

EDURANT[®]

rilpivirine hydrochloride

DATA SHEET

1. PRODUCT NAME

EDURANT[®] 25 mg film-coated tablets.

2. QUALITATIVE AND QUANTITATIVE COMPOSITION

Each film-coated tablet contains rilpivirine hydrochloride equivalent to 25 mg rilpivirine.

Excipient(s) with known effect

Each film-coated tablet contains 56 mg lactose monohydrate.

For a full list of excipients, see **section 6.1**.

3. PHARMACEUTICAL FORM

Film-coated tablet

EDURANT 25 mg tablets are white to off-white, film coated, round, biconvex, tablet of 6.4 mm, debossed with "TMC" on one side and "25" on the other side.

Rilpivirine is a non-nucleoside reverse transcriptase inhibitor (NNRTI) of human immunodeficiency virus type-1 (HIV-1).

4. CLINICAL PARTICULARS

4.1 Therapeutic indications

EDURANT, in combination with other antiretroviral medicinal products, is indicated for the treatment of human immunodeficiency virus type 1 (HIV-1) infection in antiretroviral treatment-naïve adult patients with viral load \leq 100,000 copies/mL at baseline.

This indication is based on Week 48 safety and efficacy analyses from 2 randomised double-blind, controlled Phase III trials in treatment-naïve adult patients and on Week 96 safety and efficacy analyses from the Phase IIb trial TMC278-C204 in treatment-naïve adult patients (see **section 5.1 Clinical Trials**).

4.2 Dose and method of administration

EDURANT must always be given in combination with other antiretroviral medicinal products.

Dosage

The recommended dose of EDURANT in adults is one 25 mg tablet once daily taken orally with a meal (see **section 5.2**).

Dose adjustment with rifabutin coadministration

For patients concomitantly receiving rifabutin, the EDURANT dose should be increased to 50 mg (two tablets of 25 mg each) once daily, taken with a meal. When rifabutin co-administration is

stopped, the EDURANT dose should be decreased to 25 mg once daily, taken with a meal (see **section 4.5**).

Timing of dosing

If the patient misses a dose of EDURANT within 12 hours of the time it is usually taken, the patient should take EDURANT with a meal as soon as possible and then take the next dose of EDURANT at the regularly scheduled time. If a patient misses a dose of EDURANT by more than 12 hours, the patient should not take the missed dose, but resume the usual dosing schedule.

Special populations

Elderly

No dose adjustment of EDURANT is required in elderly patients (see section 5.2).

Paediatric population

The safety and efficacy of EDURANT in paediatric patients (less than 18 years) have not been established. Treatment with EDURANT is not recommended in these populations.

Hepatic impairment

No dose adjustment of EDURANT is required in patients with mild or moderate hepatic impairment (Child Pugh score A or B). EDURANT has not been studied in patients with severe hepatic impairment (Child Pugh score C). EDURANT should be used with caution in patients with moderate to severe hepatic impairment (see **sections 5.2** and **4.4**).

Renal impairment

No dose adjustment of EDURANT is required in patients with renal impairment (see section 5.2).

Pregnancy and postpartum

The recommended dose of EDURANT in pregnant patients is one 25 mg tablet once daily taken orally with a meal. Lower exposures of rilpivirine were observed during pregnancy, therefore viral load should be monitored closely (see **sections 4.6** and **5.2**).

4.3 Contraindications

Hypersensitivity to rilpivirine or to any of the excipients.

EDURANT should not be co-administered with the following medicinal products, as significant decreases in rilpivirine plasma concentrations may occur (due to CYP3A enzyme induction or gastric pH increase), which may result in loss of therapeutic effect of EDURANT (see **section 4.5**):

- the anticonvulsants carbamazepine, oxcarbazepine, phenobarbital, phenytoin
- the antimycobacterials; rifampicin, rifapentine
- proton pump inhibitors, such as omeprazole, esomeprazole, lansoprazole, pantoprazole, rabeprazole
- the glucocorticoid systemic dexamethasone, except as a single dose treatment
- St John's wort (*Hypericum perforatum*).

4.4 Special warnings and precautions for use

Transmission of HIV

Patients should be advised that current antiretroviral therapy does not cure HIV. Appropriate precautions to prevent the transmission of HIV should continue to be employed.

Virologic failure and development of resistance

In the pooled analysis from the Phase III trials through 96 weeks, patients treated with EDURANT with a baseline viral load > 100,000 HIV-1 RNA copies/mL had a greater risk of virologic failure (18.2% with EDURANT versus 7.9% efavirenz arm) compared to patients with a baseline viral load ≤100,000 HIV-1 RNA copies/mL (5.7% with EDURANT versus 3.6% efavirenz arm). The greater risk of virologic failure for patients in the EDURANT arm was observed in the first 48 weeks of these trials while low rates of virologic failure similar between treatment arms were observed from week 48 to week 96 (see **section 5.1, Clinical trials**). Patients with a baseline viral load > 100,000 HIV-1 RNA copies/mL who experienced virologic failure exhibited a higher rate of treatment emergent resistance to the NNRTI class. More patients who failed virologically on EDURANT than who failed virologically on efavirenz developed lamivudine/emtricitabine associated resistance. This information should be taken into consideration when initiating therapy with EDURANT (see **section 5.1, Clinical trials**).

Interactions with medicinal products

Caution should be given to prescribing EDURANT with medicinal products that may reduce the exposure of rilpivirine. For information on interactions with medicinal products (see **sections 4.3** and **4.5**).

CYP3A metabolism

Rilpivirine is a CYP3A substrate. It is possible that different populations of patients have faster or slower rilpivirine metabolism because of the various isoenzymes within the CYP3A system.

Immune reconstitution inflammatory syndrome

Immune reconstitution inflammatory syndrome has been reported in patients treated with combination antiretroviral therapy, including EDURANT. During the initial phase of combination antiretroviral treatment, patients whose immune system responds may develop an inflammatory response to indolent or residual opportunistic infections (such as *Mycobacterium avium* complex, cytomegalovirus, *Pneumocystis jiroveci* pneumonia, and tuberculosis), which may necessitate further evaluation and treatment (see **section 4.8**). Autoimmune disorders such as Graves' disease and autoimmune hepatitis have also been reported to occur in the setting of immune reconstitution inflammatory syndrome; however, the time to onset is more variable, and these events can occur many months after initiation of treatment (see **section 4.8**)

Special populations

Children

Treatment with EDURANT is not recommended in paediatric patients due to insufficient data in this patient population. (see **sections 5.2** and **4.2**).

Elderly

No dose adjustment of EDURANT is required in elderly patients.

Hepatic impairment

No dose adjustment of EDURANT is required in patients with mild or moderate hepatic impairment. EDURANT has not been studied in patients with severe hepatic impairment (see **sections 5.2** and **4.2**). EDURANT should be used with caution in patients with moderate to severe hepatic impairment (see **sections 5.2** and **4.2**).

4.5 Interactions with other medicines and other forms of interaction

Medicinal products that affect rilpivirine exposure

Rilpivirine is primarily metabolised by cytochrome P450 CYP3A, and medicinal products that induce or inhibit CYP3A may thus affect the clearance of rilpivirine. Co-administration of EDURANT and medicinal products that induce CYP3A may result in decreased plasma concentrations of rilpivirine which could potentially reduce the therapeutic effect of EDURANT. Co-administration of EDURANT and medicinal products that inhibit CYP3A may result in increased plasma concentrations of rilpivirine.

Co administration of EDURANT with medicinal products that increase gastric pH may result in decreased plasma concentrations of rilpivirine which could potentially reduce the therapeutic effect of EDURANT.

Medicines that suppress gastric acid

Use of proton pump inhibitors is contraindicated as significant decreases in EDURANT plasma concentrations may occur (see **section 4.3**).

The combination of EDURANT and H_2 -receptor antagonists or antacids should be used with caution as co-administration may cause significant decreases in rilpivirine plasma concentrations (gastric pH increase). H_2 -receptor antagonists should only be administered at least 12 hours before or at least 4 hours after EDURANT and antacids should only be administered at least 2 hours before or at least 4 hours after EDURANT.

Medicinal products that are affected by the use of rilpivirine

EDURANT at a dose of 25 mg q.d. is not likely to have a clinically relevant effect on the exposure of medicinal products metabolised by CYP enzymes.

Established and theoretical interactions with selected antiretrovirals and non-antiretroviral medicinal products are listed below in **Table 1** and **Table 2**, respectively.

Interactions between rilpivirine and co-administered medicinal products are listed in the tables below (increase is indicated as " \uparrow ", decrease as " \downarrow ", no change as " \leftrightarrow ", not applicable as "NA", once daily as "q.d." and twice daily as "b.i.d.").

products					
Co-administered medicinal product	Dose of co-administered medicinal product	Medicinal product assessed	C _{max}	AUC	C _{min}
HIV NUCLEOSIDE OR	NUCLEOTIDE REVE		PTASE INHIB	ITORS (NR1	[Is/N[t]RTIs)
Didanosine*#	400 mg q.d.	didanosine	\leftrightarrow	↑ 12%	NA
		rilpivirine	\leftrightarrow	\leftrightarrow	\leftrightarrow
Tenofovir disoproxil	should be adminis 300 mg q.d.	tenofovir	ıl). ↑ 19%	↑ 23%	↑ 24%
fumarate*#	No dose adjustme tenofovir disoprox	•	↔ en EDURANT	↔ is co-admini	\leftrightarrow stered with
Other NRTIs (abacavir, emtricitabine, lamivudine, stavudine and zidovudine)	Based on the diffe NRTIs, no clinical these medicinal p	ly relevant drug-o	lrug interaction		

 Table 1: Drug interactions – Rilpivirine co-administered with antiretroviral and antiviral medicinal products

medicinal product	Dose of co-administered medicinal product	Medicinal product assessed	C _{max}	AUC	C _{min}
HIV NON-NUCLEOSIDE					
NNRTIs (delavirdine, efavirenz, etravirine, nevirapine)	It is not recommer	nded to co-admin	ister EDURAN	T with NNR	Гls.
HIV PROTEASE INHIBIT		-administration	of low dose ri	itonavir	
Darunavir/ritonavir*#	800/100 mg q.d. Concomitant use of increase in the pla enzymes). No dos co-administered w	asma concentrationse adjustment is r	ons of rilpivirine equired when E	e (inhibition of	of CYP3A
Lopinavir/ritonavir (soft gel capsules)* [#]	400/100 mg b.i.d. Concomitant use of increase in the pla enzymes). No dos co-administered w	asma concentrationse adjustment is r	ons of rilpivirine equired when E	e (inhibition o	of CYP3A
Other boosted PIs (atazanavir/ritonavir, fosamprenavir/ritonavir, saquinavir/ritonavir, tipranavir/ritonavir)	Concomitant use of in the plasma con EDURANT is not of co-administered P	of EDURANT wit centrations of rilp expected to affec	h boosted PIs r vivirine (inhibitio	on of CYP3A	enzymes).
(atazanavir/ritonavir, fosamprenavir/ritonavir, saquinavir/ritonavir,	Concomitant use of in the plasma con- EDURANT is not of co-administered P	of EDURANT wit centrations of rilp expected to affec Pls.	h boosted PIs r vivirine (inhibitio t the plasma co	on of CYP3A oncentration	enzymes).
(atazanavir/ritonavir, fosamprenavir/ritonavir, saquinavir/ritonavir, tipranavir/ritonavir) HIV PROTEASE INHIBIT Unboosted PIs (atazanavir, fosamprenavir, indinavir, nelfinavir)	Concomitant use of in the plasma con- EDURANT is not of co-administered P	of EDURANT wit centrations of rilp expected to affect Pls. t co-administrat of EDURANT with centrations of rilp expected to affect	h boosted PIs r pivirine (inhibitio at the plasma co ition of low dos n unboosted PIs ivirine (inhibitio	on of CYP3A concentration se ritonavir s may cause n of CYP3A	enzymes). s of an increase enzymes).
(atazanavir/ritonavir, fosamprenavir/ritonavir, saquinavir/ritonavir, tipranavir/ritonavir) HIV PROTEASE INHIBIT Unboosted PIs (atazanavir, fosamprenavir, indinavir, nelfinavir) CCR5 ANTAGONISTS	Concomitant use of in the plasma con- EDURANT is not of co-administered P CORS (PIs) - withou Concomitant use of in the plasma cond EDURANT is not of co-administered P	of EDURANT wit centrations of rilp expected to affect Pls. t co-administrat of EDURANT with centrations of rilp expected to affect ls.	h boosted PIs r pivirine (inhibitio at the plasma co tion of low dos n unboosted PIs ivirine (inhibition t the plasma co	on of CYP3A concentration se ritonavir s may cause n of CYP3A ncentrations	a enzymes). s of e an increas enzymes). s of
(atazanavir/ritonavir, fosamprenavir/ritonavir, saquinavir/ritonavir, tipranavir/ritonavir) HIV PROTEASE INHIBIT Unboosted PIs (atazanavir, fosamprenavir, indinavir, nelfinavir)	Concomitant use of in the plasma cont EDURANT is not of co-administered P Concomitant use of in the plasma cont EDURANT is not of	of EDURANT wit centrations of rilp expected to affect ls. t co-administrat of EDURANT with centrations of rilp expected to affect ls.	h boosted PIs r pivirine (inhibitio at the plasma co tion of low dos n unboosted PIs ivirine (inhibition t the plasma co	on of CYP3A concentration se ritonavir s may cause n of CYP3A ncentrations	a enzymes). s of e an increase enzymes). s of
(atazanavir/ritonavir, fosamprenavir/ritonavir, saquinavir/ritonavir, tipranavir/ritonavir) HIV PROTEASE INHIBIT Unboosted PIs (atazanavir, fosamprenavir, indinavir, nelfinavir) CCR5 ANTAGONISTS Maraviroc	Concomitant use of in the plasma cont EDURANT is not of co-administered P ORS (PIs) - withou Concomitant use of in the plasma conto EDURANT is not e co-administered P No clinically relevat co-administered w	of EDURANT wit centrations of rilp expected to affect Pls. t co-administrat of EDURANT with centrations of rilp expected to affect ls. ant drug-drug inte ith maraviroc.	h boosted PIs r pivirine (inhibitio at the plasma co tion of low dos n unboosted PIs ivirine (inhibition t the plasma co	on of CYP3A concentration se ritonavir s may cause n of CYP3A ncentrations	e enzymes). s of an increase enzymes). s of
(atazanavir/ritonavir, fosamprenavir/ritonavir, saquinavir/ritonavir, tipranavir/ritonavir) HIV PROTEASE INHIBIT Unboosted PIs (atazanavir, fosamprenavir, indinavir, nelfinavir) CCR5 ANTAGONISTS Maraviroc	Concomitant use of in the plasma cont EDURANT is not of co-administered P ORS (PIs) - withou Concomitant use of in the plasma conto EDURANT is not e co-administered P No clinically relevat co-administered w	of EDURANT wit centrations of rilp expected to affect Pls. t co-administrat of EDURANT with centrations of rilp expected to affect ls. ant drug-drug inte ith maraviroc. IBITORS raltegravir rilpivirine	h boosted PIs r bivirine (inhibitio et the plasma co tion of low dos n unboosted PIs ivirine (inhibitio t the plasma co eraction is experi- raction is experi- ↑ 10% ↔	n of CYP3A oncentration e ritonavir s may cause n of CYP3A ncentrations cted when E ↑ 9% ↔	a enzymes). s of e an increas enzymes). s of EDURANT is ↑ 27% ↔
(atazanavir/ritonavir, fosamprenavir/ritonavir, saquinavir/ritonavir, tipranavir/ritonavir) HIV PROTEASE INHIBIT Unboosted PIs (atazanavir, fosamprenavir, indinavir, nelfinavir) CCR5 ANTAGONISTS Maraviroc HIV INTEGRASE STRAM	Concomitant use of in the plasma cont EDURANT is not of co-administered P ORS (PIs) - withou Concomitant use of in the plasma cont EDURANT is not e co-administered P No clinically relevat co-administered w ID TRANSFER INH 400 mg b.i.d. No dose adjustme raltegravir.	of EDURANT wit centrations of rilp expected to affect Pls. t co-administrat of EDURANT with centrations of rilp expected to affect ls. ant drug-drug inte ith maraviroc. IBITORS raltegravir rilpivirine	h boosted PIs r bivirine (inhibitio et the plasma co tion of low dos n unboosted PIs ivirine (inhibitio t the plasma co eraction is experi- raction is experi- ↑ 10% ↔	n of CYP3A oncentration e ritonavir s may cause n of CYP3A ncentrations cted when E ↑ 9% ↔	a enzymes). s of e an increase enzymes). s of EDURANT is ↑ 27% ↔

The interaction between EDURANT and the drug was evaluated in a clinical study. All other drug interactions shown are predicted.

This interaction study has been performed with a dose higher than the recommended dose for EDURANT assessing the maximal effect on the co-administered drug. The dosing recommendation is applicable to the recommended dose of EDURANT 25 mg q.d.

Co-administered medicinal product	Dose of co-administered medicinal product	Medicinal product assessed	C _{max}	AUC	C _{min}
ANTIARRHYTHMICS					
Digoxin*	0.5 mg single dose	digoxin	\leftrightarrow	\leftrightarrow	NA
	No dose adjustment digoxin.	t is required when EDURA	NT is co-a	administere	ed with
ANTIDIABETICS					
Metformin*		metformin t is required when EDURA	↔ NT is co-a	↔ administere	NA ed with
	metformin.				
ANTICONVULSANTS				4	
Carbamazepine		not be used in combination			
Oxcarbazepine		n may cause significant de			
Phenobarbital		uction of CYP3A enzymes). This ma	y result in	IOSS OF
Phenytoin	therapeutic effect of	EDUKANI.			
AZOLE ANTIFUNGAL		laste e comenta		1.0.40/	1.000/
Ketoconazole*#	400 mg q.d.	ketoconazole	\leftrightarrow	↓ 24%	↓ 66%
		rilpivirine	↑ 30%	↑ 49%	↑ 76%
Fluconazole		EDURANT with azole ant			
Itraconazole		ma concentrations of rilpiv			P3A
Posaconazole		adjustment is required wh	en EDUR	ANT is	
Voriconazole	co-administered wit	h azole antifungal agents.			
ANTIMYCOBACTERIA	LS				
Rifabutin*	300 mg q.d.†	rifabutin	\leftrightarrow	\leftrightarrow	\leftrightarrow
		25-O-desacetyl-rifabutin	\leftrightarrow	\leftrightarrow	\leftrightarrow
	300 mg q.d.	rilpivirine (25 mg q.d.)	↓ 31%	↓ 42%	↓ 48%
	300 mg q.d.	rilpivirine (50 mg q.d.)	↑ 43%	↑ 16%	\leftrightarrow
	0.1	mg q.d. rilpivirine alone)			
	decreases in rilpivirir enzymes). This may Throughout co-admi dose should be incre	EDURANT with rifabutin m ne plasma concentrations (result in loss of therapeuti inistration of EDURANT with eased from 25 mg once da ration is stopped, the EDU once daily.	(induction c effect of th rifabutin ily to 50 m	of CYP3A EDURANT , the EDUF g once dai	RANT ly. When
Rifampicin*#	600 mg q.d.	rifampicin	\leftrightarrow	\leftrightarrow	NA
		25-desacetyl-rifampici n	\leftrightarrow	↓9%	NA
		rilpivirine	↓ 69%	↓ 80%	↓ 89%
Rifapentine	as co-administratior	not be used in combination n may cause significant de uction of CYP3A enzymes EDURANT.	creases ir	rilpivirine	plasma
MACROLIDE ANTIBIO	TICS				
Clarithromycin		EDURANT with clarithron			
Erythromycin		n the plasma concentratio Where possible, alternative ed.			

GLUCOCORTICOIDS

Co-administered medicinal product	Dose of co-administered medicinal product	Medicinal product assessed	C _{max}	AUC	C _{min}
Dexamethasone (systemic)	EDURANT should r dexamethasone as rilpivirine plasma co result in loss of ther	not be used in combination co-administration may ca oncentrations (induction on apeutic effect of EDURA arly for long-term use.	use signific f CYP3A er	ant decrea zymes). Tl	nis may
PROTON PUMP INHIB	ITORS				
Omeprazole*#	20 mg q.d.	omeprazole rilpivirine	↓ 14% ↓ 40%	↓ 14% ↓ 40%	NA ↓ 33%
Lansoprazole Rabeprazole Pantoprazole Esomeprazole	as co-administration	not be used in combination n may cause significant d tric pH increase). This m	ecreases in	rilpivirine p	olasma
H2-RECEPTOR ANTA	GONISTS				
Famotidine*#	40 mg single dose taken 12 hours before rilpivirine	rilpivirine	\leftrightarrow	↓ 9%	NA
	40 mg single dose taken 2 hours before rilpivirine	rilpivirine	↓ 85%	↓ 76%	NA
	40 mg single dose taken 4 hours after rilpivirine	rilpivirine	↑ 21%	↑ 13%	NA
Cimetidine Nizatidine Ranitidine	used with caution a rilpivirine plasma co	EDURANT and H2-recept s co-administration may of oncentrations (gastric pH only be administered at lo ANT.	cause signif increase). F	icant decre 12-receptor	ases in
ANTACIDS					
Antacids (e.g., aluminium or magnesium hydroxide, calcium carbonate)	as co-administration concentrations (gas	EDURANT and antacids n may cause significant d tric pH increase). Antacio irs before or at least 4 ho	ecreases in Is should or	rilpivirine p Ny be admi	olasma
NARCOTIC ANALGES	ICS				
Methadone*	60-100 mg q.d.,	R(-) methadone	↓ 14%	↓ 16%	↓ 22%
	individualised dose	S(+) methadone	↓ 13%	↓ 16%	↓21%
	methadone with ED	ts are required when initia URANT. However, clinica Itenance therapy may ne	al monitorin	g is recomi	mended
HERBAL PRODUCTS					
St John's wort (Hypericum perforatum)	St John's wort (<i>Hyp</i> significant decrease	not be used in combination ericum perforatum) as co es in rilpivirine plasma cou This may result in loss of	-administra	tion may ca (induction	ause
ANALGESICS					
Acetaminophen*# (paracetamol)	500 mg single dose	acetaminophen rilpivirine	$\leftrightarrow \\ \leftrightarrow$	$\leftrightarrow \\ \leftrightarrow$	NA ↑ 26%
	No dose adjustmen acetaminophen (pa	t is required when EDUR			
ESTROGEN-BASED C	ONTRACEPTIVES	•			
Ethinylestradiol*	0.035 mg q.d.	ethinylestradiol	↑ 17%	\leftrightarrow	\leftrightarrow
-		-			

Co-administered medicinal product	Dose of co-administered medicinal product	Medicinal product assessed	C _{max}	AUC	C _{min}
Norethindrone*	1 mg q.d.	norethindrone	\leftrightarrow	\leftrightarrow	\leftrightarrow
		t is required for the conco ogesterone-based contra		of EDURA	NT and
HMG CO-A REDUCTA	ASE INHIBITORS				
Atorvastatin*#	40 mg q.d.	atorvastatin	↑ 35%	\leftrightarrow	↓ 15%
		rilpivirine	↓ 9%	\leftrightarrow	\leftrightarrow
Fluvastatin Lovastatin Pitavastatin	No dose adjustmen an HMG Co-A redu	it is required when EDUR. ctase inhibitor.	ANT is co-a	dministere	∍d with
Pravastatin Rosuvastatin Simvastatin					
Pravastatin Rosuvastatin Simvastatin	SE TYPE 5 (PDE-5) IN	IHIBITOR			
Pravastatin Rosuvastatin Simvastatin	SE TYPE 5 (PDE-5) IN 50 mg single dose	IHIBITOR sildenafil	\leftrightarrow	\leftrightarrow	NA
Pravastatin Rosuvastatin Simvastatin PHOSPHODIESTERA	· /		\leftrightarrow	\leftrightarrow	NA ↔

* The interaction between EDURANT and the drug was evaluated in a clinical study. All other drug-drug interactions shown are predicted.

This interaction study has been performed with a dose higher than the recommended dose for EDURANT assessing the maximal effect on the co-administered drug. The dosing recommendation is applicable to the recommended dose of EDURANT 25 mg q.d.

QT prolonging drugs

There is limited information available on the potential for a pharmacodynamic interaction between rilpivirine and other medicinal products that prolong the QTc interval of the electrocardiogram. In a study of healthy subjects, supratherapeutic doses of rilpivirine (75 mg q.d. and 300 mg q.d.) have been shown to prolong the QTc interval of the electrocardiogram (see **section 5.1**). EDURANT at the recommended dose of 25 mg once daily is not associated with a clinically relevant effect on QTc. EDURANT should be used with caution when co-administered with a medicinal product with a known risk of Torsade de Pointes.

4.6 Fertility, pregnancy and lactation

Use in pregnancy

Category B1

There are no well controlled clinical or pharmacokinetic studies with EDURANT in pregnant women. Placental transfer of rilpivirine or its metabolites from dam to fetus was demonstrated in rats. Studies in animals have shown no evidence of relevant embryonic or foetal toxicity or an effect on reproductive function. There was no clinically relevant teratogenicity with rilpivirine in rats and rabbits. The exposures at the embryo foetal No Observed Adverse Effects Levels (NOAELs) in rats and rabbits were respectively 15 and 71 times higher than the exposure in humans at the recommended dose of 25 mg q.d.

To monitor maternal-fetal outcomes of pregnant women, an Antiretroviral Pregnancy Registry has been established (http://www.apregistry.com). This is a voluntary prospective, exposureregistration, observational study designed to collect and evaluate data on the outcomes of pregnancy exposures to antiretroviral products. For rilpivirine, sufficient first trimester exposures are available to allow detection of at least a two-fold increase in risk of overall birth defects. No such increases have been detected to date. Rilpivirine in combination with a background regimen was evaluated in a clinical trial of 19 pregnant women during the second and third trimesters, and postpartum. The pharmacokinetic data demonstrate that total exposure (AUC) to rilpivirine as a part of an antiretroviral regimen was approximately 30% lower during pregnancy compared with postpartum (6-12 weeks). Virologic response was preserved throughout the trial period. No mother to child transmission occurred in all 10 infants born to the mothers who completed the trial and for whom the HIV status was available. Rilpivirine was well tolerated during pregnancy and postpartum. There were no new safety findings compared with the known safety profile of rilpivirine in HIV-1 infected adults (see **section 5.2**).

EDURANT should be used during pregnancy only if the potential benefit justifies the potential risk.

Contraception in males and females

A trial to investigate the effect of EDURANT when co-administered with oral contraceptives demonstrated that EDURANT is unlikely to decrease the effectiveness of oral contraceptives. EDURANT and estrogen- and/or progesterone-based contraceptives can be used together without dose adjustments (see **section 4.5**).

Breast-feeding

It is not known whether rilpivirine is secreted in human milk. In nonclinical studies, rilpivirine was detected in the plasma of suckling rats following maternal dosing. Because of both the potential for HIV transmission and the potential for adverse events in nursing infants, mothers should be instructed not to breastfeed if they are receiving EDURANT.

Effects on fertility

No human data on the effect of rilpivirine on fertility are available (see section 5.3).

4.7 Effect on ability to drive and use machines

EDURANT has no or negligible influence on the ability to drive and use machines.

4.8 Undesirable effects

Adverse reactions from clinical trials

The safety assessment is based on the 96 week pooled data from 1368 patients in the Phase III controlled trials TMC278 C209 (ECHO) and TMC278 C215 (THRIVE) in antiretroviral treatment naïve HIV 1 infected adult patients, 686 of whom received EDURANT (25 mg q.d.) (see **Clinical trials**). The median duration of exposure for patients in the EDURANT and efavirenz arms was 104.3 and 104.1 weeks, respectively. Most adverse reactions (ARs) occurred in the first 48 weeks of treatment.

In the Phase III controlled trials ECHO and THRIVE through 96 weeks, the most frequently reported ARs ($\geq 2\%$) that were at least grade 2 in severity were depression, headache, insomnia, transaminases increased and rash (see **Table 3** for the complete list of ARs).

The majority of the ARs reported during treatment with EDURANT 25 mg once daily were grade 1 to 2 in severity. Grade 3 or 4 ARs were reported in 3.6% and 5.9% of the EDURANT and efavirenz treated patients, respectively. The most common (reported in more than 1 patient in the EDURANT arm) grade 3 or 4 ARs were transaminases increased (1.6% in the EDURANT arm and 2.9% in the efavirenz arm), depression (0.7% and 0.7% respectively), abdominal pain (0.4% and 0.1% respectively), dizziness (0.3% and 0.4% respectively) and rash (0.3% and 0.6% respectively). 1.7% of patients in the EDURANT arm discontinued treatment due to ARs compared to 4.0% of patients in the efavirenz arm. In the EDURANT arm, all ARs leading to discontinuation had an incidence < 0.5%. In the efavirenz arm, the most common ARs leading to discontinuation were rash (1.5%), transaminases increased (0.7%), depression (0.6%) and abnormal dreams (0.6%).

The most common ARs were identified in the system organ classes (SOC) of nervous system disorders (25.7% in the EDURANT arm and 42.8% in the efavirenz arm), psychiatric disorders

(23.6% in the EDURANT arm and 26.4% in the efavirenz arm) and gastrointestinal disorders (24.1% in the EDURANT arm and 22.1% in the efavirenz arm). The difference between EDURANT and the efavirenz arms observed in the SOC nervous system disorders was mainly due to the difference in dizziness experienced by patients.

ARs of at least moderate intensity (\geq grade 2) reported in adult patients treated with EDURANT are summarised in **Table 3**. The ARs are listed by system organ class (SOC) and frequency.

Table 3: ARs of at least moderate intenHIV-1 infected adult patients t			ral treatment-naïve		
System Organ Class (SOC) Adverse reaction, %	Pooled data from the week 96 analysis of the Phase III ECHO and THRIVE trials				
	EDURANT + BR N=686	efavirenz + BR N=682	Treatment Difference (95%Cl)		
Skin and subcutaneous tissue disorders	2.3%	9.5%	ND		
Rash* [#]	2.3%	9.5%	-7.2 (-9.7; -4.7)		
Psychiatric disorders	9.3%	9.7%	ND		
Depression	4.1%	3.2%	0.9 (-1.1; 2.8)		
Insomnia	3.5%	3.5%	0.0 (-2.0; 1.9)		
Abnormal dreams* [†]	1.6%	4.0%	-2.4 (-4.1; -0.6)		
Sleep disorders	1.3%	0.9%	0.4 (-0.7; 1.5)		
Depressed mood	0.4%	0.3%	0.1 (-0.5; 0.8)		
Nervous system disorders	4.5%	10.7%	ND		
Headache*	3.5%	3.8%	-0.3 (-2.3; 1.7)		
Dizziness*#	1.0%	6.7%	-5.7 (-7.7; -3.7)		
Somnolence	0.7%	1.3%	-0.6 (-1.7; 0.5)		
Gastrointestinal disorders	3.8%	5.6%	ND		
Abdominal pain	2.0%	1.9%	0.1 (-1.3; 1.6)		
Nausea*	1.3%	2.8%	-1.5 (-3.0; 0)		
Vomiting	1.0%	2.1%	-1.0 (-2.3; 0.3)		
Abdominal discomfort	0.4%	0.1%	0.3 (-0.3; 0.9)		
Metabolism and nutrition disorders	1.2%	0.6%	ND		
Decreased appetite	1.2%	0.6%	0.6 (-0.4; 1.6)		
General disorders and administration site conditions	1.6%	2.1%	ND		
Fatigue	1.6%	2.1%	-0.4 (-1.8; 0.9)		
Investigations	2.8%	4.0%	ND		
Transaminases increased	2.8%	4.0%	-1.2 (-3.1; 0.7)		

BR=background regimen; CI=confidence interval

N=total number of subjects per treatment group; ND=not determined.

* Treatment comparison was pre-specified for these ARs (Fisher's Exact Test)

† p-value < 0.01

[#] p-value < 0.0001

No new AR terms were identified in adult patients in the Phase III ECHO and THRIVE trials between 48 weeks and 96 weeks nor in the Phase IIb TMC278-C204 trial through 240 weeks.

Laboratory abnormalities

Selected treatment emergent clinical laboratory abnormalities (grade 3 or grade 4), reported in EDURANT-treated patients are shown in **Table 4**.

Table 4: Selected treatment emergent laboratory abnormalities (grade 3 or grade 4) observed in antiretroviral treatment-naïve HIV-1 infected adult patients							
Laboratory parameter abnormality, %	DAIDS toxicity range		oled data from the week 96 alysis of the Phase III ECHO and RIVE trials				
		EDURANT + BR N=686	efavirenz + BR N=682				
HEMATOLOGY							
Decreased hemoglobin	< 4.5 mmol/L < 74 g/L	0.1%	0.6%				
Decreased platelet count	< 49999/mm ³ < 49999 x 10 ⁹ /L	0.1%	0.3%				
Decreased white blood cell count	< 1499/mm ³ < 1.499 10 ⁹ /L	1.2%	1.0%				
BIOCHEMISTRY		·					
Increased creatinine	> 1.8 x ULN	0.1%	0.1%				
Increased AST	> 5.0 x ULN	2.3%	3.3%				
Increased ALT	> 5.0 x ULN	1.6%	3.7%				
Increased bilirubin	> 2.5 x ULN	0.7%	0.3%				
Increased pancreatic amylase	> 2 x ULN	3.8%	4.8%				
Increased lipase	> 3 x ULN	0.9%	1.6%				
Increased total cholesterol (fasted)*	> 7.77 mmol/L > 300 mg/dL	0.1%	3.3%				
Increased LDL cholesterol (fasted)*	≥ 4.91 mmol/L ≥ 191 mg/dL	1.5%	5.3%				
Increased Triglycerides (fasted)*	≥ 8.49 mmol/L ≥ 751 mg/dL	0.6%	3.3%				

BR=background regimen; ULN=upper limit of normal

N=number of subjects per treatment group

* $p \le 0.001$ according to Fisher's Exact test (difference in grade 3 plus 4 abnormalities between the two treatment groups).

Note: Percentages were calculated for the number of subjects with results for the analyte.

Adrenal function

In the pooled Phase III trials, at Week 96, the overall mean change from baseline in basal cortisol showed a decrease of -19.1 nmol/L in the EDURANT group, and an increase of +0.1 nmol/L in the efavirenz group. At Week 96, the mean change from baseline in ACTH-stimulated cortisol levels was lower in the EDURANT group (+18.4 \pm 8.36 nmol/L) than in the efavirenz group (+54.1 \pm 7.24 nmol/L). Mean values for both basal and ACTH-stimulated cortisol values at Week 96 were within the normal range. Overall, there were no serious adverse events, deaths, or treatment discontinuations that could clearly be attributed to adrenal insufficiency.

Serum creatinine

In the pooled Phase III trials, serum creatinine increased minimally over 96 weeks of treatment with EDURANT. Most of this increase occurred within the first four weeks of treatment with a mean change of 0.1 mg/dL (range: -0.3 mg/dL to 0.6 mg/dL) observed overall. In subjects who entered the trial with mild or moderate renal impairment, the serum creatinine increase observed was similar to that seen in subjects with normal renal function. These changes are not considered to be clinically relevant and no subject discontinued treatment due to increases in serum creatinine. Creatinine increases were comparable by background N(t)RTIs.

Serum lipids

Changes from baseline in total cholesterol, LDL-cholesterol, HDL-cholesterol and triglycerides are presented in **Table 5**. The mean changes from baseline were smaller in the EDURANT arm versus the efavirenz arm. The impact of such findings has not been demonstrated.

	Pooled data Trials	from the wee	ek 96 analysis	of the Phase	III ECHO and	d THRIVE	
	E	EDURANT + BR efavirenz + E N=686 N=682				BR	
	Baseline	Wee	ek 96	Baseline	Week 96		
Mean (95% CI)	Mean (mg/dL)	Mean (mg/dL)	Mean change* (mg/dL)	Mean (mg/dL)	Mean (mg/dL)	Mean change* (mg/dL)	
Total cholesterol (fasted) [†]	161	167	5	161	190	28	
HDL-cholesterol (fasted) [†]	41	46	4	40	51	11	
LDL-cholesterol (fasted) [†]	96	98	1	96	110	14	
Triglycerides (fasted) [†]	124	117	-7	133	148	12	

N=number of subjects per treatment group

* The change from baseline is the mean of within-patient changes from baseline for patients with both baseline and week 96 values.

† p-value < 0.001, Wilcoxon rank-sum test for treatment comparison of change from baseline

Immune reconstitution inflammatory syndrome

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 (immune reconstitution inflammatory syndrome) (see **section 4.4**). Autoimmune disorders such as Graves' disease and autoimmune hepatitis have also been reported in the context of immune reconstitution inflammatory syndrome (see **section 4.4**).

Additional information on special populations

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

In patients co-infected with hepatitis B or C virus receiving EDURANT, the incidence of hepatic enzyme elevation was higher than in patients receiving EDURANT who were not co-infected. This observation was the same in the efavirenz arm. The pharmacokinetic exposure of rilpivirine in co-infected patients was comparable to that in patients without co-infection.

Reporting of suspected adverse reactions

Reporting suspected adverse reactions after authorisation of the medicine is important. It

allows continued monitoring of the benefit/risk balance of the medicine. Healthcare professionals are asked to report any suspected adverse reactionsto https://pophealth.mv.site.com/carmreportnz/s/submit-adverse-event

4.9 Overdose

There is no specific antidote for overdose with EDURANT. Human experience of overdose with EDURANT is limited. Treatment of overdose with EDURANT consists of general supportive measures including monitoring of vital signs and ECG (QT interval) as well as observation of the clinical status of the patient. Since rilpivirine is highly bound to plasma protein, dialysis is unlikely to result in significant removal of the active substance.

For advice on the management of overdose please contact the National Poisons Centre on 0800 POISON (0800 764766).

5. PHARMACOLOGICAL PROPERTIES

5.1 Pharmacodynamic properties

Pharmacotherapeutic group: Antiviral for systemic use, non-nucleoside reverse transcriptase inhibitors, ATC code: J05AG05

Mechanism of action

Rilpivirine is a diarylpyrimidine NNRTI of HIV-1. Rilpivirine activity is mediated by non-competitive inhibition of HIV-1 reverse transcriptase (RT). Rilpivirine does not inhibit the human cellular DNA polymerases α , β and γ .

Antiviral activity in vitro

Rilpivirine exhibited activity against laboratory strains of wild type HIV-1 in an acutely infected T-cell line with a median EC_{50} value for HIV-1/IIIB of 0.73 nM (0.27 ng/mL). Although rilpivirine demonstrated limited *in vitro* activity against HIV-2 with EC_{50} values ranging from 2,510 to 10,830 nM (920 to 3,970 ng/mL), treatment of HIV-2 infection with EDURANTis not recommended in the absence of clinical data.

Rilpivirine also demonstrated antiviral activity against a broad panel of HIV-1 group M (subtype A, B, C, D, F, G, H) primary isolates with EC_{50} values ranging from 0.07 to 1.01 nM (0.03 to 0.37 ng/mL) and group O primary isolates with EC_{50} values ranging from 2.88 to 8.45 nM (1.06 to 3.10 ng/mL).

Rilpivirine showed additive antiviral activity in combination with the N(t)RTIs: abacavir, didanosine, emtricitabine, stavudine and tenofovir; the protease inhibitors: amprenavir, atazanavir, darunavir, indinavir, lopinavir, nelfinavir, ritonavir, saquinavir and tipranavir; the NNRTIs: efavirenz, etravirine and nevirapine; the fusion inhibitor enfuvirtide; and the entry inhibitor maraviroc. Rilpivirine shows additive to synergistic antiviral activity in combination with the NRTIs lamivudine and zidovudine, and the integrase inhibitor raltegravir.

Resistance

In cell culture

Rilpivirine-resistant strains were selected in cell culture starting from wild type HIV-1 of different origins and subtypes as well as NNRTI-resistant HIV-1. The most commonly observed amino acid substitutions that emerged included: L100I, K101E, V108I, E138K, V179F, Y181C, H221Y, F227C and M230I.

Resistance to rilpivirine was determined as a fold change in EC_{50} value (FC) above the biological cut-off (BCO) of the assay.

In treatment-naïve subjects

For the resistance analysis, a broader definition of virologic failure was used than in the primary efficacy analysis. In the week 48 pooled resistance analysis from the Phase III trials, 62 (of a total of 72) virologic failures in the EDURANT arm had resistance data at baseline and time of failure. In this analysis, the amino acid substitutions associated with NNRTI resistance that developed in at least 2 rilpivirine virologic failures were: V90I, K101E, E138K, E138Q, V179I, Y181C, V189I, H221Y and F227C. The most common mutations were the same in the week 48 and week 96 analyses. In the trials, the presence of the substitutions V90I and V189I, at baseline, did not affect response. The E138K substitution emerged most frequently during the rilpivirine treatment, commonly in combination with the M184I substitution.

More patients who failed virologically on EDURANT than who failed virologically on efavirenz developed lamivudine/emtricitabine associated resistance.

In the week 96 pooled resistance analysis, low rates of virologic failure, similar between the treatment arms, were observed from week 48 to week 96 (3.2% in the EDURANT arm and 2.3% in the efavirenz arm).

Considering all of the available *in vitro* and *in vivo* data, the following amino acid substitutions, when present at baseline, are likely to affect the activity of rilpivirine: K101E, K101P, E138A, E138G, E138K, E138Q, E138R, V179L, Y181C, Y181I, Y181V, Y188L, H221Y, F227C, M230I and M230L.

Cross-resistance

Site directed NNRTI mutant virus

In a panel of 67 HIV-1 recombinant laboratory strains with one amino acid substitution at RT positions associated with NNRTI resistance, including the most commonly found K103N and Y181C, rilpivirine showed antiviral activity against 64 (96%) of these strains. The single amino acid substitutions associated with a loss of susceptibility to rilpivirine were: K101P, Y181I and Y181V. The K103N substitution did not result in reduced susceptibility to rilpivirine by itself, but the combination of K103N and L100I resulted in a 7-fold reduced susceptibility to rilpivirine.

Recombinant clinical isolates

Rilpivirine retained sensitivity (FC \leq BCO) against 62% of 4786 HIV-1 recombinant clinical isolates resistant to efavirenz and/or nevirapine.

Treatment naïve HIV 1 infected patients

In the week 48 pooled analysis of the Phase III trials ECHO and THRIVE, 31 of the 62 subjects with virologic failure on EDURANT with phenotypic resistance data lost susceptibility to rilpivirine. Of these, 28 were resistant to etravirine, 27 to efavirenz, and 14 to nevirapine. These cross-resistance findings were confirmed in the week 96 pooled analyses of the Phase III clinical trials.

In the week 96 pooled analyses, among virologic failures in the EDURANT arm with baseline viral load \leq 100,000 copies/mL and with resistance to rilpivirine, there were fewer patients with phenotypic cross-resistance than among those in the EDURANT arm with baseline viral load > 100,000 copies/mL. 3, 4 and 1 rilpivirine virologic failures with baseline viral load \leq 100,000 copies/mL and with resistance to rilpivirine (N=5) had cross-resistance to efavirenz, etravirine and nevirapine, respectively, compared to 27, 28, and 15 rilpivirine virologic failures with baseline viral load > 100,000 copies/mL (N=30), respectively.

Effects on electrocardiogram

The effect of EDURANT at the recommended dose of 25 mg q.d. on the QTcF interval was evaluated in a randomised, placebo and active (moxifloxacin 400 mg once daily) controlled crossover study in 60 healthy adults, with 13 measurements over 24 hours at steady state. EDURANT at the recommended dose of 25 mg q.d. is not associated with a clinically relevant effect on QTc.

When supratherapeutic doses of 75 mg q.d. and 300 mg q.d. of EDURANT were studied in healthy adults, the maximum mean time matched (95% upper confidence bound) differences in QTcF interval from placebo after baseline correction were 10.7 (15.3) and 23.3 (28.4) ms, respectively. Steady state administration of EDURANT75 mg q.d. and 300 mg q.d. resulted in a mean C_{max} approximately 2.6 fold and 6.7 fold, respectively, higher than the mean steady state C_{max} observed with the recommended 25 mg q.d. dose of EDURANT.

Clinical trials

The evidence of efficacy of EDURANTis based on the analyses of 96 week data from 2 randomised, double-blinded, active-controlled, Phase III trials TMC278-C209 (ECHO) and TMC278-C215 (THRIVE). The trials were identical in design, with the exception of the background regimen (BR). At 96 weeks, the virologic response rate [confirmed undetectable viral load (< 50 HIV-1 RNA copies/mL)] according to the time to loss of virologic response (TLOVR) algorithm was evaluated in patients receiving EDURANT 25 mg q.d. in addition to a BR versus patients receiving efavirenz 600 mg q.d. in addition to a BR. The TLOVR imputation algorithm was used to define confirmed virologic response i.e., two consecutive viral load values below the threshold are needed to count as a response. Non-responders or failures were defined as those subjects who never responded i.e. never achieved 2 consecutive viral load values of < 50

copies/mL, or who were a rebounder (subject responded, then has two consecutive viral load values above the threshold value of 50 copies/mL), or discontinued prematurely.

Similar efficacy for EDURANT was seen in each trial demonstrating non-inferiority to efavirenz.

Antiretroviral treatment-naïve HIV-1 infected patients were enrolled who had a plasma HIV-1 RNA \geq 5000 copies/mL and were screened for susceptibility to N(t)RTIs and for absence of specific NNRTI RAMs. In ECHO, the BR was fixed to the N(t)RTIs, tenofovir disoproxil fumarate plus emtricitabine. In THRIVE, the BR consisted of 2 investigator-selected N(t)RTIs: tenofovir disoproxil fumarate plus emtricitabine <u>or</u> zidovudine plus lamivudine <u>or</u> abacavir plus lamivudine. In ECHO, randomisation was stratified by screening viral load. In THRIVE, randomisation was stratified by screening viral load.

This analysis included 690 patients in ECHO and 678 patients in THRIVE who had completed 96 weeks of treatment or discontinued earlier.

In the pooled analysis for ECHO and THRIVE, demographics and baseline characteristics were balanced between the EDURANT arm and the efavirenz arm. **Table 6** displays selected demographic and baseline disease characteristics of the patients in the EDURANT and efavirenz arms. 53.3% of patients in the EDURANT arm and 48.2% of patients in the efavirenz arm were in the \leq 100,000 copies/mL baseline viral load category. The proportion of patients with baseline viral load > 100,000 copies/mL was 46.7% and 51.8% in the EDURANT arm and efavirenz arm, respectively.

	Pooled data from the ECHO and THRIVE trials			
	EDURANT + BR	efavirenz + BR		
	N=686	N=682		
Demographic characteristics				
Median Age, years (range)	36	36		
	(18-78)	(19-69)		
Sex				
Male	76%	76%		
Female	24%	24%		
Race				
White	61%	60%		
Black/African American	24%	23%		
Asian	11%	14%		
Other	2%	2%		
Not allowed to ask per local regulations	1%	1%		
Baseline disease characteristics				
Median baseline plasma HIV-1 RNA	5.0	5.0		
(range), log ₁₀ copies/mL	(2-7)	(3-7)		
Median baseline plasma HIV-1 RNA	90,450.0	104,500.0		
(range), copies/mL	(156 – 20,800,000)	(1,010 – 4,550,000)		
Median baseline CD4+ cell count (range),	249	260		
x 10 ⁶ cells/l	(1-888)	(1-1137)		
Percentage of subjects with:				
hepatitis B/C virus co-infection	7.3%	9.5%		
Percentage of patientswith the following				
background regimens:				
tenofovir disoproxil fumarate plus	80.2%	80.1%		
emtricitabine				
zidovudine plus lamivudine	14.7%	15.1%		
abacavir plus lamivudine	5.1%	4.8%		

 Table 6: Demographic and baseline disease characteristics of antiretroviral treatment-naïve

 HIV-1 infected adult subjects in the ECHO and THRIVE trials (pooled analysis)

BR=background regimen

Table 7 below shows the efficacy results at 48 weeks and at 96 weeks for patients treated with

 EDURANT and patients treated with efavirenz from the pooled data from the ECHO and THRIVE

trials. The response rate (confirmed undetectable viral load < 50 HIV-1 RNA copies/mL) at week 96 was comparable between the EDURANT arm and the efavirenz arm. The incidence of virologic failure was higher in the EDURANT arm than the efavirenz arm at week 96; however, most of the virologic failures occurred within the first 48 weeks of treatment. Discontinuations due to adverse events were higher in the efavirenz arm at week 96 than the EDURANT arm. Most of these discontinuations occurred in the first 48 weeks of treatment.

Table 7: Virologic Outcome of Randomized Treatment in the ECHO and THRIVE Trials (Pooled Analysis at Week 48 (primary) and Week 96; ITT-TLOVR [*])								
	Outcome a	at Week 48	Outcome a	at Week 96				
%	EDURANT+ BR	efavirenz + BR	EDURANT + BR	efavirenz + BR				
	N=686	N=682	N=686	N=682				
Confirmed Undetectable Viral Load (< 50 HIV-1 RNA copies/mL) ^{§#}	84.3	82.3	77.6	77.6				
Virologic Failure [†]	9.0	4.8	11.5	5.9				
Death	0.1	0.4	0.1	0.9				
Discontinued due to adverse event (AE)	2.0	6.7	3.8	7.6				
Discontinued for non-AE reason [¶]	4.5	5.7	7.0	8.1				

N = number of subjects per treatment group

* intent-to-treat time to loss of virologic response

§ Subjects achieved virologic response (two consecutive viral loads < 50 copies/mL) and maintained it through week 48/96.

Predicted difference of response rates (95% CI) at week 48: 1.6% (-2.2%; 5.3%) and at week 96: -0.4% (-4.6%; 3.8%); both p-values < 0.0001 (non-inferiority at 12% margin) from logistic regression model, including stratification factors and study.

+ Includes subjects who were rebounder (confirmed viral load ≥ 50 copies/mL after being responder) or who were never suppressed (no confirmed viral load < 50 copies/mL, either ongoing or discontinued due to lack or loss of efficacy).

¶ e.g. lost to follow-up, non-compliance, withdrew consent

At week 96, the mean change from baseline in CD4+ cell count was +228 x 106 cells/l in the EDURANT arm and +219 x 106 cells/l in the efavirenz arm in the pooled analysis of the ECHO and THRIVE trials [estimated treatment difference (95% CI): 11.3 (-6.8; 29.4)].

A subgroup analysis of the virologic response (< 50 HIV-1 RNA copies/mL) at 48 and 96 weeks and virologic failure by baseline viral load, CD4 count and by background NRTIs (pooled data from the ECHO and THRIVE trials) is presented in **Table 8**.

Table 8: Virologic response (< 50 HIV-1 RNA copies/mL, ITT-TLOVR) and virologic failure by baseline viral load and by background NRTIs (Pooled analysis at Week 48 [primary] and Week 96 from the ECHO and THRIVE trials)								
Outcome at Week 48 Outcome at Week 96								
	EDURANT + BR Efavirenz + BR EDURANT + BR Efavire					irenz + BR N=682		
	Ν	n (%)	Ν	n (%)	Ν	n (%)	N	n (%)
Proportion of sub (copies/mL)	ojects	with HIV-1 RN	IA < 50	copies/mL at	week 4	8* by baseline	e plasm	a viral load
≤ 100,000	36 8	332 (90.2%)	330	276 (83.6%)	368	309 (84.0%)	329	263 (79.9%)
> 100,000	31 8	246 (77.4%)	352	285 (81.0%)	318	223 (70.1%)	353	266 (75.4%)
> 100,000 to ≤ 500,000	24 9	198 (79.5%)	270	223 (82.6%)	249	178 (71.5%)	270	205 (75.9%)

	Outcome at Week 48					Outcome at Week 96			
	EDURANT + BR N=686		Efavirenz + BR N=682		EDU	EDURANT + BR N=686		Efavirenz + BR N=682	
	Ν	n (%)	Ν	n (%)	Ν	n (%)	Ν	n (%)	
> 500,000	69	48 (69.6%)	82	62 (75.6%)	69	45 (65.2%)	83	61 (73.5%)	
Virologic Failure	[†] by ba	seline plasm	a viral lo	oad (copies/r	nL)				
≤ 100,000	36	14	330	11	368	21	329	12	
	8	(3.8%)		(3.3%)		(5.7%)		(3.6%)	
> 100,000	31	48	352	22	318	58	353	28	
	8	(15.1%)		(6.3%)		(18.2%)		(7.9%)	
> 100,000 to ≤	24	33	270	13	249	43	270	18	
500,000	9	(13.3%)		(4.8%)		(17.3%)		(6.7%)	
> 500,000	69	15	83	9	69	15	83	10	
		(21.7%)		(11.0%)		(21.7%)		(12.0%)	
Proportion of sul CD4 count (x 10 ⁶			IA < 50	copies/mL at	t week 4	8* and at wee	ek 96* by	/ baseline	
< 50	34	20	36	29	34	19	36	25	
		(58.8%)		(80.6%)		(55.9%)		(69.4%)	
≥ 50 - < 200	19	156	175	143	194	138	175	131	
	4	(80.4%)		(81.7%)		(71.1%)		(74.9%)	
≥ 200 - < 350	31	272	307	253	313	252	307	244	
	3	(86.9%)		(82.4%)		(80.5%)		(79.5%)	
≥ 350	14	130	164	136	144	123	164	129	
	4	(90.3%)		(82.9%)		(85.4%)		(78.7%)	
Virologic Failure	[†] by ba	seline CD4 co	ount (x '	10 ⁶ cells/l)					
<50	34	6	36	1	34	6	36	4	
		(17.6%)		(2.8%)		(17.6%)		(11.1%)	
≥ 50- <200	19	27	175	14	194	37	175	14	
	4	(13.9%)		(8.0%)		(19.1%)		(8.0%)	
≥ 200 - <350	31	21	307	14	313	26	307	15	
	3	(6.7%)		(4.6%)		(8.3%)		(4.9%)	
≥ 350	14	8	164	4	144	10	164	7	
	4	(5.6%)		(2.4%)		(6.9%)		(4.3%)	
Proportion of sub background N(t)		with HIV-1 RN	IA < 50	copies/mL at	t week 4	8* and at wee	ek 96* by	/	
tenofovir	550	459	546	450	550	423	546	422	
disoproxil		(83.5%)		(82.4%)		(76.9%)		(77.3%)	
fumarate plus									
emtricitabine									
zidovudine plus	101	88	103	83	101	82	103	79	
lamivudine		(87.1%)		(80.6%)		(81.2%)		(76.7%)	
abacavir plus	35	31	33	28	35	27	33	28	
lamivudine		(88.6%)		(84.8%)		(77.1%)		(84.8%)	

N=number of subjects per treatment group

n=number of observations

* Imputations according to the TLOVR algorithm.

+ Includes subjects who were rebounder (confirmed viral load ≥ 50 copies/mL after being responder) or who were never suppressed (no confirmed viral load < 50 copies/mL, either ongoing or discontinued due to lack or loss of efficacy).

At week 96, response rates (< 50 copies/mL [TLOVR]) in the pooled Phase III trial population were 84.0% in the EDURANT arm and 79.9% in the efavirenz arm in subjects with a baseline viral load \leq 100,000 copies/mL versus 70.1% and 75.4%, respectively, in subjects with a baseline viral load > 100,000 copies/mL. The proportion of virologic failures according to TLOVR in the pooled Phase III trial population was 5.7% in the EDURANT arm and 3.6% in the efavirenz arm, for subjects with a baseline viral load \leq 100,000 copies/mL. The proportion of virologic/mL. The proportion of virologic failures according to TLOVR in the pooled Phase III trial population was 5.7% in the EDURANT arm and 3.6% in the efavirenz arm, for subjects with a baseline viral load \leq 100,000 copies/mL. The proportion of virologic failures was higher for subjects with a baseline viral load > 100,000 copies/mL, especially in the EDURANT arm (18.2% EDURANT-treated subjects vs. 7.9% efavirenz-treated subjects).

The incidence of emergence of N(t)RTI and NNRTI RAMs in the virologic failures (according to the resistance analysis criteria) was lower in the \leq 100,000 copies/mL category than in the > 100,000 copies/mL category. This difference was observed in both treatment groups but with a lower incidence of emerging mutations in the efavirenz arm. This difference in incidence of emerging mutations between treatment groups was greater for N(t)RTI mutations. At week 48, among patients with baseline viral load \leq 100,000 copies/mL (16 patients in the rilpivirine arm and 12 patients in the efavirenz arm), 7 and 6 rilpivirine virologic failures and 2 and 5 efavirenz virologic failures had emerging N(t)RTI RAMs and NNRTI RAMs, respectively. Among patients in the efavirenz arm), 35 and 33 rilpivirine virologic failures and 7 and 10 efavirenz virologic failures had emerging N(t)RTI RAMs, respectively. These findings were confirmed in the week 96 pooled Phase III analyses.

Study TMC278-C204 was a randomised, active-controlled, Phase IIb trial in antiretroviral treatment-naïve HIV-1 infected adult patients consisting of 2 parts: an initial partially blinded dose-finding part [EDURANT doses blinded] up to 96 weeks, followed by a long-term, open label part. In the open label part of the trial, patients originally randomised to one of the 3 doses of EDURANT were all treated with EDURANT 25 mg once daily in addition to a BR, once the dose for the Phase III studies was selected. Patients in the control arm received efavirenz 600 mg once daily in addition to a BR in both parts of the study. The BR consisted of 2 investigator-selected N(t)RTIs: zidovudine plus lamivudine <u>or</u> tenofovir disoproxil fumarate plus emtricitabine.

Study TMC278-C204 enrolled 368 HIV-1 infected treatment-naïve adult patients who had a plasma HIV-1 RNA \geq 5000 copies/mL, previously received \leq 2 weeks of treatment with an N(t)RTI or protease inhibitor, had no prior use of NNRTIs, and were screened for susceptibility to N(t)RTI and for absence of specific NNRTI RAMs.

At 96 weeks, the proportion of patients with < 50 HIV-1 RNA copies/mL receiving EDURANT 25 mg (N=93) compared to patients receiving efavirenz (N=89) was 76% and 71%, respectively. The mean increase from baseline in CD4+ counts was 146 x 10^6 cells/l in patients receiving EDURANT 25 mg and 160 x 10^6 cells/l in patients receiving efavirenz.

Of those patients who were responders at week 96, 74% of patients receiving EDURANT remained with undetectable viral load (< 50 HIV-1 RNA copies/mL) at week 240 compared to 81% of patients receiving efavirenz. There were no safety concerns identified in the week 240 analyses.

Pregnancy

Rilpivirine in combination with a background regimen was evaluated in a clinical trial of 19 pregnant women during the second and third trimesters, and postpartum. The pharmacokinetic data demonstrate that total exposure (AUC) to rilpivirine as a part of an antiretroviral regimen was approximately 30% lower during pregnancy compared with postpartum (6-12 weeks). Virologic response was preserved throughout the trial period. No mother to child transmission occurred in all 10 infants born to the mothers who completed the trial and for whom the HIV status was available. Rilpivirine was well tolerated during pregnancy and postpartum. There were no new safety findings compared with the known safety profile of rilpivirine in HIV-1 infected adults.

5.2 Pharmacokinetic properties

The pharmacokinetic properties of rilpivirine have been evaluated in adult healthy subjects and in adult antiretroviral treatment naïve HIV 1 infected patients. Exposure to rilpivirine was generally lower in HIV-1 infected patients than in healthy subjects.

Absorption

After oral administration, the maximum plasma concentration of rilpivirine is generally achieved within 4-5 hours. The absolute bioavailability of EDURANT is unknown.

Effect of food on absorption

The exposure to rilpivirine was approximately 40% lower when EDURANT was taken in a fasted condition as compared to a normal caloric meal (533 kcal) or high fat high caloric meal (928 kcal). When EDURANT was taken with only a protein rich nutritional drink, exposures were 50% lower than when taken with a meal (see **section 4.2**).

Distribution

Rilpivirine is approximately 99.7% bound to plasma proteins *in vitro*, primarily to albumin. The distribution of rilpivirine into compartments other than plasma (e.g., cerebrospinal fluid, genital tract secretions) has not been evaluated in humans.

Metabolism

In vitro experiments indicate that rilpivirine primarily undergoes oxidative metabolism mediated by the cytochrome P450 (CYP) 3A system. It is possible that different populations of patients have faster or slower rilpivirine metabolism because of the various isoenzymes within the CYP3A system.

Elimination

The terminal elimination half life of rilpivirine is approximately 45 hours. After single dose oral administration of ¹⁴C rilpivirine, on average 85% and 6.1% of the radioactivity could be retrieved in faeces and urine, respectively. In faeces, unchanged rilpivirine accounted for on average 25% of the administered dose. Only trace amounts of unchanged rilpivirine (< 1% of dose) were detected in urine.

Additional information on special populations

Paediatric population

The pharmacokinetics of rilpivirine in paediatric patients (less than 18 years) have not been established.

Elderly

Population pharmacokinetic analysis in HIV infected patients showed that rilpivirine pharmacokinetics are not different across the age range (18 to 78 years, with only 2 patients aged above 65 years) evaluated. No dose adjustment of EDURANT is required in elderly patients.

Gender

Population pharmacokinetic analysis in HIV infected patients showed no clinically relevant differences in the pharmacokinetics of rilpivirine between men and women.

Race

Population pharmacokinetic analysis of rilpivirine in HIV infected patients indicated that race had no clinically relevant effect on the exposure to rilpivirine.

Hepatic impairment

Rilpivirine is primarily metabolised and eliminated by the liver. In a study comparing 8 patients with mild hepatic impairment (Child Pugh score A) to 8 matched controls, and 8 patients with moderate hepatic impairment (Child Pugh score B) to 8 matched controls. The mean steady-state exposure to rilpivirine was higher in subjects with mild hepatic impairment (27% higher for C_{max} and 47% higher for AUC) than in healthy controls. However, rilpivirine exposure in subjects with moderate hepatic impairment (5% lower for C_{max} and 5% higher for AUC) was similar to healthy controls. The mean apparent elimination half-life of rilpivirine was longer in subjects with mild (81 hours versus 61 hours respectively) and moderate (91 hours versus 56 hours, respectively) hepatic impairment compared to healthy controls. No dose adjustment is required in patients with

mild or moderate hepatic impairment. EDURANT has not been studied in patients with severe hepatic impairment (Child Pugh score C) (see **section 4.4**).

Renal impairment

The pharmacokinetics of rilpivirine have not been studied in patients with renal insufficiency. Renal elimination of rilpivirine is negligible. Therefore, the impact of renal impairment on rilpivirine elimination is expected to be minimal. As rilpivirine is highly bound to plasma proteins, it is unlikely that it will be significantly removed by haemodialysis or peritoneal dialysis.

Hepatitis B and/or hepatitis C virus co infection

Population pharmacokinetic analysis indicated that hepatitis B and/or C virus co infection had no clinically relevant effect on the exposure to rilpivirine.

Pregnancy and postpartum

The exposure to total rilpivirine after intake of rilpivirine 25 mg once daily as part of an antiretroviral regimen was lower during pregnancy (similar for the 2nd and 3rd trimester) compared with postpartum (see Table 9). The decrease in unbound (i.e., active) rilpivirine pharmacokinetic parameters during pregnancy compared to postpartum was less pronounced than for total rilpivirine.

In women receiving rilpivirine 25 mg once daily during the 2^{nd} trimester of pregnancy, mean intraindividual values for total rilpivirine C_{max} , AUC_{24h} and C_{min} values were, respectively, 21%, 29% and 35% lower as compared to postpartum; during the 3^{rd} trimester of pregnancy, C_{max} , AUC_{24h} and C_{min} values were, respectively, 20%, 31% and 42% lower as compared to postpartum.

Table 9:	Pharmacokinetic Results of Total Rilpivirine After Administration of Rilpivirine 25 mg	
Once Daily as Part of an Antiretroviral Regimen, During the 2 nd Trimester of Pregnancy, the 3 rd		
Trimester o	of Pregnancy and Postpartum	

$\begin{array}{l} \mbox{Pharmacokinetics of total} \\ \mbox{rilpivirine} \\ \mbox{(mean} \pm \mbox{SD}, \mbox{t}_{max} \mbox{: median [range])} \end{array}$	Postpartum (6-12 Weeks) (n=11)	2 nd Trimester of pregnancy (n=15)	3 rd Trimester of pregnancy (n=13)
C _{min} , ng/mL	84.0 ± 58.8	54.3 ± 25.8	52.9 ± 24.4
C _{max} , ng/mL	167 ± 101	121 ±45.9	123 ± 47.5
t _{max} , h	4.00 (2.03-25.08)	4.00 (1.00-9.00)	4.00 (2.00-24.93)
AUC _{24h} , ng.h/mL	2714 ± 1535	1792 ± 711	1762 ± 662

5.3 Preclinical safety data

Carcinogenicity

Rilpivirine was evaluated for carcinogenic potential by oral gavage administration to mice and rats up to 104 weeks. Daily doses of 20, 60 and 160 mg/kg/day were administered to mice and doses of 40, 200, 500 and 1500 mg/kg/day were administered to rats. An increase in the incidences of hepatocellular adenomas and carcinomas was observed in mice and rats. An increase in the incidences of follicular cell adenomas and/or carcinomas in the thyroid gland was observed in rats. Administration of rilpivirine did not cause a statistically significant increase in the incidence of any other benign or malignant neoplasm in mice or rats. The observed hepatocellular findings in mice and rats are considered to be rodent specific, associated with liver enzyme induction. A similar mechanism does not exist in humans; hence, these tumors are not relevant for humans. The follicular cell findings are considered to be rat specific, associated with increased clearance of thyroxine and are not considered to be relevant for humans. At the lowest tested doses in the carcinogenicity studies, the systemic exposures (based on AUC) to rilpivirine were 21 fold (mice) and 3 fold (rats), relative to those observed in humans at the recommended dose (25 mg q.d.).

Genotoxicity

Rilpivirine has tested negative in the *in vitro* Ames reverse mutation assay and *in vitro* clastogenicity mouse lymphoma assay, tested in the absence and presence of a metabolic activation system. Rilpivirine did not induce chromosomal damage in the *in vivo* micronucleus test in mice.

Fertility

In a study conducted in rats, there were no effects on mating or fertility with rilpivirine up to 400 mg/kg/day, a dose of rilpivirine that showed maternal toxicity. This dose is associated with an exposure that is approximately 40 times higher than the exposure in humans at the recommended dose of 25 mg q.d.

6. PHARMACEUTICAL PARTICULARS

6.1 List of excipients

Lactose monohydrate Croscarmellose sodium Povidone Polysorbate 20 Silicified microcrystalline cellulose (a combination of microcrystalline cellulose and silicon dioxide) Magnesium stearate Hypromellose Titanium dioxide Macrogol 3000 Triacetin

6.2 Incompatibilities

Not applicable

6.3 Shelf life

3 years.

6.4 Special precautions for storage

Store below 30°C. Store in the original bottle. Protect from light.

6.5 Nature and contents of container

EDURANT tablets are provided in a high density polyethylene (HDPE) bottle with a polypropylene (PP) child resistant closure and induction seal liner.

One bottle contains 30 tablets.

6.6 Special precautions for disposal and other handling

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

7. MEDICINE SCHEDULE

Prescription Medicine

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8. SPONSOR

Janssen-Cilag (New Zealand) Ltd Auckland, NEW ZEALAND Telephone: 0800 800 806 Email: <u>medinfo@janau.jnj.com</u>

9. DATE OF FIRST APPROVAL

26 January 2012

10. DATE OF REVISION OF THE TEXT

12 May 2025

Summary table of changes:

Section changes	Summary of new information	
4.2	Added paediatric age for clarity	
5.2	Added paediatric age for clarity	