September 2022

SUMMARY OF PRODUCT CHARACTERISTICS

1. NAME OF THE MEDICINAL PRODUCT

Ritonavir Tablets USP 100 mg

2. QUALITATIVE AND QUANTITATIVE COMPOSITION

Each tablet contains 100 mg ritonavir.

For the list of excipients, see section 6-1.

3. PHARMACEUTICAL FORM

Tablets.

White to off-white, capsule-shaped, film-coated tablets, debossed with 'H' on one side and 'R9' on the other side.

4. CLINICAL PARTICULARS

4.1 Therapeutic indications

Ritonavir Tablets USP 100 mg is indicated as a pharmacokinetic enhancer for protease inhibitors when these are used in combination therapy with other antiretroviral agents for the treatment of HIV-1 infected patients.

Consideration should be given to official treatment guidelines for HIV-1 infection (e.g. those of the WHO).

This product is intended for use in children. Nonetheless, safety information is provided with respect to adult health issues such as liver disease, pregnancy and breastfeeding, to allow full access to all relevant information.

4.2 Posology and method of administration

Therapy should be initiated by a health care provider experienced in the management of HIV infection.

Posology

Ritonavir Tablets USP 100 mg should be taken with food. The tablets should be swallowed whole and not be chewed, broken or crushed.

As Ritonavir Tablets USP 100 mg is used as a pharmacokinetic enhancer with other protease inhibitors, the productinformation for the particular protease inhibitor must be consulted.

The following HIV-1 protease inhibitors can be used with ritonavir as a pharmacokinetic enhancer at the noted doses.

Adults and adolescents:

- Amprenavir 600 mg twice daily with ritonavir 100 mg twice daily
- Atazanavir 300 mg once daily with ritonavir 100 mg once daily
- Fosamprenavir 700 mg twice daily with ritonavir 100 mg twice daily
- Lopinavir co-formulated with ritonavir (lopinavir/ritonavir) 400 mg/100 mg or 800 mg/200 mg
- Saquinavir 1000 mg twice daily with ritonavir 100 mg twice daily in antiretroviral treatment (ART)experienced patients. Initiate treatment with saquinavir 500 mg twice daily with ritonavir 100 mg twice
 daily for the first 7 days, then saquinavir 1000 mg twice daily with ritonavir 100 mg twice daily in ARTnaïve patients.
- Tipranavir 500 mg twice daily with ritonavir 200 mg twice daily. Tipranavir with ritonavir should not be used in treatment-naïve patients.
- Darunavir 600 mg twice daily with ritonavir 100 mg twice daily in ART-experienced patients.
- Darunavir 800 mg once daily with ritonavir 100 mg once daily may be used in some ART-experienced patients. Refer to the darunavir product information for further information on once daily dosing in ART-experienced patients.
- Darunavir 800 mg once daily with ritonavir 100 mg once daily in ART-naïve patients.

[†] Trade names are not prequalified by WHO. This is the national medicines regulatory agency's responsibility.

Paediatric patients weighing 25 kg or more:

The recommended dose is 100 mg ritonavir once or two times per day, depending on the concurrently used protease inhibitor.

For children who are undergoing anti-tuberculosis treatment with rifampicin, higher dosages of ritonavir may be needed for pharmacokinetic enhancement of the combined protease inhibitor. Please refer to the product information of the protease inhibitors approved for co-administration with ritonavir.

Special populations

Renal impairment:

Depending on the specific protease inhibitor with which it is co-administered, ritonavir may be appropriate for use with caution in patients with renal insufficiency. For specific dosing information in patients with renal impairment, refer to the product information of the co-administered protease inhibitor.

Hepatic impairment:

Ritonavir should not be given to patients with decompensated liver disease, (see section 4.3). In the absence of pharmacokinetic studies in patients with stable severe hepatic impairment (Child Pugh grade C) without decompensation, caution should be exercised when ritonavir is used as a pharmacokinetic enhancer as increased levels of the co-administered protease inhibitor may occur. Specific recommendations for use of ritonavir as a pharmacokinetic enhancer in patients with hepatic impairment are dependent on the protease inhibitor with which it is co-administered. The product information of the co-administered protease inhibitor should be reviewed for specific dosing information in this patient population.

Paediatric population:

Ritonavir Tablets USP 100 mg should not be used in children weighing less than 25 kg. For these patients, more suitable formulations containing a lower amount of the active substance may be available.

4.3 Contraindications

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

Consult the product information of the co-administered medicine for other possible contraindications.

Ritonavir should not be given to patients with decompensated liver disease.

In vitro and *in vivo* studies have demonstrated that ritonavir is a potent inhibitor of CYP3A- and CYP2D6mediated biotransformations. The following medicines are contraindicated when used with ritonavir and, unless otherwise noted, the contraindication is based on the potential for ritonavir to inhibit metabolism of the co-administered drug, resulting in increased exposure to the co-administered drug and risk of clinically significant adverse effects.

The enzyme-modulating effect of ritonavir may be dose dependent.

Drug class	Drugs within class	Rationale
Concomitant drug leve	els increased or decreased	
α1-Adrenoreceptor Antagonist	alfuzosin	Increased plasma concentrations of alfuzosin which may lead to severe hypotension (see section 4.5).
Analgesics	pethidine, piroxicam, propoxyphene	Increased plasma concentrations of norpethidine, piroxicam and propoxyphene. Thereby, increasing the risk of serious respiratory depression or haematologic abnormalities, or other serious adverse effects from these agents.
Antianginal	ranolazine	Increased plasma concentrations of ranolazine which may increase the potential for serious and/or life-threatening reactions (see section 4.5).
Anticancer	neratinib	Increased plasma concentrations of neratinib which may increase the potential for serious and/or life-threatening reactions including hepatotoxicity (see section 4.5).
	venetoclax	Increased plasma concentrations of venetoclax. Increased risk of tumor lysis syndrome at the dose initiation and during the dose-titration phase (see section 4.5).
Antiarrhythmics	amiodarone, bepridil, dronedarone, encainide, flecainide, propafenone, quinidine	Increased plasma concentrations of amiodarone, bepridil, dronedarone, encainide, flecainide, propafenone, quinidine. Thereby, increasing the risk of arrhythmias or other serious adverse effects from these agents.
Antibiotic	fusidic acid	Increased plasma concentrations of fusidic acid and ritonavir.
Antihistamines	astemizole, terfenadine	Increased plasma concentrations of astemizole and terfenadine. Thereby, increasing the risk of serious arrhythmias from these agents.
Anti-gout	colchicine	Potential for serious and/or life-threatening reactions in patients with renal and/or hepatic impairment (see sections 4.4 and 4.5).
Antipsychotics/ Neuroleptics	lurasidone	Increased plasma concentrations of lurasidone which may increase the potential for serious and/or life-threatening reactions (see section 4.5).
	clozapine, pimozide	Increased plasma concentrations of clozapine and pimozide. Thereby, increasing the risk of serious haematologic abnormalities, or other serious adverse effects from these agents.
	quetiapine	Increased plasma concentrations of quetiapine which may lead to coma. The concomitant administration with quetiapine is contraindicated (see section 4.5).
Ergot Derivatives	dihydroergotamine, ergonovine, ergotamine, methylergonovine	Increased plasma concentrations of ergot derivatives leading to acute ergot toxicity, including vasospasm and ischaemia.
GI motility agent	cisapride	Increased plasma concentrations of cisapride. Thereby, increasing the risk of serious arrhythmias from this agent.

Drug class	Drugs within class	Rationale
Concomitant drug levels in	ncreased or decreased	
Lipid-modifying agents		
HMG Co-A Reductase Inhibitors	lovastatin, simvastatin	Increased plasma concentrations of lovastatin and simvastatin; thereby, increasing the risk of myopathy including rhabdomyolysis (see section 4.5).
Microsomal triglyceride transfer protein (MTTP) inhibitor	lomitapide	Increased plasma concentrations of lomitapide (see section 4.5).
PDE5 inhibitor	avanafil	Increased plasma concentrations of avanafil (see section 4.4. and 4.5).
	sildenafil	Contraindicated when used for the treatment of pulmonary arterial hypertension (PAH) only. Increased plasma concentrations of sildenafil. Thereby, increasing the potential for sildenafil-associated adverse events (which include hypotension and syncope). See section 4.4 and section 4.5 for co-administration of sildenafil in patients with erectile dysfunction.
	vardenafil	Increased plasma concentrations of vardenafil (see section 4.4. and 4.5).
Sedatives/hypnotics	clorazepate, estazolam, flurazepam, oral midazolam and triazolam	Increased plasma concentrations of clorazepate, estazolam, flurazepam, oral midazolam and triazolam. Thereby, increasing the risk of extreme sedation and respiratory depression from these agents. (For caution on parenterally administered midazolam, see section 4.5).
Ritonavir level decreased		
Herbal preparation	St. John's wort	Herbal preparations containing St John's wort (Hypericum perforatum) due to the risk of decreased plasma concentrations and reduced clinical effects of ritonavir (see section 4.5).

4.4 Special warnings and precautions for use

Patients receiving ritonavir or any other antiretroviral therapy may still develop opportunistic infections and other complications of HIV infection. Therefore patients should remain under close clinical observation by health care providers experienced in the treatment of these associated HIV diseases.

While effective viral suppression with antiretroviral therapy has been proven to substantially reduce the risk of sexual transmission, a residual risk cannot be excluded. Precautions to prevent transmission should be taken in accordance with national guidelines.

As ritonavir is used as a pharmacokinetic enhancer with other protease inhibitors, full details on the warnings and precautions relevant to that particular protease inhibitor should be considered.

Some of the below warnings originate from the use of ritonavir as antiretroviral agent at higher doses than those recommended for pharmacokinetic enhancement. The effects of ritonavir when used as a pharmacokinetic enhancer might hence be less pronounced.

Patients with chronic diarrhoea or malabsorption

Extra monitoring is recommended when diarrhoea occurs. The relatively high frequency of diarrhoea during treatment with ritonavir may compromise the absorption and efficacy (due to decreased compliance) of ritonavir or other concurrent medicinal products. Serious persistent vomiting and/or diarrhoea associated

with ritonavir use might also compromise renal function. It is advisable to monitor renal function in patients with renal function impairment.

Patients with haemophilia

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

Weight, blood lipids and glucose

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

Pancreatitis

Pancreatitis should be considered if clinical symptoms (nausea, vomiting, abdominal pain) or abnormalities in laboratory values (such as increased serum lipase or amylase values) suggestive of pancreatitis should occur. Patients who exhibit these signs or symptoms should be evaluated and ritonavir therapy should be discontinued if a diagnosis of pancreatitis is made (see section 4.8).

Immune reconstitution inflammatory syndrome

When starting combination antiretroviral therapy (CART) in patients with severe immune deficiency, an inflammatory reaction to asymptomatic or residual opportunistic pathogens may arise and cause serious clinical conditions, or aggravate symptoms. Typically, such reactions occur within the first weeks or months of starting CART. Relevant examples are cytomegalovirus retinitis, generalised or focal mycobacterial infections and pneumonia caused by *Pneumocystis jirovecii* (formerly known as *Pneumocystis carinii*). Any inflammatory symptoms should be evaluated and treated when necessary.

Autoimmune disorders (such as Graves' disease and autoimmune hepatitis) have also been reported in the setting of immune reactivation; however, the time to onset is more variable and can occur many months after starting treatment.

Liver disease

Ritonavir should not be given to patients with decompensated liver disease. For patients with stable severe hepatic impairment (Child Pugh grade C) without decompensation see section 4.2. 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 to the relevant product information for these medicines.

Patients with 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 disease

Since the renal clearance of ritonavir is negligible, decrease in the total body clearance is not expected in patients with renal impairment. For specific dosing information in patients with renal impairment, refer to the product information of the co-administered protease inhibitor. See also section 4.2.

Renal failure, renal impairment, elevated creatinine, hypophosphataemia and proximal tubulopathy (including Fanconi syndrome) have been reported with concomitant use of tenofovir disoproxil fumarate in clinical practice (see section 4.8).

Osteonecrosis

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

PR interval prolongation

Ritonavir has been shown to cause modest asymptomatic prolongation of the PR interval in some healthy adult subjects. Rare reports of 2nd or 3rd degree atrioventricular block in patients with underlying structural heart disease and pre-existing conduction system abnormalities or in patients receiving medicinal products known to prolong the PR interval (such as verapamil or atazanavir) have been reported in patients receiving ritonavir. Ritonavir should be used with caution in such patients (see section 5.1).

Interactions with other medicinal products

Full details on the warnings and precautions relevant to the protease inhibitor ritonavir is used with must be considered, therefore section 4.4 of the product information for the particular protease inhibitor must be consulted to determine if the information below is applicable. Furthermore, some information may only apply to ritonavir used as an antiretroviral agent.

PDE5 inhibitors: Particular caution should be used when prescribing sildenafil or tadalafil for the treatment of erectile dysfunction in patients receiving ritonavir. Co-administration of ritonavir with these medicinal products is expected to substantially increase their concentrations and may result in associated adverse reactions such as hypotension and prolonged erection (see section 4.5).

Concomitant use of avanafil or vardenafil with ritonavir is contraindicated. Concomitant use of sildenafil with ritonavir is contraindicated in pulmonary arterial hypertension patients (see section 4.3).

HMG-CoA reductase inhibitors: The HMG-CoA reductase inhibitors simvastatin and lovastatin are highly dependent on CYP3A for metabolism, thus concomitant use of ritonavir with simvastatin or lovastatin is not recommended due to an increased risk of myopathy including rhabdomyolysis. Caution must also be exercised and reduced doses should be considered if ritonavir is used concurrently with atorvastatin, which is metabolised to a lesser extent by CYP3A. While rosuvastatin elimination is not dependent on CYP3A, an elevation of rosuvastatin exposure has been reported with ritonavir co-administration. The mechanism of this interaction is not clear, but may be the result of transporter inhibition. When used with ritonavir, the lowest doses of atorvastatin or rosuvastatin should be administered. The metabolism of pravastatin and fluvastatin is not dependent of CYP3A, and interactions are not expected with ritonavir. If treatment with an HMG-CoA reductase inhibitor is indicated, pravastatin or fluvastatin is recommended (see section 4.5).

Colchicine: Life-threatening and fatal drug interactions have been reported in patients treated with colchicine and strong inhibitors of CYP3A like ritonavir (see sections 4.3 and 4.5).

Digoxin: Particular caution should be used when prescribing ritonavir in patients taking digoxin since coadministration of ritonavir with digoxin is expected to increase digoxin levels. The increased digoxin levels may lessen over time (see section 4.5).

In patients who are already taking digoxin when ritonavir is introduced, the digoxin dose should be reduced to one-half of the patients' normal dose and patients need to be followed more closely than usual for several weeks after initiating co-administration of ritonavir and digoxin.

In patients who are already taking ritonavir when digoxin is introduced, digoxin should be introduced more gradually than usual. Digoxin levels should be monitored more intensively than usual during this period, with dose adjustments made, as necessary, based on clinical, electrocardiographic and digoxin level findings.

Ethinyl estradiol: Barrier or other non-hormonal methods of contraception should be considered when administering ritonavir at therapeutic or low doses as ritonavir is likely to reduce the effect and change the uterine bleeding profile when co-administered with estradiol-containing contraceptives.

Glucocorticoids: Concomitant use of 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).

Trazodone: Particular caution should be used when prescribing ritonavir in patients using trazodone. Trazodone is a CYP3A4 substrate and co-administration of ritonavir is expected to increase trazodone levels. Adverse reactions of nausea, dizziness, hypotension and syncope have been observed in single dose interaction studies in healthy volunteers (see section 4.5).

Rivaroxaban: It is not recommended to use ritonavir in patients receiving rivaroxaban, due to the risk of increased bleeding (see section 4.5).

Riociguat: The concomitant use of ritonavir is not recommended due to potential increase in riociguat exposure (see section 4.5).

Vorapaxar: The concomitant use of ritonavir is not recommended due to potential increase in vorapaxar exposure (see section 4.5).

Bedaquiline: Strong CYP3A4 inhibitors such as protease inhibitors may increase bedaquiline exposure which could potentially increase the risk of bedaquiline-related adverse reactions. Therefore, combination of bedaquiline with ritonavir should be avoided. However, if the benefit outweighs the risk, co-administration of bedaquiline with ritonavir must be done with caution. More frequent electrocardiogram monitoring and monitoring of transaminases is recommended (see section 4.5 and refer to the bedaquiline product information).

Delamanid: Co-administration of delamanid with a strong inhibitor of CYP3A (ritonavir) may increase exposure to delamanid metabolite, which has been associated with QTc prolongation. Therefore, if co-administration of delamanid with ritonavir is considered necessary, very frequent ECG monitoring throughout the full delamanid treatment period is recommended (see section 4.5 and refer to the delamanid product information).

Saquinavir: Doses of ritonavir higher than 100 mg twice daily should not be used. Higher doses of ritonavir have been shown to be associated with an increased incidence of adverse reactions.

Co-administration of saquinavir and ritonavir has led to severe adverse reactions, mainly diabetic ketoacidosis and liver disorders, especially in patients with pre-existing liver disease.

Saquinavir/ritonavir should not be given together with rifampicin, due to the risk of severe hepatotoxicity (presenting as increased hepatic transaminases) if the three medicines are given together (see section 4.5).

Tipranavir: Co-administration of tipranivir with 200 mg of ritonavir has been associated with reports of clinical hepatitis and hepatic decompensation including some fatalities. Extra vigilance is warranted in patients with chronic hepatitis B or hepatitis C co-infection, as these patients have an increased risk of hepatotoxicity.

Doses of ritonavir lower than 200 mg twice daily should not be used as they might alter the efficacy profile of the combination.

Fosamprenavir: Co-administration of fosamprenavir with ritonavir in doses greater than 100 mg twice daily has not been clinically evaluated. The use of higher ritonavir doses might alter the safety profile of the combination and therefore is not recommended.

Atazanavir: 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. Refer to the atazanavir product information for further details.

4.5 Interaction with other medicinal products and other forms of interaction

Ritonavir has a high affinity for several cytochrome P450 (CYP) isoforms and may inhibit oxidation with the following ranked order: CYP3A4 > CYP2D6. Co-administration of ritonavir and medicinal products primarily metabolised by CYP3A may result in increased plasma concentrations of the other medicinal product, which could increase or prolong its therapeutic and adverse effects. For select medicinal products (e.g. alprazolam) the inhibitory effects of ritonavir on CYP3A4 may decrease over time. Ritonavir also has a high affinity for P-glycoprotein and may inhibit this transporter. The inhibitory effect of ritonavir (with or without other protease inhibitors) on P-gp activity may decrease over time (e.g. digoxin and fexofenadine - see table "Ritonavir effects on non-antiretroviral medicinal products" below). Ritonavir may induce glucuronidation and oxidation by CYP1A2, CYP2C8, CYP2C9 and CYP2C19 thereby increasing the biotransformation of some medicinal products, which could decease or shorten their therapeutic effect. Important information regarding medicinal product interactions when ritonavir is used as a pharmacokinetic enhancer is also contained in the summary of product characteristics of the co-administered protease inhibitor.

Medicinal products that affect ritonavir levels

Serum levels of ritonavir can be reduced by concomitant use of herbal preparations containing St John's wort (*Hypericum perforatum*). This is due to the induction of medicinal product metabolising enzymes by St John's wort. Herbal preparations containing St John's wort must not be used in combination with ritonavir. If a patient is already taking St John's wort, stop St John's wort and if possible check viral levels. Ritonavir levels may increase on stopping St John's wort. The dose of ritonavir may need adjusting. The inducing effect may persist for at least 2 weeks after cessation of treatment with St John's wort (see section 4.3).

Serum levels of ritonavir may be affected by certain co-administered medicinal products (eg delavirdine, efavirenz, phenytoin and rifampicin). These interactions are noted in the medicinal product interaction tables below.

Interaction table

Interactions between ritonavir and protease inhibitors, antiretroviral agents other than protease inhibitors and other non-antiretroviral medicinal products are listed in the tables below. This list is not intended to be inclusive or comprehensive. The product information of the medicines used concomitantly with ritonavir should be consulted.

Co-administered drug	Dose Co- administered drug (mg)	Ritonavir dose (mg)	Drug assessed	AUC	Cmin	
Amprenavir	600 q12h	100 q12h	Amprenavir ¹	↑64%	↑5 fold	
	Ritonavir increases the serum levels of amprenavir as a result of CYP3A4 inhibition. Clinical trials confirmed the safety and efficacy of 600 mg amprenavir twice daily with ritonavir 100 mg twice daily. For further information, physicians should refer to the amprenavir product information.					
Atazanavir	300 q24h	100 q24h	Atazanavir	↑86%	↑11 fold	
			Atazanavir ²	↑2 fold	↑3-7 fold	
	Ritonavir increases the serum levels of atazanavir as a result of CYP3A4 inhibition. Clinical trials confirmed the safety and efficacy of 300 mg atazanavir once daily with ritonavir 100 mg once daily in treatment experienced patients. For further information physicians should refer to the product information for atazanavir products.					
Darunavir	600, single	100 q12h	Darunavir	↑ 14 fold		
			evels of darunavir as a re itonavir to ensure its the			

Medicinal Product Interactions – Ritonavir with Protease Inhibitors

			have not been studied we the information for daruna		-
Fosamprenavir	700 q12h	100 q12h	Amprenavir	↑ 2.4 fold	\uparrow 11 fold
	CYP3A4 inhi effect. Clinica with ritonavir been studied	bition. Fosamprer al trials confirmed 100 mg twice dat	evels of amprenavir (fro navir must be given with the safety and efficacy o ily. Ritonavir doses high ir. For further informatio tion.	ritonavir to ensure it of fosamprenavir 700 er than 100 mg twice	ts therapeutic mg twice daily daily have not
Indinavir	800 q12h	100 q12h	Indinavir ³	↑ 178%	ND
			Ritonavir	↑ 72%	ND
	400 q12h	400 q12h	Indinavir ³	\leftrightarrow	↑4 fold
			Ritonavir	\leftrightarrow	\leftrightarrow
	achieved with ritonavir (100	doses higher that	ritonavir-mediated phar n 100 mg twice daily. In and indinavir (800 mg tw be increased. Saquinavir ⁴	cases of co-administ	ration of
Saquinavir	1000 q1211	100 q120	-	· · · · · · · · · · · · · · · · · · ·	
	400 101	400 101	Ritonavir	↔ • 17 C 11	↔
	400 q12h	400 q12h	Saquinavir ⁴ Ritonavir	$\uparrow 17 \text{ fold}$	ND ↔
	Saquinavir sh daily with sac hours similar without ritona In a clinical s saquinavir 10 hepatocellula normal after	ould only be give quinavir 1000 mg to or greater than avir. tudy investigating 00 mg with ritona r toxicity with tran 1 to 5 days of co-a	evels of saquinavir as a n n in combination with ri twice daily provides saq those achieved with saq the interaction of rifam vir 100 mg twice daily i nsaminase elevations up idministration was noted	tonavir. Ritonavir 10 uinavir systemic expe uinavir 1200 mg three picin 600 mg once da n healthy volunteers, to > 20-fold the uppe . Due to the risk of se	0 mg twice osure over 24 ee times daily aily and severe er limit of evere
		-	vir should not be given to ians should refer to the s		
Fipranavir	500 q12h	200 q12h	Tipranavir	↑ 11 fold	↑ 29 fold
		200 4120	Ritonavir	↓ 40%	ND
	Tipranavir m ritonavir less the efficacy o	ust be given with 1 than 200 mg twice	evels of tipranavir as a re low dose ritonavir to ens e daily should not be use . For further information	esult of CYP3A inhib sure its therapeutic ef ed with tipranavir as t	pition. fect. Doses of hey might alter

ND: Not determined.

1. Based on cross-study comparison to 1200 mg amprenavir twice daily alone.

Based on cross-study comparison to 400 mg atazanavir once daily alone.
 Based on cross-study comparison to 800 mg indinavir three times daily alone.

4. Based on cross-study comparison to 600 mg saquinavir three times daily alone.

Co-administered drug	Dose Co- administered drug (mg)	Ritonavir dose (mg)	Drug assessed	AUC	Cmin		
Didanosine	200 q12h	600 q12h 2 h later	Didanosine	↓ 13%	\leftrightarrow		
			be taken with food and does separated by 2.5 h. Do				
Delavirdine	400 q8h	600 q12h	Delavirdine ¹	\leftrightarrow	\leftrightarrow		
			Ritonavir	↑ 50%	↑ 75%		
		y ritonavir. Whe	cal data, the pharmacokin en used in combination w				
Efavirenz	600 q24h	500 q12h	Efavirenz	↑ 21%			
			Ritonavir	17%			
	A higher frequency of adverse reactions (eg, dizziness, nausea, paraesthesia) and laboratory abnormalities (elevated liver enzymes) has been observed when efavirenz is co-administered with ritonavir dosed as an antiretroviral agent.						
Maraviroc	100 q12h	100 q12h	Maraviroc	↑ 161%	↑ 28%		
	Maraviroc may	be given with ri	evels of maraviroc as a re tonavir to increase the ma t information for maravir	araviroc exposure.			
Nevirapine	200 q12h	600 q12h	Nevirapine	\leftrightarrow	\leftrightarrow		
			Ritonavir	\leftrightarrow	\leftrightarrow		
			vith nevirapine does not l nevirapine or ritonavir.	ead to clinically re	levant changes		
Raltegravir	400 single	100 q12h	Raltegravir	↓ 16%	↓ 1%		
Raltegravir			nd ralteoravir results in a	minor reduction in	n raltagravir		
Raltegravir	Co-administrat levels.	ion of ritonavir a	ind fancegravit results in a		ii raitegravii		
Raltegravir		300 q6h	Zidovudine	↓ 25%	ND		

Medicinal product interactions - ritonavir with antiretroviral agents other than protease inhibitors

ND: Not determined

1. Based on parallel group comparison.

Ritonavir effects on non-antiretroviral co-administered medicinal products

Co-administered drug	Dose Co-administered drug (mg)	Ritonavir dose (mg)	AUC	Cmax
Alpha1-Adrenoreceptor	· Antagonist			
Alfuzosin		stration is likely to res	1	lasma concentrations of
Amphetamine Derivativ	/es			

ND: Not determined

Amphetamine	Ritonavir dosed as an antiretroviral agent is likely to inhibit CYP2D6 and as a result is expected to increase concentrations of amphetamine and its derivatives. Careful monitoring of therapeutic and adverse effects is recommended when these medicines are concomitantly administered with antiretroviral doses of ritonavir (see section 4.4).					
Analgesics						
Buprenorphine	16 q24h	100 q12h	↑ 57%	↑ 77%		
Norbuprenorphine			↑ 33%	↑ 108%		
Glucuronide metabolites			\leftrightarrow	\leftrightarrow		
	clinically significant patients. Adjustment necessary when the t another protease inhi	pharmacodynamic ch to the dose of bupren wo are dosed together bitor and buprenorphi	anges in a popul orphine or ritona r. When ritonavir ine, the product i	ive metabolite did not lead to ation of opioid tolerant wir may therefore not be is used in combination with nformation of the co- cific dosing information.		
Pethidine, piroxicam, propoxyphene	Ritonavir co-administration is likely to result in increased plasma concentrations of pethidine, piroxicam, and propoxyphene and is therefore contraindicated (see section 4.3).					
Fentanyl	Ritonavir dosed as a pharmacokinetic enhancer or as an antiretroviral agent inhibits CYP3A4 and as a result is expected to increase the plasma concentrations of fentanyl. Careful monitoring of therapeutic and adverse effects (including respiratory depression) is recommended when fentanyl is concomitantly administered with ritonavir.					
Methadone ¹	5, single dose	500 q12h,	↓ 36%	↓ 38%		
	ritonavir dosed as an induction of glucuror patient's clinical resp	antiretroviral agent o nidation. Dose adjustr onse to methadone th	r as a pharmacok nent should be co nerapy.	itantly administered with inetic enhancer due to onsidered based on the		
Morphine		be decreased due to r dosed as an antiretro		curonidation by co- a pharmacokinetic enhancer.		
Antianginal						
Ranolazine	Due to CYP3A inhibition by ritonavir, concentrations of ranolazine are expected to increase. The concomitant administration with ranolazine is contraindicated (see section 4.3).					
Antiarrthymics						
Amiodarone, bepridil, dronedarone, encainide, flecanide, propafenone, quinidine		, dronedarone, encain	ide, flecanide, pr	plasma concentrations of opafenone, and quinidine and		
Lidocaine	Coadministration may increase lidocaine exposure and a dose adjustment may be needed. The clinical effect should be monitored.					
Digoxin	0.5 single IV dose	300 q12h, 3 days	↑ 86%	ND		
	0.4 single oral dose	200 q12h, 13 days	↑ 22%	\leftrightarrow		
	by ritonavir dosed as	an antriretroviral age red in patients receiving	nt or as a pharma	ein mediated digoxin efflux acokinetic enhancer. Increased lessen over time as induction		
Antiasthmatic						
Theophylline ¹	3 mg/kg q8h	500 q12h	↓ 43%	↓ 32%		

	An increased dose of due to induction of C		quired when co	- administered with ritonavir,		
Anticancer agents						
Afatinib	20 mg, single dose	200 q12h/1h before	↑ 48%	↑ 39%		
	40 mg, single dose	200 q12h/ coadministered	↑ 19%	↑ 4%		
	40 mg, single dose	200 q12h/6h after	↑ 11%	↑ 5%		
	and acute P-gp inhib on the timing of ritor	ition by ritonavir. The e navir administration. Ca rir Tablets USP 100 mg	xtent of increa ution should be	the Resistance Protein (BCRP) see in AUC and C_{max} depends the exercised in administering fatinib product information).		
Abemaciclib	Serum concentrations may be increased due to CYP3A4 inhibition by ritonavir. Co-administration of abemaciclib and ritonavir should be avoided. If this co- administration is judged unavoidable, refer to the abemaciclib product information for dosage adjustment recommendations. Monitor for ADRs related to abemaciclib.					
Apalutamide	Apalutamide is a moderate to strong CYP3A4 inducer and this may lead to a decreased exposure of ritonavir and potential loss of virologic response. In addition, serum concentrations may be increased when co-administered with ritonavir resulting in the potential for serious adverse events including seizure.					
		itonavir with apalutami				
Ceritinib	Serum concentrations may be increased due to CYP3A and P-gp inhibition by ritonavir. Caution should be exercised in administering ceritinib with ritonavir. Refer to the ceritinib product information for dosage adjustment recommendations. Monitor for ADRs related to ceritinib.					
Dasatinib, nilotinib, vincristine, vinblastine	Serum concentrations may be increased when co-administered with ritonavir resulting in the potential for increased incidence of adverse reactions.					
Encorafenib	Serum concentrations may be increased when co-administered with ritonavir which may increase the risk of toxicity, including the risk of serious adverse events such as QT interval prolongation. Co-administration of encorafenib and ritonavir should be avoided. If the benefit is considered to outweigh the risk and ritonavir must be used, patients should be carefully monitored for safety.					
Fostamatinib	Co-administration of fostamatinib with ritonavir may increase fostamatinib metabolite R406 exposure resulting in dose-related adverse events such as hepatotoxicity, neutropenia, hypertension, or diarrhoea. Refer to the fostamatinib product information for dose reduction recommendations if such events occur.					
Ibrutinib	Serum concentrations of ibrutinib may be increased due to CYP3A inhibition by ritonavir, resulting in increased risk for toxicity including risk of tumour lysis syndrome. Co-administration of ibrutinib and ritonavir should be avoided. If the benefit is considered to outweigh the risk and ritonavir must be used, reduce the ibrutinib dose to 140 mg and monitor patient closely for toxicity.					
Neratinib	Serum concentrations may be increased due to CYP3A4 inhibition by ritonavir.					
		neratinib with ritonavir i reactions including hep		ted due to serious and/or life- e section 4.3).		
Venetoclax	in increased risk of t		t the dose initia	ibition by ritonavir, resulting ation and during the ramp-up nformation).		
	venetoclax, reduce th		least 75% whe	e on a steady daily dose of en used with strong CYP3A dosing instructions).		

Anticoagulants						
Rivaroxaban	10, single dose	600 q12h	↑ 153%	↑ 55%		
	effects of rivaroxab	an which may lead		s and pharmacodynamic ng risk. Therefore, the use aban.		
Vorapaxar	Serum concentrations may be increased due to CYP3A inhibition by ritonavir. The co- administration of vorapaxar with ritonavir is not recommended (see section 4.4 and refer to the vorapaxar product information).					
Warfarin	5, single dose	400 q12h				
S-Warfarin			↑9%	↓ 9%		
R-Warfarin			↓ 33%	\leftrightarrow		
	pharmacokinetic eff Decreased R-warfar recommended that a	fect is noted on S-v rin levels may lead anticoagulation par	ad to decreased levels of varfarin when co-admin to reduced anticoagula ameters are monitored n antiretroviral agent of	tion, therefore it is when warfarin is co-		
Anticonvulsants						
Carbamazepine	Ritonavir dosed as a pharmacokinetic enhancer or as an antiretroviral agent inhibits CYP3A4 and as a result is expected to increase the plasma concentrations of carbamazepine. Careful monitoring of therapeutic and adverse effects is recommended when carbamazepine is concomitantly administered with ritonavir.					
Divalproex, lamotrigine, phenytoin	Ritonavir dosed as a pharmacokinetic enhancer or as an antiretroviral agent induces oxidation by CYP2C9 and glucuronidation and as a result is expected to decrease the plasma concentrations of anticonvulsants. Careful monitoring of serum levels or therapeutic effects is recommended when these medicines are concomitantly administered with ritonavir. Phenytoin may decrease serum levels of ritonavir.					
Oxcarbamazepine		dose adjustment m	ure of the antiretroviral ay be needed. Monitor	drug, although to a clinical effect. Alternative		
Antidepressants						
Amitriptyline, fluoxetine, imipramine, nortriptyline, paroxetine, sertraline	expected to increase fluoxetine, paroxeti	e concentrations of ne or sertraline. Ca nen these medicine	imipramine, amitriptyl reful monitoring of the s are concomitantly adu	rapeutic and adverse effect		
Desipramine	100, single oral dose	500 q12h	↑ 145%	↑ 22%		
		e reduction of desi		ed 15 and 67%, ed when co-administered		
Trazodone	50, single dose	200 q12h	↑ 2.4-fold	↑ 34%		
Trazodone	An increase in the in administered with r enhancer. If trazodo	ncidence in trazodo itonavir dosed as a one is co-administe ing trazodone at th	one-related adverse rea n antiretroviral agent o	ctions was noted when co- r as a pharmacokinetic ombination should be used		

1. Sulfamethoxazole was co-administered with trimethoprim.

Colchicine	Concentrations of colchicine are expected to increase when coadministered with ritonavir. Life-threatening and fatal drug interactions have been reported in patients treated with colchicine and ritonavir (CYP3A4 and P-gp inhibition) in patients with rena and/or hepatic impairment (see sections 4.3 and 4.4). Refer to the colchicine product information.					
Antihistamines						
Astemizole, terfenadine	Ritonavir co-administration is likely to result in increased plasma concentrations of astemizole and terfenadine and is therefore contraindicated (see section 4.3).					
Fexofenadine	Ritonavir may modify P-glycoprotein mediated fexofenadine efflux when dosed as an antriretroviral agent or as a pharmacokinetic enhancer resulting in increased concentrations of fexofenadine. Increased fexofenadine levels may lessen over time as induction develops.					
Loratadine	Ritonavir dosed as a pharmacokinetic enhancer or as an antiretroviral agent inhibits CYP3A and as a result is expected to increase the plasma concentrations of loratadine. Careful monitoring of therapeutic and adverse effects is recommended when loratidine is concomitantly administered with ritonavir.					
Anti-infectives						
Fusidic Acid	Ritonavir co-administration is likely to result in increased plasma concentrations of both fusidic acid and ritonavir and is therefore contraindicated (see section 4.3).					
25-O-desacetyl rifabutin metabolite	↑ 38-fold ↑ 16-fold					
	Due to the large increase in rifabutin AUC, the concomitant use of rifabutin with ritonavir dosed as an antiretroviral agent is contraindicated (see section 4.3). The					
	reduction of the rifabutin dose to 150 mg 3 times per week may be indicated for select PIs when co-administered with ritonavir as a pharmacokinetic enhancer. The product information of the co-administered protease inhibitor should be consulted for specific recommendations. Consideration should be given to official guidance on the appropriate treatment of tuberculosis in HIV-infected patients.					
Rifampicin	PIs when co-administered with ritonavir as a pharmacokinetic enhancer. The product information of the co-administered protease inhibitor should be consulted for specific recommendations. Consideration should be given to official guidance on the appropriate					
Rifampicin Voriconazole	 PIs when co-administered with ritonavir as a pharmacokinetic enhancer. The product information of the co-administered protease inhibitor should be consulted for specific recommendations. Consideration should be given to official guidance on the appropriate treatment of tuberculosis in HIV-infected patients. Although rifampicin may induce metabolism of ritonavir, limited data indicate that when high doses of ritonavir (600 mg twice daily) is co-administered with rifampicin, the additional inducing effect of rifampicin (next to that of ritonavir itself) is small and may have no clinical relevant effect on ritonavir levels in high-dose ritonavir therapy. The 					
_	PIs when co-administered with ritonavir as a pharmacokinetic enhancer. The product information of the co-administered protease inhibitor should be consulted for specific recommendations. Consideration should be given to official guidance on the appropriate treatment of tuberculosis in HIV-infected patients. Although rifampicin may induce metabolism of ritonavir, limited data indicate that when high doses of ritonavir (600 mg twice daily) is co-administered with rifampicin, the additional inducing effect of rifampicin (next to that of ritonavir itself) is small and may have no clinical relevant effect on ritonavir levels in high-dose ritonavir therapy. The effect of ritonavir on rifampicin is not known.					
_	PIs when co-administered with ritonavir as a pharmacokinetic enhancer. The product information of the co-administered protease inhibitor should be consulted for specific recommendations. Consideration should be given to official guidance on the appropriate treatment of tuberculosis in HIV-infected patients. Although rifampicin may induce metabolism of ritonavir, limited data indicate that when high doses of ritonavir (600 mg twice daily) is co-administered with rifampicin, the additional inducing effect of rifampicin (next to that of ritonavir itself) is small and may have no clinical relevant effect on ritonavir levels in high-dose ritonavir therapy. The effect of ritonavir on rifampicin is not known. 200 q12h 100 q12h \downarrow 39% \downarrow 24% Co-administration of voriconazole and ritonavir dosed as a pharmacokinetic enhancer should be avoided, unless an assessment of the benefit/risk to the patient justifies the use					
Voriconazole	PIs when co-administered with ritonavir as a pharmacokinetic enhancer. The product information of the co-administered protease inhibitor should be consulted for specific recommendations. Consideration should be given to official guidance on the appropriate treatment of tuberculosis in HIV-infected patients.Although rifampicin may induce metabolism of ritonavir, limited data indicate that when high doses of ritonavir (600 mg twice daily) is co-administered with rifampicin, the additional inducing effect of rifampicin (next to that of ritonavir itself) is small and may have no clinical relevant effect on ritonavir levels in high-dose ritonavir therapy. The effect of ritonavir on rifampicin is not known.200 q12h100 q12h \downarrow 39% \downarrow 24%Co-administration of voriconazole and ritonavir dosed as a pharmacokinetic enhancer should be avoided, unless an assessment of the benefit/risk to the patient justifies the use of voriconazole.Ritonavir dosed as a pharmacokinetic enhancer or as an antiretroviral agent induces glucuronidation and as a result is expected to decrease the plasma concentrations of atovaquone. Careful monitoring of serum levels or therapeutic effects is recommended					

14-OH clarithromycin metabolite			↓ 100%	↓ 99%	
	necessary in patients per day should not b as a pharmacokinetic dose reduction should	s with normal renal e co-administered c enhancer. For pat ld be considered: fo uld be reduced by 5	with ritonavir dosed as a ients with renal impairn or patients with creatinin 50%, for patients with c	in doses greater than 1 g an antiretroviral agent or nent, a clarithromycin ne clearance of 30 to 60	
Delamanid	interaction study of a twice daily for 14 da increased. Due to the administration of de	delamanid 100 mg ays, the exposure of e risk of QTc prolo lamanid with ritona but the full delaman	the delamanid metabol ngation associated with wir is considered necess id treatment period is re	ir/ritonavir 400/100 mg ite DM-6705 was 30%	
Erythromycin, itraconazole	CYP3A4 and as a re erythromycin and itr	sult is expected to a aconazole. Careful	nhancer or as an antiretr increase the plasma con- monitoring of therapeu raconazole is used conce	centrations of tic and adverse effects is	
Ketoconazole	200 daily	500 q12h	↑ 3.4-fold	↑ 55%	
	incidence of gastroir	ntestinal and hepati be considered whe	etabolism of ketoconazo c adverse reactions, a do en co-administered with netic enhancer.	ose reduction of	
Sulfamethoxazole/ Trimethoprim ¹	800/160, single dose	500 q12h	↓ 20% / ↑ 20%	\leftrightarrow	
	Dose alteration of su should not be necess		methoprim during conco	omitant ritonavir therapy	
Antipsychotics/Neurolep	tics				
Clozapine, pimozide			result in increased plasr contraindicated (see sec		
Haloperidol, risperidone, thioridazine	Ritonavir dosed as an antiretroviral agent is likely to inhibit CYP2D6 and as a result is expected to increase concentrations of haloperidol, risperidone and thioridazine. Careful monitoring of therapeutic and adverse effects is recommended when these medicines are concomitantly administered with antiretroviral doses of ritonavir (see section 4.3).				
Lurasidone	Due to CYP3A inhibition by ritonavir, concentrations of lurasidone are expected to increase. The concomitant administration with lurasidone is contraindicated (see section 4.3).				
Quetiapine	Due to CYP3A inhibition by ritonavir, concentrations of quetiapine are expected to increase. Concomitant administration of ritonavir and quetiapine is contraindicated as it may increase quetiapine-related toxicity.				
β2-agonist (long acting)					
Salmetarol			ult a pronounced increa d. Therefore concomitar		
Calcium channel antago	nists				
Amlodipine, diltiazem, nifedipine			nhancer or as an antiret		

	channel antagonists. Careful monitoring of therapeutic and adverse effects is recommended when these medicines are concomitantly administered with ritonavir.
Contraceptives/HRT	
<i>HRT</i> Dydrogesterone, levonorgestrel, medroxyprogesterone (oral), norethisterone (norethindrone)	Coadministration may increase comedication exposure. The clinical significance of this increase in terms of overall risk of deep vein thrombosis, pulmonary embolism, stroke and myocardial infarction in postmenopausal women receiving substitution hormones in unknown. Postmenopausal women should be re-evaluated periodically to determine if treatment is still necessary.
Drospirenone	Coadministration may increase drospirenone exposure. The clinical significance of this increase in terms of overall risk of deep vein thrombosis, pulmonary embolism, stroke and myocardial infarction in postmenopausal women receiving substitution hormones in unknown. Postmenopausal women should be re-evaluated periodically to determine if treatment is still necessary. Clinical monitoring is recommended due to the potential risk for hyperkalaemia.
Estradiol	Coadministration may decrease comedication exposure. Monitor for signs of hormone deficiency.
Endothelin antagonists	
Bosentan	Co-administration of bosentan and ritonavir may increase steady state bosentan maximum concentr ations (C_{max}) and area under the curve (AUC).
Riociguat	Serum concentrations may be increased due to CYP3A and P-gp inhibition by ritonavir. The co-administration of riociguat with ritonavir is not recommended (see section 4.4 and refer to riociguat product information).
Ergot Derivatives	
Dihydroergotamine, ergonovine, ergotamine, methylergonovine	Ritonavir co-administration is likely to result in increased plasma concentrations of ergot derivatives and is therefore contraindicated (see section 4.3).
GI motility agent	
Cisapride	Ritonavir co-administration is likely to result in increased plasma concentrations of cisapride and is therefore contraindicated (see section 4.3).
HCV Direct Acting Anti	viral
Glecaprevir/pibrentasvir	Seurm concentrations may be increased due to P-glycoprotein, BCRP and OATP1B inhibition by ritonavir.
	Concomitant administration of glecaprevir/pibrentasvir and ritonavir is not recommended due to an increased risk of ALT elevations associated with increased glecaprevir exposure.
HMG Co-A Reductase I	nhibitors
Atorvastatin, fluvastatin, lovastatin, pravastatin, rosuvastatin, simvastatin	HMG-CoA reductase inhibitors which are highly dependent on CYP3A metabolism, such as lovastatin and simvastatin, are expected to have markedly increased plasma concentrations when co-administered with ritonavir dosed as an antiretroviral agent or as a pharmacokinetic enhancer. Since increased concentrations of lovastatin and simvastatin may predispose patients to myopathies, including rhabdomyolysis, the combination of these medicinal products with ritonavir is contraindicated (see section 4.3). Atorvastatin is less dependent on CYP3A for metabolism. While rosuvastatin elimination is not dependent on CYP3A, an elevation of rosuvastatin exposure has been reported with ritonavir co-administration. The mechanism of this interaction is not clear, but may be the result of transporter inhibition. When used with ritonavir dosed as a pharmacokinetic enhancer or as an antiretroviral agent, the lowest possible doses of atorvastatin or rosuvastatin should be administered. The metabolism of pravastatin and fluvastatin is not

	dependent on CYP3A an HMG-CoA reduct recommended.			n ritonavir. If treatment with r fluvastatin is
Hormonal contraceptive				
Ethinyl estradiol	50 µg, single dose	500 q12h	↓ 40%	↓ 32%
	methods of contracep dosed as an antiretroy	tion should be considered agent or as a pheeding profile and re	idered with concom armacokinetic enha	or other non-hormonal nitant ritonavir use when ancer. Ritonavir is likely to ness of estradiol-containing
Immunosupressants				
Cyclosporine, tacrolimus, everolimus	CYP3A4 and as a res cyclosporine, tacrolir	ult is expected to in nus or everolimus.	crease the plasma c Careful monitoring	retroviral agent inhibits oncentrations of of therapeutic and adverse antly administered with
Lipid-modifying agents				
Lomitapide	exposure approximat	ely 27-fold. Due to ected to increase. C	CYP3A inhibition boncomitant use of r	n strong inhibitors increasing by ritonavir, concentrations itonavir with lomitapide is e section 4.3).
Phosphodiesterase (PDE	5) inhibitors			
Avanafil	50, single dose	600 q12h	↑ 13-fold	↑ 2.4-fold
	Concomitant use of a	vanafil with ritonav	ir is contraindicated	l (see section 4.3).
Sildenafil	100, single dose	500 q12h	↑ 11-fold	↑ 4-fold
	dosed as an antiretroy and in no instance sho	viral agent or as a ph ould sildenafil doses e of sildenafil with r	armacokinetic enha	vsfunction with ritonavir ancer should be with caution 8 hours (see also section dicated in pulmonary arterial
Tadalafil	20, single dose	200 q12h	↑ 124%	\leftrightarrow
	pharmacokinetic enha mg tadalafil every 72	ancer should be with hours with increase s used concurrently	n caution at reduced d monitoring for ac with ritonavir in pa	ntiretroviral agent or as a l doses of no more than 10 lverse reactions (see section ttients with pulmonary on.
Vardenafil	5, single dose	600 q12h	↑ 49-fold	↑ 13-fold
	The concomitant use	of vardenafil with r	itonavir is contraine	licated (see section 4.3).
Sedatives/hynoptics				
Clorazepate, diazepam, estazolam, flurazepam, oral and parenteral midazolam and triazolam	clorazepate, estazolar Midazolam is extensi may cause a large inc product interaction st benzodiazepines. Bas midazolam are expec Therefore, ritonavir s (see section 4.3), whe	m and flurazepam ar vely metabolised by rease in the concent udy has been perfor sed on data for other ted to be significant hould not be co-adn ereas caution should	nd is therefore contr CYP3A4. Co-adm ration of this benzo med for the co-adm CYP3A4 inhibitor ly higher when mid ninistered with oral be used with co-ad	asma concentrations of raindicated (see section 4.3). inistration with ritonavir diazepine. No medicinal inistration of ritonavir with s, plasma concentrations of lazolam is given orally. ly administered midazolam ministration of ritonavir and eral midazolam with other

	ritonavir is co-admir care unit (ICU) or si acisappropriate med sedation. Dosage adj	nistered with parenteral milar setting which ens	midazolam, it shou ures close clinical e of respiratory de a should be conside	pression and/or prolonged
Diazepam		ay increase diazepam ex effect should be monit		adjustment may be
Triazolam	0.125, single dose	200, 4 doses	$\uparrow > 20$ fold	↑ 87%
		stration is likely to resu efore contraindicated (s		ma concentrations of
Pethidine	50, oral single dose	500 q12h	↓ 62%	↓ 59%
Norpethidine metabolite			↑ 47%	↑ 87%
	of the metabolite, no	rpethidine, which has b	oth analgesic and	e increased concentrations CNS stimulant activity. CNS effects (eg, seizures)
Alprazolam	1, single dose	200 q12h, 2 days	↑2.5 fold	\leftrightarrow
		500 q12h, 10 days	↓ 12%	↓ 16%
Buspirone	ritonavir dosed as an antiretroviral agent or as a pharmacokinetic enhancer, before induction of alprazolam metabolism develops. Ritonavir dosed as a pharmacokinetic enhancer or as an antiretroviral agent inhibits CYP3A and as a result is expected to increase the plasma concentrations of buspirone. Careful monitoring of therapeutic and adverse effects is recommended when buspirone concomitantly administered with ritonavir.			
Sleeping agent				
Zolpidem	5	200, 4 doses	↑ 28%	↑ 22%
-	Zolpidem and ritona sedative effects.	vir may be co-administ	ered with careful n	nonitoring for excessive
Smoke cessation				
Bupropion	150	100 q12h	↓ 22%	↓ 21%
	150	600 q12h	↓ 66%	↓ 62%
	Bupropion is primarily metabolised by CYP2B6. Concurrent administration of bupropion with repeated doses of ritonavir is expected to decrease bupropion levels. These effects are thought to represent induction of bupropion metabolism. However, because ritonavir has also been shown to inhibit CYP2B6 in vitro, the recommended dose of bupropion should not be exceeded. In contrast to long-term administration of ritonavir, there was no significant interaction with bupropion after short-term administration of low doses of ritonavir (200 mg twice daily for 2 days), suggesting reductions in bupropion concentrations may have onset several days after initiation of ritonavir co-administration.			
Steroids				
Inhaled, injectable or intranasal fluticasone				and adrenal suppression or intranasal fluticasone
	-	Page 19 of 28		

propionate, budesonide, triamcinolone	propionate; similar effects could also occur with other corticosteroids metabolised by CYP3A e.g., budesonide and triamcinolone. Consequently, concomitant administration of ritonavir dosed as an antiretroviral agent or as a pharmacokinetic enhancer and these glucocorticoids is not recommended unless the potential benefit of treatment outweighs the risk of systemic corticosteroid effects (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 that is not a substrate for CYP3A4 (e.g., beclomethasone). Moreover, in case of withdrawal of glucocorticoids, progressive dose reduction may be required over a longer period.			
Dexamethasone	Ritonavir dosed as a pharmacokinetic enhancer or as an antiretroviral agent inhibits CYP3A and as a result is expected to increase the plasma concentrations of dexamethasone. Careful monitoring of therapeutic and adverse effects is recommended when dexamethasone is concomitantly administered with ritonavir.			
Prednisolone	20	200 q12h	↑ 28%	↑ 9%
	Careful monitoring of therapeutic and adverse effects is recommended when prednisolone is concomitantly administered with ritonavir. The AUC of the metabolite prednisolone increased by 37 and 28% after 4 and 14 days ritonavir, respectively.			
Thyroid hormone replacement therapy				
Levorthyroxine	Post-marketing cases have been reported indicating a potential interaction between ritonavir containing products and levothyroxine. Thyroid-stimulating hormone (TSH) should be monitored in patients with levothyroxine at least the first month after starting and/or ending ritonavir treatment.			

Cardiac and neurologic events have been reported when ritonavir has been co-administered with disopyramide, mexiletine or nefazodone. The possibility of medicinal product interaction cannot be excluded.

In addition to the interactions listed above, as ritonavir is highly protein bound, the possibility of increased therapeutic and toxic effects due to protein binding displacement of concomitant medicinal products should be considered.

Further information regarding medicinal product interactions when ritonavir is used a pharmacokinetic enhancer is also contained in the product information of the coadministered protease inhibitor.

Proton pump inhibitors and H2-receptor antagonists (e.g. omeprazole or ranitidine) may reduce concentrations for co-administered protease inhibitors. For specific information regarding the impact of co-administration of acid-reducing agents, refer to the product information of the co-administered protease inhibitor. Based on interaction studies with the ritonavir boosted protease inhibitors (lopinavir/ritonavir, atazanavir), concurrent administration of omeprazole or ranitidine does not significantly modify ritonavir efficacy as a pharmacokinetic enhancer despite a slight change of exposure (about 6-18%).

4.6 Fertility, pregnancy and breastfeeding

Pregnancy

A large number of pregnant women (corresponding to 6100 live births) were exposed to ritonavir during pregnancy; of these, 2800 live births were exposed during the first trimester. These data largely refer to exposure of ritonavir used as a booster for protease inhibitors in combination therapy. There was no increase in the rate of birth defects compared to rates in population-based birth defect surveillance systems. Animal data have shown reproductive toxicity (see section 5.3).

Ritonavir Tablets USP 100 mg can be used during pregnancy if clinically needed.

Ritonavir interacts with oral contraceptives. Therefore, an alternative, effective and safe method of contraception should be used during treatment.

Breast-feeding

Ritonavir has been detected in human milk. There is no information on the effects of ritonavir on the breastfed infant or the effects of the medicine on milk production. Current recommendations on HIV and breastfeeding (e.g. those from the WHO) should be consulted before advising patients on this matter. Preferred options may vary depending on the local circumstances.

Fertility

No human data on the effect of ritonavir on fertility are available. Animal studies do not indicate harmful effects of ritonavir on fertility (see section 5.3).

4.7 Effects on ability to drive and use machines

No studies on the effects on the ability to drive and use machines have been performed. Dizziness is a known undesirable effect that should be borne in mind when considering a patient's ability to drive or operate machinery (see section 4.8).

4.8 Undesirable effects

Adverse reactions associated with the use of ritonavir as a pharmacokinetic enhancer are dependent on the specific co-administered protease inhibitor. For information on adverse reactions refer to the product information of the specific co-administered protease inhibitor.

The following adverse reactions were reported from clinical trials and post-marketing experience in adult patients with ritonavir dosed as antiretroviral agent.

Summary of the safety profile

The most frequent adverse reactions among patients receiving ritonavir alone or in combination with other antiretroviral drugs were gastrointestinal (including diarrhea, nausea, vomiting, abdominal pain [upper and lower]), neurological disturbances (including paresthesia and oral paresthesia) and fatigue/asthenia.

Tabulated list of adverse reactions

The following adverse reactions of moderate to severe intensity with possible or probable relationship to ritonavir have been reported. Within each frequency grouping, undesirable effects are presented in order of decreasing seriousness: very common ($\geq 1/10$), common ($\geq 1/100$ to < 1/10), uncommon ($\geq 1/1,000$ to < 1/1,000 to < 1/1,000 to < 1/1,000) and not known (frequency cannot be estimated from the available data).

Events noted as having frequency not known were identified via post-marketing surveillance.

MedDRA system organ class	Adverse reaction
Frequency category	
Blood and lymphatic system disorders	
common	decreased white blood cells, decreased haemoglobin, decreased neutrophils, increased eosinophils, thrombocytopenia
uncommon	increased neutrophils
Immune system disorders	
common	hypersenstitivity including urticaria, and face oedema
rare	anaphylaxis
Metabolism and nutrition disorders	

Adverse reactions in clinical trials and post-marketing

common	hypercholesterolaemia, hypertriglyceridaemia, gout, oedema and peripheral oedema, dehydration (usually associated with gastrointestinal symptoms)	
uncommon	diabetes mellitus	
rare	hyperglycaemia	
Nervous system disorders		
very common	dysgeusia, oral and peripheral paraesthesia, headache, dizziness, peripheral neuropathy	
common	insomnia, anxiety, confusion, disturbance in attention, syncope, seizure	
Eye disorders		
common	blurred vision	
Cardiac disorders		
uncommon	myocardial infarction	
Vascular disorders		
common	hypertension, hypotension including orthostatic hypotension, peripheral coldness	
Respiratory, thoracic and mediastinal a	disorders	
very common	pharyngitis, oropharyngeal pain, cough	
Gastrointestinal disorders		
very common	abdominal pain (upper and lower), nausea, diarrhoea (including severe with electrolyte imbalance), vomiting, dyspepsia	
common	anorexia, flatulence, mouth ulcer, gastrointestinal haemorrhage, gastroesophageal reflux disease, pancreatitis	
Hepatobiliary disorders		
common	hepatitis (including increased AST, ALT, GGT), blood bilirubin increased (including jaundice)	
Skin and subcutaneous tissue disorders		
very common	pruritus, rash (including erythematous and maculopapular)	
common	acne	
rare	Stevens Johnson syndrome, toxic epidermal necrolysis (TEN)	
Musculoskeletal and connective tissue a	disorders	
very common	arthralgia and back pain	
common	myositis, rhabdomyolysis, myalgia, myopathy/CPK increased	
Renal and urinary disorders		
common	increased urination, renal impairment (e.g. oliguria, elevated creatinine)	
uncommon	acute renal failure	
not known	nephrolithiasis	
Reproductive system and breast disorde	ers	
common	menorrhagia	

General disorders and administration site conditions		
very common	fatigue including asthenia, flushing, feeling hot	
common	fever, weight loss	
Investigations		
common	increased amylase, decreased free and total thyroxin	
uncommon	increased glucose, increased magnesium, increased alkaline phosphatase	

Description of selected adverse reactions

Hepatotoxicity

Hepatic transaminase elevations exceeding five times the upper limit or normal, clinical hepatitis, and jaundice have occurred in patients receiving ritonavir alone or in combination with other antiretrovirals.

Metabolic parameters

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

Immune reconstitution inflammatory syndrome

In patients with severe immune deficiency at the time of initiation of 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 time to onset is more variable and these events can occur many months after starting treatment (see section 4.4).

Pancreatitis

Pancreatitis has been observed in patients receiving ritonavir therapy, including those who developed hypertriglyceridaemia. In some cases fatalities have been observed. Patients with advanced HIV disease may be at risk of elevated triglycerides and pancreatitis (see section 4.4).

Osteonecrosis

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).

Paediatric populations

The safety profile of ritonavir in children 2 years of age and older is similar to that seen in adults.

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. Health care providers are asked to report any suspected adverse reactions to the marketing authorisation holder, or, if available, via the national reporting system.

4.9 Overdose

Symptoms

Human experience of acute overdose with ritonavir is limited. One patient in clinical trials took ritonavir 1500 mg/day for two days and reported paraesthesia, which resolved after the dose was decreased. A case of renal failure with eosinophilia has been reported.

The signs of toxicity observed in animals (mice and rats) included decreased activity, ataxia, dyspnoea and tremors.

Management

There is no specific antidote for overdose with ritonavir. Treatment of overdose with ritonavir should consist of general supportive measures including monitoring of vital signs and observation of the clinical status of the patient. Due to the solubility characteristics and possibility of transintestinal elimination, it is proposed that management of overdose could entail gastric lavage and administration of activated charcoal. Since ritonavir is extensively metabolised by the liver and is highly protein bound, dialysis is unlikely to be beneficial in significant removal of the medicine.

5. PHARMACOLOGICAL PROPERTIES

5.1 Pharmacodynamic properties

Pharmacotherapeutic group: Antivirals for systemic use, protease inhibitors, ATC code: J05AE03.

Mechanism of action

Pharmacokinetic enhancement by ritonavir is based on ritonavir's activity as a potent inhibitor of CYP3Amediated metabolism. The degree of enhancement is related to the metabolic pathway of the co-administered protease inhibitor and the impact of the co-administered protease inhibitor on the metabolism of ritonavir. Maximal inhibition of metabolism of darunavir is generally achieved with ritonavir doses of 100 mg daily to 200 mg twice daily. For additional information on the effect of ritonavir on co-administered protease inhibitor metabolism, see section 4.5 and consult the product information of the particular co-administered protease inhibitor.

Effects on the electrocardiogram

QTcF interval was evaluated in a randomised, placebo and active (moxifloxacin 400 mg once daily) controlled crossover study in 45 healthy adults, with 10 measurements over 12 hours on Day 3. The maximum mean (95% upper confidence bound) difference in QTcF from placebo was 5.5 (7.6) for 400 mg twice daily ritonavir. The Day 3 ritonavir exposure was approximately 1.5 fold higher than that observed with the 600 mg twice daily dose at steady state. No subject experienced an increase in QTcF of \geq 60 msec from baseline or a QTcF interval exceeding the potentially clinically relevant threshold of 500 msec.

Modest prolongation of the PR interval was also noted in subjects receiving ritonavir in the same study on Day 3. The mean changes from baseline in PR interval ranged from 11.0 to 24.0 msec in the 12 hour interval post dose. Maximum PR interval was 252 msec and no second or third degree heart block was observed (see section 4.4).

Resistance

Ritonavir-resistant isolates of HIV-1 have been selected in vitro and isolated from patients treated with therapeutic doses of ritonavir.

Reduction in the antiretroviral activity of ritonavir is primarily associated with the protease mutations V82A/F/T/S and I84V. Accumulation of other mutations in the protease gene (including at positions 20, 33, 36, 46, 54, 71, and 90) can also contribute to ritonavir resistance. In general, as mutations associated with ritonavir resistance accumulate, susceptibility to select other protease inhibitors may decrease due to cross-resistance. The summary of product characteristics of other protease inhibitors or official continuous updates should be consulted for specific information regarding protease mutations associated with reduced response to these agents.

Clinical efficacy and safety data

Ritonavir was the first protease inhibitor (approved in 1996) for which efficacy was proven in a study with clinical endpoints. The effects of ritonavir (alone or combined with other antiretroviral agents) on biological markers of disease activity such as CD4 cell count and viral RNA were evaluated in several studies involving HIV-1 infected patients. However, due to ritonavir's metabolic inhibitory properties its use as a pharmacokinetic enhancer of other protease inhibitors is the prevalent use of ritonavir in clinical practice (see section 4.2).

5.2 Pharmacokinetic properties

The absorption characteristics of Ritonavir Tablets USP 100 mg have been determined after administration of oneritonavir 100 mg tablet in healthy volunteers in the fasted state as follows:

Pharmacokinetic variable	Mean value* (±standard deviation)
	Ritonavir
Maximum concentration (C _{max})	798 ± 366 ng/mL
Area under the curve (AUC $_{0-\infty}$), a measure of the extent of absorption	$6536 \pm 2856 \text{ ng} \cdot \text{h/mL}$
Time to attain maximum concentration (T _{max})	$3.44 \pm 1.28 \text{ h}$

*arithmetic mean

Pharmacokinetics of Ritonavir

	Ritonavir		
General			
Absorption			
Oral bioavailability	NA		
Food effect	Food slightly decreases the bioavailability of ritonavir tablets.		
	A single oral dose of ritonavir 100 mg with a moderate fat meal (857 kcal, 31% calories from fat) or a high fat meal (907 kcal, 52% calories from fat) was associated with a mean decrease of 20-23% in ritonavir AUC and C_{max} .		
Distribution			
Volume of distribution (mean ± SD)	After single 600 mg dose: approximately 20–40L		
Plasma protein	Approximately 98–99% and is constant over the concentration range of $1-100 \ \mu g/ml$.		
binding in vitro	Ritonavir binds to both human alpha 1-acid glycoprotein (AAG) and human serum albumin (HSA) with comparable affinities.		
Tissue distribution	Studies in rats showed highest concentrations of ritonavir in the liver, adrenals, pancreas, kidneys and thyroid.		
	Tissue to plasma ratios of approximately 1 measured in rat lymph nodes suggest that ritonavir distributes into lymphatic tissues.		
	Ritonavir penetrates minimally into the brain.		
Metabolism			
	Primarily oxidative metabolism according to animal studies and <i>in vitro</i> experiments with human liver microsomes (HLMs).		
	Four ritonavir metabolites have been identified in man. The isopropylthiazole oxidation metabolite (M-2) is the major metabolite.		
	Low doses of ritonavir have shown profound effects on the pharmacokinetics of other protease inhibitors (and other products metabolised by CYP3A4) and other protease inhibitors may influence the pharmacokinetics of ritonavir (see section 4.5).		

Active metabolite(s)	M-2 has antiviral activity similar to that of parent compound but its AUC was approximately 3% of the AUC of parent compound.
Elimination	
Elimination half life	NA
Mean systemic clearance (Cl/F)	NA
% of dose excreted in urine	Renal clearance of ritonavir is negligible
% of dose excreted in faeces	86%; part of which is expected to be unabsorbed ritonavir
Drug interactions (in vitro)
Transporters	P-glycoprotein and anion-transporting polypeptides
Metabolising enzymes	Hepatic CYP system, primarily by the CYP3A isozyme family and to a lesser extent by the CYP2D6 isoform

Pharmacokinetics in special populations

Paediatric population

Ritonavir steady-state pharmacokinetic parameters were evaluated in HIV-infected children above 2 years of age receiving doses ranging from 250 mg/m² twice daily to 400 mg/m² twice daily. Ritonavir concentrations obtained after 350 to 400 mg/m² twice daily in paediatric patients were comparable to those obtained in adults receiving 600 mg (approximately 330 mg/m²) twice daily. Across dose groups, ritonavir oral clearance (CL/F/m²) was approximately 1.5 to 1.7 times faster in paediatric patients above 2 years of age than in adult subjects.

Ritonavir steady-state pharmacokinetic parameters were evaluated in HIV infected children less than 2 years of age receiving doses ranging from 350 to 450 mg/m² twice daily. Ritonavir concentrations in this study were highly variable and somewhat lower than those obtained in adults receiving 600 mg (approximately 330 mg/m²) twice daily. Across dose groups, ritonavir oral clearance (CL/F/m²) declined with age with median values of 9.0 L/h/m² in children less than 3 months of age, 7.8 L/h/m² in children between 3 and 6 months of age and 4.4 L/h/m² in children between 6 and 24 months of age.

Elderly

Plasma exposures in patients 50–70 years of age when dosed 100 mg in combination with lopinavir or at higher doses in the absence of other protease inhibitors is similar to that observed in younger adults.

Gender

No clinically significant differences in AUC or C_{max} were noted between males and females.

Renal impairment

Ritonavir pharmacokinetic parameters have not been studied in patients with renal impairment. However, since the renal clearance of ritonavir is negligible, no changes in the total body clearance are expected in patients with renal impairment.

Hepatic impairment

After multiple dosing to healthy volunteers (500 mg twice daily) and subjects with mild to moderate hepatic impairment (Child Pugh Class A and B, 400 mg twice daily) exposure to ritonavir after dose normalisation was not significantly different between the two groups.

5.3 Preclinical safety data

Repeated dose toxicity studies in animals identified major target organs as the liver, retina, thyroid gland and kidney. Hepatic changes involved hepatocellular, biliary and phagocytic elements and were accompanied by increases in hepatic enzymes. Hyperplasia of the retinal pigment epithelium (RPE) and retinal degeneration have been seen in all of the rodent studies conducted with ritonavir, but have not been seen in dogs. Ultrastructural evidence suggests that these retinal changes may be secondary to phospholipidosis. All thyroid changes were reversible upon discontinuation of ritonavir. Renal changes including tubular degeneration, chronic inflammation and proteinurea were noted in rats and are felt to be attributable to species-specific spontaneous disease.

Developmental toxicity observed in rats (embryolethality, decreased fetal body weight and ossification delays and visceral changes, including delayed testicular descent) occurred mainly at a maternally toxic dosage. Developmental toxicity in rabbits (embryolethality, decreased litter size and decreased fetal weights) occurred at a maternally toxic dosage.

Ritonavir was not found to be mutagenic or clastogenic in a battery of *in vitro* and *in vivo* assays including the Ames bacterial reverse mutation assay using *S. typhimurium* and *E. coli*, the mouse lymphoma assay, the mouse micronucleus test and chromosomal aberration assays in human lymphocytes.

Long-term carcinogenicity studies of ritonavir in mice and rats revealed tumourigenic potential specific for these species, but are regarded as of no relevance for humans.

6. PHARMACEUTICAL PARTICULARS

6.1 List of excipients *Core tablet:* Copovidone Colloidal silicon dioxide Sorbitan monolaurate Dibasic calcium phosphate anhydrous Sodium stearyl fumarate

Film coat: Hypromellose Titanium dioxide Macrogol/PEG Hydroxypropyl cellulose Talc Colloidal anhydrous silica Polysorbate 80

6.2 Incompatibilities

Not applicable.

6.3 Shelf life

24 months

6.4 Special precautions for storage

Do not store above 30°C. Protect from moisture.

6.5 Nature and contents of container

40cc heavy weight high density polyethylene (HDPE) bottles closed with 33mm polypropylene ribbed child-resistant plastic caps with a pulp liner and heat seal liner. Pack Size: 30 Tablets.

6.6 Special precautions for disposal and other handling

No special requirements.

7. SUPPLIER

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8. MARKETING AUTHORISATION NUMBER

06417/08321/NMR/2020

9. DATE OF AUTHORISATION/RENEWAL OF THE AUTHORISATION

Jul 26, 2021

10. DATE OF REVISION OF THE TEXT

May 2022

References

General reference sources for this SmPC include:

World Health Organization (2021) Consolidated guidelines on HIV prevention, testing, treatment, service delivery and monitoring: recommendations for a public health approach, available at https://www.who.int/publications/i/item/9789240031593

World Health Organization (2016) Consolidated guidelines on the use of antiretroviral drugs for treating and preventing HIV infection: recommendations for a public health approach, 2nd ed. World Health Organization https://apps.who.int/iris/handle/10665/208825

European SmPC, Norvir, available at:

https://www.ema.europa.eu/en/documents/product-information/norvir-epar-product-information_en.pdf

All weblinks were last accessed on 10 May 2022.

Detailed information on this medicine is available on the World Health Organization (WHO) website: <u>https://extranet.who.int/pgweb/medicines</u>