46	5649.10	
Geometric mean (CV%)	(39)	(48)	(31)	
AUC ng•h/mL	34399.1	34251.5	60532.7	
Geometric mean (CV%)	(37)	(43)	(33)	
C _{min} ng/mL ^b Geometric mean (CV%)	663.78	668.48	1420.64	
	(36)	(50)	(47)	

^a Atazanavir peak concentrations and AUCs were found to be approximately 26-40% higher during the postpartum period (4-12 weeks) than those observed historically in HIV-infected, non-pregnant patients. Atazanavir plasma trough concentrations were approximately 2-fold higher during the postpartum period when compared to those observed historically in HIV-infected non-pregnant patients.

Paediatric population

There is a trend toward a higher clearance in younger children when normalised for body weight. As a result, greater peak to trough ratios are observed, however at recommended doses, geometric mean atazanavir exposures (C_{min} , C_{max} and AUC) in paediatric patients are expected to be similar to those observed in adults.

5.3 Preclinical safety data

In repeat-dose toxicity studies, conducted in mice, rats, and dogs, atazanavir-related findings were generally confined to the liver and included generally minimal to mild increases in serum bilirubin and liver enzymes, hepatocellular vacuolation and hypertrophy, and, in female mice only, hepatic single- cell necrosis. Systemic exposures of atazanavir in mice (males), rats, and dogs at doses associated with hepatic changes were at least equal to that observed in humans given 400 mg once daily. In female mice, atazanavir exposure at a dose that produced single-cell necrosis was 12 times the exposure in humans given 400 mg once daily. Serum cholesterol and glucose were minimally to mildly increased in rats but not in mice or dogs.

During *in vitro* studies, cloned human cardiac potassium channel (hERG), was inhibited by 15% at a concentration (30 µM) of atazanavir corresponding to 30-fold the free drug concentration at Cmax in humans. Similar concentrations of atazanavir increased by 13% the action potential duration (APD₉₀) in rabbit Purkinje fibres study. Electrocardiographic changes (sinus bradycardia, prolongation of PR interval, prolongation of QT interval, and prolongation of QRS complex) were observed only in an initial 2-week oral toxicity study performed in dogs. Subsequent 9-month oral toxicity studies in dogs showed no drug-related electrocardiographic changes. The clinical relevance of these non-clinical data is unknown. Potential cardiac effects of this product in humans cannot be ruled out (see sections 4.4 and 4.8). The potential for PR prolongation should be considered in cases of overdose (see section 4.9).

In a fertility and early embryonic development study in rats, atazanavir altered oestrus cycling with no effects on mating or fertility. No teratogenic effects were observed in rats or rabbits at maternally toxic doses. In pregnant rabbits, gross lesions of the stomach and intestines were observed in dead or moribund does at maternal doses 2 and 4 times the highest dose administered in the definitive embryo- development study. In the pre- and postnatal

^b C_{min} is concentration 24 hours post-dose.

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development assessment in rats, atazanavir produced a transient reduction in body weight in the offspring at a maternally toxic dose. Systemic exposure to atazanavir at doses that resulted in maternal toxicity was at least equal to or slightly greater than that observed in humans given 400 mg once daily.

Atazanavir was negative in an Ames reverse-mutation assay but did induce chromosomal aberrations *in vitro* in both the absence and presence of metabolic activation. In *in vivo* studies in rats, atazanavir did not induce micronuclei in bone marrow, DNA damage in duodenum (comet assay), or unscheduled DNA repair in liver at plasma and tissue concentrations exceeding those that were clastogenic *in vitro*.

In long-term carcinogenicity studies of atazanavir in mice and rats, an increased incidence of benign hepatic adenomas was seen in female mice only. The increased incidence of benign hepatic adenomas in female mice was likely secondary to cytotoxic liver changes manifested by single-cell necrosis and is considered to have no relevance for humans at intended therapeutic exposures. There were no tumorigenic findings in male mice or in rats.

Atazanavir increased opacity of bovine corneas in an *in vitro* ocular irritation study, indicating it may be an ocular irritant upon direct contact with the eye.

6. PHARMACEUTICAL PARTICULARS

6.1 List of excipients

100 mg, 150 mg, 200 mg hard capsules Capsule content:
Lactose monohydrate
Crospovidone (type A) (E 1202)
Silica, colloidal anhydrous (E 551)
Magnesium stearate (E 470b)

Capsule shell:

Gelatin

Titanium dioxide (E 171)

Indigotine (E 132) (contains sodium)

Printing ink, white:

Shellac

Titanium dioxide (E 171)

Propylene glycol (E 1520)

300 mg hard capsules

Capsule content:

Lactose monohydrate

Crospovidone (type A) (E 1202)

Silica, colloidal anhydrous (E 551)

Magnesium stearate (E 470b)

Capsule shell:

Gelatin

Titanium dioxide (E 171)

Indigotine (E 132) (contains sodium)

Red iron oxide (E 172)

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Printing ink, white:

Shellac

Titanium dioxide (E 171)

Propylene glycol (E 1520)

6.2 **Incompatibilities**

Not applicable.

6.3 Shelf life

36 months

Shelf life after first opening:

Bottles:

[NL/H/4117/001-002-003, NL/H/4118/001-002]

4 months

[NL/H/4117/004, NL/H/4118/003]

2 months

6.4 Special precautions for storage

Do not store above 30°C.

6.5 Nature and contents of container

[NL/H/4117/001-002-003-004]

The hard capsules are packed in Aluminium-OPA/Alu/PVC unit dose perforated blisters, Aluminium-OPA/Alu/PVC blisters or high-density polyethylene (HDPE) bottles closed with child-resistant polypropylene closure.

[NL/H/4118/001-002-003]

The hard capsules are packed in Aluminium-OPA/Alu/PVC unit dose perforated blisters or high-density polyethylene (HDPE) bottles closed with child-resistant polypropylene closure.

Pack sizes:

[NL/H/4117/001-002-003]

Unit dose blister: 60 x 1 hard capsules; 10 blister cards of 6 x 1 hard capsules each

Blister: 60 hard capsules; 10 blister cards of 6 hard capsules each

Bottle: 60 hard capsules

[NL/H/4117/004]

Unit dose blister:

30 x 1 hard capsules; 5 blister cards of 6 x 1 hard capsules each multipack containing 60 x 1 (2 packs of 30 x 1) hard capsules multipack containing 90 x 1 (3 packs of 30 x 1) hard capsules multipack containing 120 x 1 (4 packs of 30 x 1) hard capsules Blister:

30 hard capsules; 5 blister cards of 6 hard capsules each multipack containing 60 (2 packs of 30) hard capsules multipack containing 90 (3 packs of 30) hard capsules multipack containing 120 (4 packs of 30) hard capsules Bottles:

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30 hard capsules

multipack containing 60 (2 packs of 30) hard capsules multipack containing 90 (3 packs of 30) hard capsules multipack containing 120 (4 packs of 30) hard capsules

[NL/H/4118/001-002]

Unit dose blister: 60 x 1 hard capsules; 10 blister cards of 6 x 1 hard capsules each.

Bottle: 60 hard capsules

[NL/H/4118/003]

Unit dose blister:

30 x 1 hard capsules; 5 blister cards of 6 x 1 hard capsules each multipack containing 90 x 1 (3 packs of 30 x 1) hard capsules

Bottle:

30 hard capsules

multipack containing 90 (3 packs of 30) hard capsules

Not all pack sizes may be marketed.

6.6 Special precautions for disposal

Any unused medicinal product or waste material should be disposed of in accordance with local requirements.

7. HOUDER VAN DE VERGUNNING VOOR HET IN DE HANDEL BRENGEN

Sandoz B.V. Hospitaaldreef 29 1315 RC Almere Nederland

NUMMER(S) VAN DE VERGUNNING VOOR HET IN DE HANDEL 8. BRENGEN

150 mg: RVG 121568 200 mg: RVG 121569 300 mg: RVG 121570

9. DATUM VAN EERSTE VERLENING VAN DE VERGUNNING/VERLENGING VAN DE VERGUNNING

Datum van eerste verlening van de vergunning: 12 november 2018 Datum laatste verlenging: 19 september 2023

10. DATUM VAN HERZIENING VAN DE TEKST

Laatste gedeeltelijke wijziging betreft de rubrieken 4.3 & 4.5: 18 december 2024