ANNEX I SUMMARY OF PRODUCT CHARACTERISTICS

1. NAME OF THE MEDICINAL PRODUCT

Viekirax 12.5 mg/75 mg/50 mg film-coated tablets

2. QUALITATIVE AND QUANTITATIVE COMPOSITION

Each film-coated tablet contains 12.5 mg of ombitasvir, 75 mg of paritaprevir and 50 mg of ritonavir.

For the full list of excipients, see section 6.1.

3. PHARMACEUTICAL FORM

Film-coated tablet (tablet).

Pink, oblong, biconvex, film-coated tablets of dimensions 18.8 mm x 10.0 mm, debossed on one side with 'AV1'.

4. CLINICAL PARTICULARS

4.1 Therapeutic indications

Viekirax is indicated in combination with other medicinal products for the treatment of chronic hepatitis C (CHC) in adults (see sections 4.2, 4.4, and 5.1).

For hepatitis C virus (HCV) genotype specific activity, see sections 4.4 and 5.1.

4.2 Posology and method of administration

Treatment with Viekirax should be initiated and monitored by a physician experienced in the management of chronic hepatitis C.

Posology

The recommended oral dose of Viekirax is two 12.5 mg / 75 mg / 50 mg tablets once daily with food.

Viekirax should be used in combination with other medicinal products for the treatment of HCV (see Table 1).

Table 1. Recommended co-administered medicinal product(s) and treatment duration for Viekirax by patient population

Patient population	Treatment*	Duration	
Genotype 1b, without cirrhosis or with compensated cirrhosis	Viekirax + dasabuvir	8 weeks may be considered in previously untreated genotype 1b-infected patients with minimal to moderate fibrosis** (see section 5.1, GARNET study)	
Genotype 1a, without cirrhosis	Viekirax + dasabuvir + ribavirin*	12 weeks	
Genotype 1a, with compensated cirrhosis	Viekirax + dasabuvir + ribavirin*	24 weeks (see section 5.1.)	
Genotype 4, without cirrhosis or with compensated cirrhosis	Viekirax + ribavirin	12 weeks	

^{*}Note: Follow the genotype 1a dosing recommendations in patients with an unknown genotype 1 subtype or with mixed genotype 1 infection.

For specific dosage instructions for dasabuvir and ribavirin, including dose modification, refer to the respective Summaries of Product Characteristics.

Missed doses

In case a dose of Viekirax is missed, the prescribed dose can be taken within 12 hours. If more than 12 hours have passed since Viekirax is usually taken, the missed dose should NOT be taken and the patient should take the next dose per the usual dosing schedule. Patients should be instructed not to take a double dose.

Special populations

HIV-1 Co-infection

The dosing recommendations in Table 1 should be followed. For dosing recommendations with HIV antiviral medicinal products, refer to sections 4.4 and 4.5. See sections 4.8 and 5.1 for additional information.

Liver transplant recipients

Viekirax and dasabuvir in combination with ribavirin is recommended for 24 weeks in liver transplant recipients with genotype 1 HCV infection. Viekirax in combination with ribavirin is recommended in genotype 4 infection. Lower ribavirin dose at initiation may be appropriate. In the post-liver transplant study, ribavirin dosing was individualized and most subjects received 600 to 800 mg per day (see section 5.1). For dosing recommendations with calcineurin inhibitors see section 4.5.

Elderly

No dose adjustment of Viekirax is warranted in elderly patients (see section 5.2).

^{**} When assessing severity of liver disease using non-invasive methods, a combination of blood biomarkers or the combination of liver stiffness measurement and a blood test improves accuracy and should be undertaken prior to 8 week treatment in all patients with moderate fibrosis.

Renal impairment

No dose adjustment of Viekirax is required for patients with mild, moderate, or severe renal impairment, or end-stage-renal disease on dialysis (see section 5.2). For patients that require ribavirin, refer to the ribavirin Summary of Product Characteristics for information regarding use in patients with renal impairment.

Hepatic impairment

No dose adjustment of Viekirax is required in patients with mild hepatic impairment (Child-Pugh A). Viekirax is contraindicated in patients with moderate to severe hepatic impairment (Child-Pugh B or C) (see sections 4.3 and 5.2).

Paediatric population

The safety and efficacy of Viekirax in children less than 18 years of age have not been established. No data are available.

Method of administration

The film-coated tablets are for oral use. Patients should be instructed to swallow the tablets whole (i.e. patients should not chew, break or dissolve the tablet). To maximise absorption, Viekirax tablets should be taken with food, without regard to fat and calorie content (see section 5.2).

4.3 Contraindications

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

Patients with moderate to severe hepatic impairment (Child-Pugh B or C) (see section 5.2).

Use of ethinyloestradiol-containing medicinal products such as those contained in most combined oral contraceptives or contraceptive vaginal rings (see sections 4.4 and 4.5).

Medicinal products that are highly dependent on CYP3A for clearance and for which elevated plasma levels are associated with serious events must not be co-administered with Viekirax (see section 4.5). Examples are provided below.

CYP3A4 substrates:

- alfuzosin hydrochloride
- amiodarone, disopyramide, dronedarone, quinidine, ranolazine
- astemizole, terfenadine
- cisapride
- colchicine in patients with renal or hepatic impairment
- ergotamine, dihydroergotamine, ergonovine, methylergometrine
- fusidic acid
- lomitapide
- lovastatin, simvastatin, atorvastatin
- lurasidone
- oral midazolam, triazolam
- pimozide
- quetiapine
- salmeterol
- sildenafil (when used for the treatment of pulmonary arterial hypertension)
- ticagrelor

Co-administration of Viekirax with or without dasabuvir with medicinal products that are strong or moderate enzyme inducers is expected to decrease ombitasvir, paritaprevir, and ritonavir plasma concentrations and reduce their therapeutic effect and must not be co-administered (see section 4.5). Examples of contraindicated strong or moderate enzyme inducers are provided below.

Enzyme inducers:

- carbamazepine, phenytoin, phenobarbital
- efavirenz, nevirapine, etravirine
- apalutamide, enzalutamide
- mitotane
- rifampicin
- St. John's Wort (Hypericum perforatum)

Co-administration of Viekirax with or without dasabuvir with medicinal products that are strong inhibitors of CYP3A4 is expected to increase paritaprevir plasma concentrations and must not be co-administered with Viekirax (see section 4.5). Examples of contraindicated strong CYP3A4 inhibitors are provided below.

CYP3A4 inhibitors:

- cobicistat
- indinavir, lopinavir/ritonavir, saquinavir, tipranavir,
- itraconazole, ketoconazole, posaconazole, voriconazole
- clarithromycin, telithromycin
- conivaptan

4.4 Special warnings and precautions for use

General

Viekirax is not recommended for administration as monotherapy and must be used in combination with other medicinal products for the treatment of hepatitis C infection (see sections 4.2 and 5.1).

Risk of hepatic decompensation and hepatic failure in patients with cirrhosis

Hepatic decompensation and hepatic failure, including liver transplantation or fatal outcomes, have been reported postmarketing in patients treated with Viekirax with and without dasabuvir and with and without ribavirin. Most patients with these severe outcomes had evidence of advanced or decompensated cirrhosis prior to initiating therapy. Although causality is difficult to establish due to background advanced liver disease, a potential risk cannot be excluded.

Viekirax is contraindicated in patients with moderate to severe hepatic impairment (Child-Pugh B or C) (see sections 4.2, 4.3, 4.8 and 5.2).

For patients with cirrhosis:

- Monitoring should be performed for clinical signs and symptoms of hepatic decompensation (such as ascites, hepatic encephalopathy, variceal haemorrhage).
- Hepatic laboratory testing including direct bilirubin levels should be performed at baseline, during the first 4 weeks of starting treatment and as clinically indicated thereafter.
- Treatment should be discontinued in patients who develop evidence of hepatic decompensation.

ALT elevations

During clinical trials with Viekirax and dasabuvir with or without ribavirin, transient elevations of ALT to greater than 5 times the upper limit of normal occurred in approximately 1% of subjects (35 of 3,039). ALT elevations were asymptomatic and generally occurred during the first 4 weeks of treatment, without concomitant elevations of bilirubin, and declined within approximately two weeks of onset with continued dosing of Viekirax and dasabuvir with or without ribavirin.

These ALT elevations were significantly more frequent in the subgroup of subjects who were using ethinyloestradiol-containing medicinal products such as combined oral contraceptives or contraceptive vaginal rings (6 of 25 subjects); (see section 4.3). In contrast, the rate of ALT elevations in subjects using other types of oestrogens as typically used in hormonal replacement therapy (i.e., oral and topical oestradiol and conjugated oestrogens) was similar to the rate observed in subjects who were not using oestrogen-containing products (approximately 1% in each group).

Patients who are taking ethinyloestradiol-containing medicinal products (i.e. most combined oral contraceptives or contraceptive vaginal rings) must switch to an alternative method of contraception (e.g., progestin only contraception or non-hormonal methods) prior to initiating Viekirax with or without dasabuvir therapy (see sections 4.3 and 4.5).

Although ALT elevations associated with Viekirax and dasabuvir have been asymptomatic, patients should be instructed to watch for early warning signs of liver inflammation, such as fatigue, weakness, lack of appetite, nausea and vomiting, as well as later signs such as jaundice and discoloured faeces, and to consult a doctor without delay if such symptoms occur. Routine monitoring of liver enzymes is not necessary in patients that do not have cirrhosis (for cirrhotics, see above). Early discontinuation may result in drug resistance, but implications for future therapy are not known.

Pregnancy and concomitant use with ribavirin

Also see section 4.6.

Extreme caution must be taken to avoid pregnancy in female patients and female partners of male patients when Viekirax is taken in combination with ribavirin, see section 4.6 and refer to the Summary of Product Characteristics for ribavirin for additional information.

Use with tacrolimus, sirolimus and everolimus

Co-administration of Viekirax and dasabuvir with systemic tacrolimus, sirolimus or everolimus increases the concentrations of the immunosuppressant due to CYP3A inhibition by ritonavir (see section 4.5). Serious and/or life threatening events have been observed with co-administration of Viekirax and dasabuvir with systemic tacrolimus, and a similar risk can be expected with sirolimus and everolimus.

Avoid concomitant use of tacrolimus or sirolimus with Viekirax and dasabuvir unless the benefits outweigh the risks. If tacrolimus or sirolimus are used together with Viekirax and dasabuvir, caution is advised, and recommended doses and monitoring strategies can be found in section 4.5. Everolimus cannot be used due to lack of suitable dose strengths for dose adjustments.

Tacrolimus or sirolimus whole blood concentrations should be monitored upon initiation and throughout co-administration with Viekirax and dasabuvir and the dose and/or dosing frequency should be adjusted as needed. Patients should be monitored frequently for any changes in renal function or tacrolimus or

sirolimus associated adverse reactions. Refer to the tacrolimus or sirolimus Summary of Product Characteristics for additional dosing and monitoring instructions.

Genotype-specific activity

Concerning recommended regimens with different HCV genotypes, see section 4.2. Concerning genotype- specific virological and clinical activity, see section 5.1.

The efficacy of Viekirax has not been established in patients with HCV genotypes 2, 3, 5 and 6; therefore Viekirax should not be used to treat patients infected with these genotypes.

Co-administration with other direct-acting antivirals against HCV

Viekirax safety and efficacy have been established in combination with dasabuvir and/or ribavirin. Co-administration of Viekirax with other antivirals has not been studied and, therefore, cannot be recommended.

Retreatment

The efficacy of Viekirax in patients previously exposed to Viekirax, or to medicinal products of the same classes as those of Viekirax (NS3/4A inhibitors or NS5A inhibitors), has not been demonstrated. Concerning cross-resistance, see also section 5.1.

Use with glucocorticoids metabolised by CYP3A (e.g. fluticasone)

Caution should be used when administering Viekirax with fluticasone or other glucocorticoids that are metabolised by CYP3A4. Concomitant use of inhaled glucocorticoids metabolised with CYP3A can increase systemic exposures of the glucocorticoids, and cases of Cushing's syndrome and subsequent adrenal suppression have been reported with ritonavir-containing regimens. Concomitant use of Viekirax and glucocorticoids, particularly long-term use, should only be initiated if the potential benefit of treatment outweighs the risk of systemic corticosteroid effects (see section 4.5).

Use with colchicine

The interaction between Viekirax with or without dasabuvir and colchicine has not been evaluated. A reduction in colchicine dosage or an interruption of colchicine treatment is recommended in patients with normal renal or hepatic function if treatment with Viekirax with or without dasabuvir is required (see section 4.5). In patients with renal or hepatic impairment, use of colchicine with Viekirax with or without dasabuvir is contraindicated (see sections 4.3 and 4.5).

Use with statins

Simvastatin, lovastatin and atorvastatin are contraindicated (see sections 4.3 and 4.5).

Rosuvastatin

Viekirax with dasabuvir is expected to increase the exposure to rosuvastatin more than 3-fold. If rosuvastatin treatment is required during the treatment period, the maximum daily dose of rosuvastatin should be 5 mg (see section 4.5, Table 2). The increase in rosuvastatin when combined with Viekirax without dasabuvir is less pronounced. In this combination, the maximum daily dose of rosuvastatin should be 10 mg (see section 4.5, Table 2).

Pitavastatin and fluvastatin

The interactions between pitavastatin and fluvastatin and Viekirax have not been investigated. Theoretically, Viekirax with and without dasabuvir is expected to increase the exposure to pitavastatin and fluvastatin. A temporary suspension of pitavastatin/fluvastatin is recommended for the duration of treatment with Viekirax. If statin treatment is required during the treatment period, a switch to a reduced dose of pravastatin/rosuvastatin is possible (see section 4.5, Table 2).

Treatment of patients with HIV co-infection

Low dose ritonavir, which is part of the fixed dose combination Viekirax, may select for PI resistance in HIV co-infected patients without ongoing antiretroviral therapy. HIV co-infected patients without suppressive antiretroviral therapy should not be treated with Viekirax.

Drug interactions need to be carefully taken into account in the setting of HIV co-infection (for details see section 4.5, Table 2).

Atazanavir can be used in combination with Viekirax and dasabuvir, if administered at the same time. To be noted, atazanavir should be taken without ritonavir, since ritonavir 100 mg once daily is provided as part of Viekirax. The combination carries an increased risk for hyperbilirubinemia (including ocular icterus), in particular when ribavirin is part of the hepatitis C regimen.

Darunavir, dosed 800 mg once daily, if administered at the same time as Viekirax and dasabuvir, can be used in the absence of extensive PI resistance (darunavir exposure lowered). To be noted, darunavir should be taken without ritonavir, since ritonavir 100 mg once daily is provided as part of Viekirax.

HIV protease inhibitors other than atazanavir and darunavir (e.g., indinavir, saquinavir, tipranavir, lopinavir/ritonavir) are contraindicated (see section 4.3).

Raltegravir exposure is substantially increased (2-fold). The combination was not linked to any particular safety issues in a limited set of patients treated for 12-24 weeks.

Rilpivirine exposure is substantially increased (3-fold) when rilpivirine is given in combination with Viekirax and dasabuvir, with a consequent potential for QT-prolongation. If an HIV protease inhibitor is added (atazanavir, darunavir), rilpivirine exposure may increase even further and is, therefore, not recommended. Rilpivirine should be used cautiously, in the setting of repeated ECG monitoring.

NNRTIs other than rilpivirine (efavirenz, etravirine and nevirapine) are contraindicated (see section 4.3).

Hepatitis B virus reactivation

Cases of hepatitis B virus (HBV) reactivation, some of them fatal, have been reported during or after treatment with direct-acting antiviral medicinal products. HBV screening should be performed in all patients before initiation of treatment. HBV/HCV co-infected patients are at risk of HBV reactivation, and should, therefore, be monitored and managed according to current clinical guidelines.

Depression or psychiatric illness

Cases of depression and more rarely of suicidal ideation and suicide attempt have been reported with Viekirax with or without dasabuvir treatment in combination with ribavirin in the majority of the cases. Although some cases had previous history of depression, psychiatric illness and/or substance abuse, a causal relation with Viekirax with or without dasabuvir treatment cannot be excluded. Caution should be used in patients with a pre-existing history of depression or psychiatric illness. Patients and caregivers

should be instructed to notify the prescriber of any changes in behaviour or mood and of any suicidal ideation.

Use in diabetic patients

Diabetics may experience improved glucose control, potentially resulting in symptomatic hypoglycaemia, after initiating HCV direct acting antiviral treatment. Glucose levels of diabetic patients initiating direct acting antiviral therapy should be closely monitored, particularly within the first 3 months, and their diabetic medicinal products modified when necessary. The physician in charge of the diabetic care of the patient should be informed when direct acting antiviral therapy is initiated.

4.5 Interaction with other medicinal products and other forms of interaction

Viekirax may be administered with or without dasabuvir. When co-administered, they exert mutual effects on each other (see section 5.2). Therefore, the interaction profile of the compounds must be considered as a combination.

Pharmacodynamic interactions

Coadministration with enzyme inducers may increase the risk of adverse reactions and ALT elevations (see Table 2). Coadministration with ethinyloestradiol may increase the risk of ALT elevations (see sections 4.3 and 4.4). Examples of contraindicated enzyme inducers are provided in section 4.3.

Pharmacokinetic interactions

Potential for Viekirax to affect the pharmacokinetics of other medicinal products
In vivo drug interaction studies evaluated the net effect of the combination treatment, including ritonavir.

The following section describes the specific transporters and metabolizing enzymes that are affected by Viekirax with or without dasabuvir. See Table 2 for guidance regarding potential interactions with other medicinal products and dosing recommendations.

Medicinal products metabolised by CYP3A4

Ritonavir is a strong inhibitor of CYP3A. Co-administration of Viekirax with or without dasabuvir with medicinal products primarily metabolized by CYP3A may result in increased plasma concentrations of these medicinal products. Medicinal products that are highly dependent on CYP3A for clearance and for which elevated plasma levels are associated with serious events are contraindicated (see section 4.3 and Table 2).

CYP3A substrates evaluated in drug interaction studies which may require dose adjustment and/or clinical monitoring include (see Table 2) ciclosporin, sirolimus, tacrolimus, amlodipine, rilpivirine and alprazolam. Examples of other CYP3A4 substrates which may require dose adjustment and/or clinical monitoring include calcium channel blockers (e.g. nifedipine), and trazodone. Although buprenorphine and zolpidem are also metabolized by CYP3A, drug interaction studies indicate that no dose adjustment is needed when co-administering these medicinal products with Viekirax with or without dasabuvir (see Table 2).

Medicinal products transported by the OATP family and OCT1

Paritaprevir is an inhibitor of the hepatic uptake transporters OATP1B1 and OATP1B3, and paritaprevir and ritonavir are inhibitors of OATP2B1. Ritonavir is an *in vitro* inhibitor of OCT1, but the clinical relevance is unknown. Co-administration of Viekirax with or without dasabuvir with medicinal products

that are substrates of OATP1B1, OATP1B3, OATP2B1 or OCT1 may increase plasma concentrations of these transporter substrates, potentially requiring dose adjustment/clinical monitoring. Such medicinal products include some statins (see Table 2), fexofenadine, repaglinide and angiotensin II receptor antagonists (e.g., valsartan).

OATP1B1/3 substrates evaluated in drug interaction studies include pravastatin and rosuvastatin (see Table 2).

Medicinal products transported by BCRP

Paritaprevir, ritonavir and dasabuvir are inhibitors of BCRP *in vivo*. Co-administration of Viekirax with or without dasabuvir together with medicinal products that are substrates of BCRP may increase plasma concentrations of these transporter substrates, potentially requiring dose adjustment/clinical monitoring. Such medicinal products include sulfasalazine, imatinib and some of the statins (see Table 2).

BCRP substrates evaluated in drug interaction studies include rosuvastatin (see Table 2).

Medicinal products transported by P-gp in the intestine

While paritaprevir, ritonavir and dasabuvir are *in vitro* inhibitors of P-gp, no significant change was observed in the exposure of the P-gp substrate digoxin when administered with Viekirax and dasabuvir. However, co-administration of digoxin with Viekirax without dasabuvir may result in increased plasma concentrations (see Table 2). Viekirax may increase the plasma exposure to medicinal products that are sensitive for changed intestinal P-gp activity (such as dabigatran etexilate).

Medicinal products metabolised by glucuronidation (UGT1A1)

Paritaprevir, ombitasvir and dasabuvir are inhibitors of UGT1A1. Co-administration of Viekirax with or without dasabuvir with medicinal products that are primarily metabolized by UGT1A1 result in increased plasma concentrations of such medicinal products; routine clinical monitoring is recommended for narrow therapeutic index medicinal products (i.e. levothyroxine). See also Table 2 for specific advice on raltegravir and buprenorphine, which have been evaluated in drug interaction studies.

Medicinal products metabolised by CYP2C19

Co-administration of Viekirax with or without dasabuvir can decrease exposures of medicinal products that are metabolized by CYP2C19 (e.g. lansoprazole, esomeprazole, s-mephenytoin), which may require dose adjustment/clinical monitoring. CYP2C19 substrates evaluated in drug interaction studies include omeprazole and escitalopram (see Table 2).

Medicinal products metabolised by CYP2C9

Viekirax administered with or without dasabuvir did not affect the exposures of the CYP2C9 substrate, warfarin. Other CYP2C9 substrates (NSAIDs (e.g. ibuprofen), antidiabetics (e.g. glimepiride, glipizide) are not expected to require dose adjustments.

Medicinal products metabolised by CYP2D6 or CYP1A2

Viekirax administered with or without dasabuvir did not affect the exposures of the CYP2D6/CYP1A2 substrate, duloxetine. Exposures of cyclobenzaprine, a CYP1A2 substrate, were decreased. Clinical monitoring and dose adjustment may be needed for other CYP1A2 substrates (e.g. ciprofloxacin, cyclobenzaprine, theophylline and caffeine). CYP2D6 substrates (e.g. desipramine, metoprolol and dextromethorphan) are not expected to require dose adjustments.

Medicinal products renally excreted via transport proteins

Ombitasvir, paritaprevir, and ritonavir do not inhibit organic anion transporter (OAT1) *in vivo* as shown by the lack of interaction with tenofovir (OAT1 substrate). *In vitro* studies show that ombitasvir, paritaprevir, and ritonavir are not inhibitors of organic cation transporters (OCT2), organic anion

transporters (OAT3), or multidrug and toxin extrusion proteins (MATE1 and MATE2K) at clinically relevant concentrations.

Therefore, Viekirax with or without dasabuvir is not expected to affect medicinal products which are primarily excreted by the renal route via these transporters (see section 5.2).

Potential for other medicinal products to affect the pharmacokinetics of ombitasvir, paritaprevir, and dasabuvir

Medicinal products that inhibit CYP3A4

Co-administration of Viekirax with or without dasabuvir with strong inhibitors of CYP3A may increase paritaprevir concentrations (see section 4.3 and Table 2).

Enzyme inducers

Co-administration of Viekirax and dasabuvir with medicinal products that are moderate or strong enzyme inducers is expected to decrease ombitasvir, paritaprevir, ritonavir and dasabuvir plasma concentrations and reduce their therapeutic effect. Contraindicated enzyme inducers are provided in section 4.3 and Table 2.

Medicinal products that inhibit CYP3A4 and transport proteins

Paritaprevir is eliminated via CYP3A4 mediated metabolism and biliary excretion (substrate of the hepatic transporters OATP1B1, P-gp and BCRP). Caution is advised if co-administering Viekirax with medicinal products that are both moderate inhibitors of CYP3A4 and inhibitors of multiple transporters (P-gp, BCRP and/or OATP1B1/OATP1B3). These medicinal products may show clinically relevant increases in exposures of paritaprevir (e.g., ritonavir with atazanavir, erythromycin, diltiazem or verapamil).

Medicinal products that inhibit transport proteins

Potent inhibitors of P-gp, BCRP, OATP1B1 and/or OATP1B3 have the potential to increase the exposure to paritaprevir. Inhibition of these transporters is not expected to show clinically relevant increases in exposures of ombitasvir and dasabuvir.

Patients treated with vitamin K antagonists

As liver function may change during treatment with Viekirax administered with or without dasabuvir, a close monitoring of International Normalised Ratio (INR) values is recommended.

Drug interaction studies

Recommendations for co-administration of Viekirax with and without dasabuvir for a number of medicinal products are provided in Table 2.

If a patient is already taking medicinal product(s) or initiating a medicinal product while receiving Viekirax with or without dasabuvir for which potential for drug interaction is expected, dose adjustment of the concomitant medicinal product(s) or appropriate clinical monitoring should be considered (Table 2).

If dose adjustments of concomitant medicinal products are made due to treatment with Viekirax or Viekirax with dasabuvir, doses should be re-adjusted after administration of Viekirax or Viekirax with dasabuvir is completed.

Table 2 provides the Least Squares Means Ratio (90% Confidence Interval) effect on concentration of Viekirax with or without dasabuvir and concomitant medicinal products.

The magnitude of interaction when administered with medicinal products listed in Table 2 are similar (≤25% difference in the Least Square Means ratio) for Viekirax with or without dasabuvir, unless otherwise noted. Drug interactions were evaluated for the Viekirax and dasabuvir regimen, but not for the Viekirax without dasabuvir, with carbamazepine, furosemide, zolpidem, darunavir twice daily, darunavir (evening administration), atazanavir (evening administration), rilpivirine, abacavir/lamivudine, dolutegravir, metformin, sulfamethoxazole/trimethoprim, cyclobenzaprine, carisoprodol, hydrocodone/paracetamol or diazepam. Thus, for these medicinal products, results and dosing recommendations of the Viekirax and dasabuvir regimen can be extrapolated to Viekirax without dasabuvir.

The direction of the arrow indicates the direction of the change in exposures (C_{max} , and AUC) in paritaprevir, ombitasvir, dasabuvir and the co-administered medicinal product (\uparrow = *increase* (*more than* 20%), \downarrow = *decrease* (*of more than* 20%), \leftrightarrow = *no change* or change less than 20%). This is not an exclusive list.

Table 2. Interactions between Viekirax with or without dasabuvir and other medicinal products

Medicinal	GIVEN	EFFECT	Cmax	AUC	Ctrough	Clinical Comments
Product/Poss	WITH					
ible						
Mechanism						
of						
Interaction						
		PTOR ANTAG				T
Alfuzosin	Viekirax	Not studied. E	xpected			Concomitant use is
Mechanism:	with or					contraindicated (see section
CYP3A	without	↑ alfuzosin				4.3).
inhibition by	dasabuvir					
ritonavir						
AMINOSALIO Sulfasalazine	Viekirax	Not Studied. E	Synactad:			Caution should be used
Sulfasalazine	with or	Not Studied. I	expected.			when sulfasalazine is co-
36.1		↑ sulfasalazine	2			
Mechanism:	without dasabuvir					administered with Viekirax
BCRP	dasabuvir					with or without dasabuvir.
inhibition by						
paritaprevir,						•
ritonavir and						
dasabuvir.						
		OR BLOCKER				T
Valsartan	Viekirax	Not Studied. E	expected:			Clinical monitoring and
Losartan	with or	↑ valsartan				dose reduction is
Candesartan	without	† losartan				recommended for
	dasabuvir	† candesartan			angiotensin receptor	
Mechanism:		Candesarian		blockers when co-		
CYP3A4						administered with Viekirax
and/or						with or without dasabuvir.
OATP1B						
inhibition by						
paritaprevir.						
ANTIANGINA	\/ANTIARRY	YTHMICS				

Medicinal	GIVEN	EFFECT	Cmax	AUC	Ctrough	Clinical Comments
Product/Poss	WITH					
ible Mechanism						
of						
Interaction						
Amiodarone	Viekirax	Not studied. E	xpected:			Concomitant use is
Disopyramide	with or					contraindicated (see section
Dronedarone	without	↑ amiodarone	L			4.3).
Quinidine	dasabuvir	↑ disopyramid ↑ dronedarone				
Ranolazine		↑ quinidine				
Mechanism: CYP3A4		↑ ranolazine				
inhibition by						
ritonavir.						
Digoxin	Viekirax +	↔ digoxin	1.15	1.16	1.01	While no dose adjustment
	dasabuvir		(1.04-1.27)	(1.09-1.23)	(0.97-1.05)	is necessary for digoxin,
0.5 mg single		↔ ombitasvir	1.03 (0.97-1.10)	1.00 (0.98-1.03)	0.99 (0.96-1.02)	appropriate monitoring of
dose		↔	0.92	0.94	0.92	serum digoxin levels is
		paritaprevir	(0.80-1.06)	(0.81-1.08)	(0.82-1.02)	recommended.
Mechanism:		↔ dasabuvir	0.99	0.97	0.99	
P-gp inhibition by	x7' 1'	↑ 4::	(0.92-1.07) 1.58	(0.91-1.02) 1.36	(0.92-1.07)	D 1' ' 1 1
paritaprevir,	Viekirax without	↑ digoxin	(1.43-1.73)	(1.21-1.54)	(1.07-1.43)	Decrease digoxin dose by
ritonavir and	dasabuvir	\leftrightarrow		de of interaction		30-50%. Appropriate monitoring of serum
dasabuvir.	uasaouvii	ombitasvir		ed with Viekirax		digoxin levels is
		↔				recommended.
ANTIBIOTICS	 S (SVSTFMI)	paritaprevir	RATION)			
Clarithromycin	Viekirax	Not Studied. E				Concomitant use is
	with or		-			contraindicated (see
Telithromycin	without	↑ clarithromy				section 4.3).
	dasabuvir	↑ telithromyci	n			
Mechanism:		↑ paritaprevir				
CYP3A4/P- gp inhibition		↑ dasabuvir				
by						
clarithromyci						
n and						
ritonavir. Erythromycin	Viekirax	Not Studied. E	Expected:			Administration of Viekirax
Liyanomyem	with or	1.5. Stadion I	p			with or without dasabuvir
Mechanism:	without	↑ erythromyci	n			with erythromycin may
CYP3A4/P-	dasabuvir	^ ma=:t=== :				result in increased
gp inhibition		↑ paritaprevir ↑ dasabuvir				concentrations of
by		dasabuvii				erythromycin and
erythromycin,						paritaprevir. Caution is
paritaprevir, ritonavir and						advised.
dasabuvir.						
Fusidic Acid	Viekirax	Not studied. E	xpected:			Concomitant use is
1 dordie 1 tota	with or		1			contraindicated (see
	without	↑ fusidic acid				section 4.3).
	dasabuvir					

Medicinal	GIVEN	EFFECT	Cmax	AUC	Ctrough	Clinical Comments
Product/Poss	WITH		- 		o trough	
ible						
Mechanism						
of						
Interaction						
Mechanism:					1	
CYP3A4						
inhibition by						
ritonavir.						
Sulfameth-	Viekirax +	↑ Sulfameth-	1.21	1.17	1.15	No dose adjustment needed
oxazole,	dasabuvir	oxazole,	(1.15-1.28)	(1.14-1.20)	(1.10-1.20)	for Viekirax with or
Trimethoprim		↑ trimetho-	1.17	1.22	1.25	without dasabuvir.
		prim	(1.12-1.22)	(1.18-1.26)	(1.19-1.31)	
800/160 mg		↔	0.88	0.85	NA	
twice daily		ombitasvir	(0.83-0.94) 0.78	(0.80-0.90) 0.87	NA	
		↓ paritaprevir	(0.61-1.01)	(0.72-1.06)	INA	
Mechanism:		↑ dasabuvir	1.15	1.33	NA	
increase in			(1.02-1.31)	(1.23-1.44)		
dasabuvir	Viekirax		Not st			
possibly due	without	Similar effe		s observed with	Viekirax +	
to CYP2C8	dasabuvir		dasab	ouvir.		
inhibition by						
trimethoprim						
ANTICANCE						
Encorafenib	Viekirax	Not studied. E	xpected:			Co-administration may
34 1 .	with or					result in increased risk for
Mechanism: CYP3A4	without dasabuvir	↑ encorafenib				adverse events. Refer to the prescribing information
inhibition by	dasaouvii					of encorafenib for details
ritonavir.						on co-administration with
						strong CYP3A inhibitors.
Apalutamide	Viekirax	Not studied. E	xpected:			Concomitant use is
	with or					contraindicated (see
Enzalutamide	without dasabuvir	↓ombitasvir				section 4.3).
Mitotane	dasabuvir	↓ paritaprevir				
Willotane		↓ dasabuvir				
Mechanism:						
CYP3A4						
induction						
apalutamide						
enzalutamide or mitotane.						
Fostamatinib	Viekirax	Not Studied. E	Expected			Co-administration may
1 Ostalliatillio	with or	110t Studied. L	peetea		result in increased risk for	
Mechanism:	without	↑ fostamatinib		adverse events. Refer to		
CYP3A4	dasabuvir			the prescribing information		
inhibition by						of fostamatinib for details
ritonavir.						on co-administration with
						strong CYP3A inhibitors.
Ibrutinib	Viekirax	Not studied. E	xpected:			Co-administration may
	with or					result in increased risk for
		↑ ibrutinib				adverse events. Refer to
<u> </u>	1	<u> </u>				E. S.

Medicinal	GIVEN	EFFECT	Cmax	AUC	Ctrough	Clinical Comments
Product/Poss	WITH					
ible						
Mechanism						
of						
Interaction						
Mechanism:	without					the prescribing information
CYP3A4	dasabuvir					of ibrutinib for details on
inhibition by						co-administration with
ritonavir.						strong CYP3A inhibitors.
Imatinib	Viekirax	Not Studied. I	Expected:			Clinical monitoring and
34 1 .	with or	A				lower doses of imatinib are
Mechanism: BCRP	without dasabuvir	↑ imatinib				recommended.
inhibition by	dasabuvir					
paritaprevir,						
ritonavir and						
dasabuvir.						
ANTICOAGU		T	Γ		1	T
Warfarin	Viekirax +	↔ D	1.05	0.88	0.94	While no change to the
	dasabuvir	R-warfarin ↔	(0.95-1.17) 0.96	(0.81-0.95) 0.88	(0.84-1.05) 0.95	pharmacokinetics of
5 mg single		→ S-warfarin	(0.85-1.08)	(0.81-0.96)	(0.88-1.02)	warfarin is expected, close
dose and		5-warrariii	0.94	0.96	0.98	monitoring of INR is
other vitamin		ombitasvir	(0.89-1.00)	(0.93-1.00)	(0.95-1.02)	recommended with all
K antagonists		\leftrightarrow	0.98	1.07	0.96	vitamin K antagonists. This
		paritaprevir	(0.82-1.18)	(0.89-1.27)	(0.85-1.09)	is due to liver function
		↔	0.97	0.98	1.03	changes during treatment
	77' 1'	dasabuvir	(0.89-1.06)	(0.91-1.06)	(0.94-1.13)	with Viekirax \pm dasabuvir.
	Viekirax	↔ R-warfarin		de of interaction ed with Viekirax		
	without	K-wariariii ↔	to that observe	ed with vickinas	dasabuvii.	
	dasabuvir	S-warfarin				
		\leftrightarrow				
		paritaprevir				
		\leftrightarrow				
		ombitasvir			T	
Dabigatran	Viekirax	Not Studied. I	expected:			Viekirax without dasabuvir
etexilate	with or	↑ dabigatran e	texilate			may increase the plasma
Mechanism:	without	duoiganun c	textitute			concentrations of dabigatran
Intestinal P-	dasabuvir					etexilate. Use with caution.
gp inhibition						
by						
paritaprevir						
and ritonavir.	I CANTO	l				
ANTICONVU. Carbamaze-	Viekirax +	↔ carba-	1.10	1.17	1.35	Concomitant use is
	dasabuvir	→ carba- mazepine	(1.07-1.14)	(1.13-1.22)	(1.27-1.45)	contraindicated (see
pine	uasaouvir	↓ carbamaze	0.84	0.75	0.57	section 4.3).
200 ===================================		pine 10, 11-	(0.82-0.87)	(0.73-0.77)	(0.54-0.61)	, '
200 mg once		epoxide	ŕ			
daily followed by 200 mg		↓	0.69	0.69	NA	
twice daily		ombitasvir	(0.61-0.78)	(0.64-0.74)		
twice daily			0.34	0.30	NA	
		paritaprevir	(0.25-0.48)	(0.23-0.38)		

Medicinal Product/Poss	GIVEN WITH	EFFECT	Cmax	AUC	Ctrough	Clinical Comments
ible						
Mechanism						
of Interaction						
The action		Ţ	0.45	0.30	NA	
Mechanism:		dasabuvir	(0.41-0.50)	(0.28-0.33)		
CYP3A4	Viekirax	Not studied		expected as obs	erved with	
induction by	without		Viekirax +	dasabuvir.		
carbamazepine	dasabuvir					
Phenobarbital	Viekirax	Not Studied. I	Expected:			Concomitant use is contraindicated (see
Mechanism:	with or without	↓ ombitasvir				section 4.3).
CYP3A4	dasabuvir	↓ paritaprevir				
induction by	uasabuvii	↓ dasabuvir				
phenobarbital.		V				
Phenytoin	Viekirax	Not Studied. I	Expected:			Concomitant use is
Mechanism:	with or	1.1.				contraindicated (see section 4.3).
CYP3A4	without	↓ ombitasvir				section 4.5).
induction by	dasabuvir	↓ paritaprevir↓ dasabuvir				
phenytoin.		↓ dasaouvii				
S-	Viekirax	Not studied. E	Expected:			Clinical monitoring and
mephenytoin	with or		•			dose adjustment maybe
	without	↓ S-mepheny	toin			needed for s-mephenytoin.
Mechanism:	dasabuvir					
CYP2C19						
induction by						
ritonavir. ANTIDEPRES	CANTC					
Escitalopram	Viekirax +	↔ es-	1.00	0.87	NA	No dose adjustment is
-	dasabuvir	citalopram	(0.96-1.05)	(0.80-0.95)	1,11	necessary for escitalopram.
10 mg single		↑ S-	1.15	1.36	NA	
dose		Desmethyl	(1.10-1.21)	(1.03-1.80)		
		citalopram ↔	1.09	1.02	0.97	
		ombitasvir	(1.01-1.18)	(1.00-1.05)	(0.92-1.02)	
		\leftrightarrow	1.12	0.98	0.71	
		paritaprevir	(0.88-1.43)	(0.85-1.14)	(0.56-0.89)	
		↔ dasabuvir	1.10 (0.95-1.27)	1.01 (0.93-1.10)	0.89 (0.79-1.00)	
	Viekirax	↓ es-		e of interaction		
	without	citalopram		ed with Viekirax		
	dasabuvir			T	_	
		↔ S-	1.17	1.07	NA	
		Desmethyl citalopram	(1.08-1.26)	(1.01-1.13)		
		↔	The magnitu	de of interaction	was similar	
		ombitasvir		ed with Viekirax		
		↔				
Duloxetine	Viekirax +	paritaprevir	0.79	0.75	NA	No dose adjustment is
Duioactine	dasabuvir	duloxetine	(0.67-0.94)	(0.67-0.83)	11/1	necessary for duloxetine.

Medicinal Product/Poss ible Mechanism of	GIVEN WITH	EFFECT	Cmax	AUC	Ctrough	Clinical Comments
Interaction						
60 mg single		\leftrightarrow	0.98	1.00	1.01	
dose		ombitasvir	(0.88-1.08)	(0.95-1.06)	(0.96-1.06)	No dose adjustment needed
		\downarrow	0.79	0.83	0.77	for Viekirax with or
		paritaprevir	(0.53-1.16)	(0.62-1.10)	(0.65-0.91)	without dasabuvir.
		←	0.94	0.92	0.88	William dusus avii.
	Viekirax	dasabuvir ↔	(0.81-1.09)	(0.81-1.04) de of interaction	(0.76-1.01)	
	without	↔ duloxetine		ed with Viekirax		
	dasabuvir	↔		de of interaction		1
		ombitasvir		ed with Viekirax		
		\leftrightarrow	1.07	0.96	0.93	
		paritaprevir	(0.63-1.81)	(0.70-1.32)	(0.76-1.14)	
Trazodone	Viekirax	.				Trazodone should be used
Mechanism:	with or	Not studied. I	Expected:			with caution and a lower
CYP3A4	without	↑ trazodone				dose of trazodone may be
inhibition by	dasabuvir	trazodone				considered.
ritonavir.						
ANTI-DIURE	TIC HORMO	ONE				
Conivaptan Mechanism: CYP3A4/P- gp inhibition by conivaptan and	Viekirax with or without dasabuvir	Not studied. If †conivaptan † paritaprevir † dasabuvir	Expected:			Concomitant use is contraindicated (see section 4.3).
paritaprevir/ ritonavir/ombi tasvir						
ANTIFUNGAL	LS	1				
Ketoconazole 400 mg once	Viekirax with	↑ keto- conazole	1.15 (1.09-1.21)	2.17 (2.05-2.29)	NA	Concomitant use is contraindicated (see
daily	dasabuvir	↔ ombitasvir	0.98 (0.90-1.06)	1.17 (1.11-1.24)	NA	section 4.3).
Mechanism:		1	1.37	1.98	NA	
CYP3A4/P-		paritaprevir	(1.11-1.69)	(1.63-2.42)		
gp inhibition		 	1.16	1.42	NA	
by	Viekirax	dasabuvir ↑ keto-	(1.03-1.32)	(1.26-1.59) de of interaction	woo cimilan	-
ketoconazole	without	conazole		de of interaction ed with Viekirax		
and	dasabuvir	†		de of interaction		-
paritaprevir/		ombitasvir		ed with Viekirax		
ritonavir/		↑	1.72	2.16	NA	
ombitasvir		paritaprevir	(1.32-2.26)	(1.76-2.66)		

Medicinal Product/Poss	GIVEN WITH	EFFECT	Cmax	AUC	Ctrough	Clinical Comments
ible Mechanism of Interaction Itraconazole Posaconazole Mechanism: CYP3A4 and/or P-gp inhibition by itraconazole, posaconazole	Viekirax + dasabuvir Viekirax without dasabuvir	Not Studied. E † itraconazole † posaconazol † paritaprevir † dasabuvir	-			Concomitant use is contraindicated (see section 4.3).
and paritaprevir/ritonavir/ombi						
Voriconazole Mechanism: CYP2C19 induction and CYP3A4 inhibition by ritonavir	Viekirax with or without dasabuvir	Metabolisers: ↓ voriconazole ↑ paritaprevir ↑ dasabuvir	e xpected in CYI	P2C19 Extensive		Concomitant use is contraindicated (see section 4.3).
ANTI-GOUT		partapre vii				
Colchicine Mechanism: CYP3A4 inhibition by ritonavir.	Viekirax with or without dasabuvir	Not Studied. E	Expected:			A reduction in colchicine dosage or an interruption of colchicine treatment is recommended in patients with normal renal or hepatic function if treatment with Viekirax with or without dasabuvir is required. Use of colchicine is contraindicated with Viekirax with or without dasabuvir in patients with renal or hepatic impairment (see sections 4.3 and 4.4).
	WINES Viekirax	Not Studied T	Typoatod:			Concomitant use is
Astemizole Terfenadine Mechanism: CYP3A4	with or without dasabuvir	Not Studied. E	_			contraindicated (see section 4.3).

Medicinal Product/Poss	GIVEN WITH	EFFECT	Cmax	AUC	Ctrough	Clinical Comments
ible						
Mechanism						
of						
Interaction						<u> </u>
inhibition by						
ritonavir.			_			
Fexofenadine	Viekirax with or	Not Studied. F	•			Caution should be used when Viekirax with or
Mechanism:	without	↑ fexofenadin	e			without dasabuvir is
OATP1B1	dasabuvir					coadministered with
inhibition by						fexofenadine.
paritaprevir.						
ANTIHYPERI		CS	1.01	1.20	37.4	
Gemfibrozil	Paritaprevir/ ritonavir +	†	1.21 (0.94-1.57)	1.38 (1.18-1.61)	NA	Concomitant use of Viekirax with dasabuvir is
600 mg twice	dasabuvir	paritaprevir	2.01	11.25	NA	contraindicated (see
daily		↑ dasabuvir	(1.71-2.38)	(9.05-13.99)	NA	section 4.3).
Mechanism: Increase in	Viekirax		Not st	udied;		No dose adjustment of
dasabuvir	without	No interac		hen gemfibrozil	is used in	gemfibrozil is necessary.
exposure is	dasabuvir	combin	ation with Viek	irax without dasa	abuvir.	germierezh is necessary.
possibly due						No dose adjustment
to CYP2C8						needed for Viekirax.
inhibition and						100 000 101 (10111 011
increase in						
paritaprevir possibly due						
to OATP1B1						
inhibition by						
gemfibrozil.						
Lomitapide	Viekirax with or	Not Studied. H	Expected:			Concomitant use is
Mechanism:	without	1				contraindicated (see
CYP3A4	dasabuvir	lomitapide				section 4.3).
inhibition by						
ritonavir.						
ANTIMYCOB	ACTERIALS	<u>S</u>				
Rifampicin	Viekirax	Not Studied. I	Expected:			Concomitant use is
	with or					contraindicated (see section
Mechanism:	without	↓ ombitasvir				4.3).
CYP3A4	dasabuvir	↓ paritaprevir				
induction by		↓ dasabuvir				
rifampicin.						
ANTIPSYCH						
Lurasidone	Viekirax	Not Studied. E	Expected:			Concomitant use is
Pimozide	with or					contraindicated (see section
Quetiapine	without	↑ pimozide				4.3).
M1	dasabuvir	↑ quetiapine				
Mechanism:		↑ lurasidone				
CYP3A4						

Medicinal	GIVEN	EFFECT	Cmax	AUC	Ctrough	Clinical Comments
Product/Poss	WITH	EFFECT	Ciliax	1100	Ctrough	Chinear Comments
ible	,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,					
Mechanism						
of						
Interaction						
inhibition by						
ritonavir.						
ANTITPLATE						
Ticagrelor	Viekirax	Not studied. E	xpected:			Concomitant use is
Mechanism:	with or	↑ ticagrelor				contraindicated (see section 4.3).
CYP3A4	without	ticagicioi				T.3).
inhibition by	dasabuvir					
ritonavir						
BIGUANIDE		HYPERGLYC				
Metformin	Viekirax +	↓ metformin	0.77	0.90		No dose adjustment
	dasabuvir	1.74	(0.71-0.83)	(0.84-0.97)	NA 1.01	needed for metformin
500 mg single		↔ ombitasvir	0.92 (0.87-0.98)	1.01 (0.97-1.05)	(0.98-1.04)	when co-administered with
dose		↓ paritaprevir	0.63	0.80	1.22	Viekirax with and without
		↓ partapre (ii	(0.44-0.91)	(0.61-1.03)	(1.13-1.31)	dasabuvir.
		↔ dasabuvir	0.83	0.86	0.95	1
			(0.74-0.93)	(0.78 - 0.94)	(0.84-1.07)	
	Viekirax	g: 11 or	Not st		7. 1.	
	without dasabuvir	Similar eff	ect expected as dasab	observed with V	/iekirax +	
		O GLEED G	dasac	ouvir.		
CALCIUM CH	Viekirax +	OCKERS	1.26	2.57	1	Dagmaga amladining daga
Amlodipine	dasabuvir	amlodipine	(1.11-1.44)	(2.31-2.86)	NA	Decrease amlodipine dose by 50% and monitor
5 mg single	aususu vii	<u> </u>	1.00	1.00	1.00	patients for clinical effects.
dose		ombitasvir	(0.95-1.06)	(0.97-1.04)	(0.97-1.04)	
		\downarrow	0.77	0.78	0.88	
Mechanism:		paritaprevir	(0.64-0.94)	(0.68-0.88)	(0.80-0.95)	
CYP3A4		↔ dasabuvir	1.05 (0.97-1.14)	1.01 (0.96-1.06)	0.95 (0.89-1.01)	
inhibition by ritonavir.	Viekirax		Not sti		(0.89-1.01)	_
monavir.	without	Similar eff		observed with \	/iekirax +	
	dasabuvir		dasab			
Diltiazem	Viekirax	Not studied. E	xpected:			Caution is advised due to
Verapamil	with or					the expected increase in
	without	↑ diltiazem, ve	rapamil			paritaprevir exposures.
Mechanism:	dasabuvir					
CYP3A4/P-		↑ paritaprevir				Dose decrease and clinical
gp inhibition.		†/↔ dasabuvir	•			monitoring of calcium
						channel blockers is
						recommended when co-
						administered with Viekirax
						with and without
2710 11 1						dasabuvir.
Nifedipine	Viekirax	Not studied. E	xpected:			Dose decrease and clinical
36 1 '	with or	↑ nifedipine				monitoring of calcium
Mechaniem						channel blockers is
Mechanism: CYP3A4	without dasabuvir	Intersections				recommended when co-

Medicinal	GIVEN	EFFECT	Cmax	AUC	Ctrough	Clinical Comments
Product/Poss	WITH					
ible						
Mechanism						
of						
Interaction						1 ' ' . 1 ' . 1 77' 1 '
						administered with Viekirax
						with and without
CONTRACER						dasabuvir.
CONTRACEP	Viekirax	T	1.16	1.06	1.12	Editoria actualitat
Ethinyloestra diol/	with or	↔ ethinyloestra	(0.90-1.50)	(0.96-1.17)	(0.94-1.33)	Ethinyloestradiol- containing oral
norgestimate	without	diol	(0.50-1.50)	(0.50-1.17)	(0.74-1.55)	contraceptives are
8	dasabuvir		Norgestimate	metabolites:	I	contraindicated (see
0.035/0.25 mg		↑ norgestrel	2.26	2.54	2.93	section 4.3).
once daily			(1.91-2.67)	(2.09-3.09)	(2.39-3.57)	
•		↑ nor-	2.01	2.60	3.11	
Mechanism:		elgestromine	(1.77-2.29)	(2.30-2.95)	(2.51-3.85)	
possibly due		\leftrightarrow	1.05	0.97	1.00	
to UGT		ombitasvir	(0.81-1.35)	(0.81-1.15)	(0.88- 1.12)	
inhibition by		1	0.70	0.66	0.87	
paritaprevir, ombitasvir		paritaprevir	(0.40-1.21)	(0.42-1.04)	(0.67-1.14)	
and		↓ dasabuvir	0.51	0.48	0.53	
dasabuvir.		Ť	(0.22-1.18)	(0.23-1.02)	(0.30-	
					0.95)	
Nor-	Viekirax +	↔ nor-	0.83	0.91	0.85	No dose adjustment is
ethindrone	dasabuvir	ethindrone	(0.69-1.01)	(0.76-1.09)	(0.64-1.13)	necessary for
(progestin		↔	1.00	0.99	0.97	norethindrone or Viekirax
only pill) 0.35 mg once		ombitasvir	(0.93-1.08)	(0.94-1.04)	(0.90-1.03)	with or without dasabuvir.
daily		paritaprevir	(0.95-1.62)	(0.96-1.57)	(1.13-1.80)	
		↔ dasabuvir	1.01	0.96	0.95	
			(0.90-1.14)	(0.85-1.09)	(0.80-1.13)	
	Viekirax		Not st			
	without	Similar eff		observed with V	/iekirax +	
	dasabuvir		dasab	ouvir.		
DIURETICS	_	Ţ		.		,
Furosemide	Viekirax +		1.42	1.08	NA	Patients should be
	dasabuvir	furosemide	(1.17-1.72)	(1.00-1.17)	1.10	monitored for clinical
20 mg single		↔ ombitogvir	1.14	1.07	1.12	effects; a decrease in furosemide dose of up to
dose		ombitasvir ↔	(1.03-1.26)	(1.01-1.12) 0.92	(1.08-1.16)	50% may be required.
		paritaprevir	(0.63-1.36)	(0.70-1.21)	(1.16-1.38)	5070 may be required.
Mechanism:		→ dasabuvir	1.12	1.09	1.06	
possibly due			(0.96-1.31)	(0.96-1.23)	(0.98-1.14)	No dose adjustment
to UGT1A1	Viekirax		Not st			needed for Viekirax with
inhibition by	without	Similar eff		observed with V	/iekirax +	or without dasabuvir.
paritaprevir,	dasabuvir		dasab	ouvir.		
ombitasvir						
and						
dasabuvir.						
ERGOT ALK	ALOIDS					

Medicinal	GIVEN	EFFECT	Cmax	AUC	Ctrough	Clinical Comments
Product/Poss	WITH					
ible						
Mechanism						
of						
Interaction						
Ergotamine	Viekirax	Not studied. Ex	xpected:			Concomitant use is
Dihydroergot	with or					contraindicated (see section
amine	without	↑ ergot derivati	ives			4.3).
Ergonovine	dasabuvir					
Methylergom						
etrine						
Mechanism:						
CYP3A4						
inhibition by						
ritonavir.						
GLUCOCORT	TICOIDS (IN	HALED)				
Fluticasone	Viekirax	Not studied. Ex	xpected:			Concomitant use of
	with or					fluticasone can increase
Mechanism:	without	↑ fluticasone				systemic exposures of
CYP3A4	dasabuvir					fluticasone. Concomitant
inhibition by						use of Viekirax and
ritonavir.						fluticasone particularly
						long-term use, should only
						be initiated if the potential
						benefit of treatment
						outweighs the risk of
						systemic corticosteroid
						effects (see section 4.4).
GASTROINTI	ESTINAL PR	ODUCTS (PRO	OPULSIVE)			
Cisapride	Viekirax	Not studied. Ex	xpected:			Concomitant use is
Mechanism:	with or					contraindicated (see section
CYP3A4	without	↑ cisapride				4.3).
inhibition by	dasabuvir					
ritonavir.						
HCV ANTIVI	RALS					
Sofosbuvir	Viekirax +	↑ sofosbuvir	1.61	2.12	NA	No dose adjustment
	dasabuvir	<u> </u>	(1.38-1.88)	(1.91-2.37)		needed for sofosbuvir
400 mg once		↑ GS-331007	1.02	1.27	NA	when administered with
daily			(0.90-1.16)	(1.14-1.42)		Viekirax with or without
		↔ ombitasvir	0.93	0.93	0.92	dasabuvir.
Mechanism:			(0.84-1.03)	(0.87-0.99)	(0.88-0.96)	_
BCRP and P-		↔ paritaprevir	0.81 (0.65-1.01)	0.85 (0.71-1.01)	0.82 (0.67-1.01)	
gp inhibition		ornaprevir	1.09	1.02	0.85	-
by		· · dasaouvii	(0.98-1.22)	(0.95-1.10)	(0.76-0.95)	
paritaprevir,	Viekirax	The magnitud		was similar to t		1
ritonavir and	without		with Viekirax			
dasabuvir	dasabuvir					
HERBAL PR	ODUC19					

Medicinal	GIVEN	EFFECT	Cmax	AUC	Ctrough	Clinical Comments
Product/Poss	WITH					
ible						
Mechanism						
of						
Interaction						
St. John's Wort	St. John's Wort (hypericum		or without	Not studied. Expected:		Concomitant use is
perforatum)		dasabuvir				contraindicated (see section
,				↓ dasabuvir		4.3).
Mechanism:				↓ ombitasvir		
CYP3A4 induction by St.				↓ paritaprevir		
John's Wort						
					l.	

HIV ANTIVIRALS: PROTEASE INHIBITORS

For a general comment on treatment of HIV co-infected patients, including a discussion on different antiretroviral regimens that may be used, please see section 4.4 (Treatment of HIV co-infected patients).

dasabuvir das	Atazanavir	Viekirax +	\leftrightarrow	0.91	1.01	0.90	The recommended dose of
300 mg once daily (given at the same time) Mechanism: Increase in paritaprevir exposures may be due to inhibition of OATPIBI/B3 and CYP3A by atazanavir. ↓ ombitasvir 0.77 0.83 0.89 (0.74-0.94) (0.78-1.02) ↑ 1.46 1.94 3.26 ↑ 2.74 2.87 3.71 without ritonavir, in combination with Vickirax with dasabuvir. Ritonavir, in combination with Vickirax with dasabuvir. Ritonavir dose in Vickirax with dasabuvir. Ritonavir dose in Vickirax with dasabuvir. Treatment with atazanavir + Vickirax with dasabuvir. Treatment with atazanavir + Vickirax without dasabuvir is not recommended-(↑ paritaprevir). The combination of at the same time as Vickirax with dasabuvir. Treatment with atazanavir + Vickirax without dasabuvir is not recommended-(↑ paritaprevir). The combination of at the same time as Vickirax with dasabuvir. Treatment with atazanavir + Vickirax without dasabuvir is not recommended-(↑ paritaprevir). The combination of at the same time as Vickirax with dasabuvir. Treatment with atazanavir + Vickirax with dasabuvir is not recommended-(↑ paritaprevir). The combination with Vickirax with dasabuvir. Treatment with atazanavir + Vickirax with dasabuvir is not recommended-(↑ paritaprevir). The combination of attention was similar to that observed with Vickirax + dasabuvir.			atazanavir	(0.84-0.99)	(0.93-1.10)	(0.81-1.01)	
daily (given at the same time) Mechanism: Increase in paritaprevir exposures may be due to inhibition of OATP1B1/B3 and CYP3A by atazanavir. ↓ ombitasvir 0.77 (0.70-0.85) (0.74-0.94) (0.78-1.02) ↑ (0.70-0.85) (0.71-0.94) (0.78-1.02) ↑ (0.71-0.96) (0.71-0.94) (0.66-0.94) ↑ (0.71-0.94) (0.66-0.94) ↑ (0.71-0.96) ↑ (0.71-0.94) (0.66-0.94) ↑ (0.71-0.94) ↑ (0.66-0.94) ↑ (0.71-0.94) ↑ (0.66-0.94) ↑ (0.71-0.96) ↑ (0.71-0.94) ↑ (0.66-0.94) ↑ (0.71-0.96) ↑ (0.71-0.94) ↑ (0.66-0.94) ↑ (0.71-0.96) ↑ (0.71-0.94	300 mg once						<u> </u>
the same time) Mechanism: Increase in paritaprevir exposures may be due to inhibition of OATP1B1/B3 and CYP3A by atazanavir. ↓ ombitasvir 0.77 0.83 0.89 (0.74-0.94) (0.78-1.02) ↑ 1.46 1.94 3.26 paritaprevir (1.06-1.99) (1.34-2.81) (2.06-5.16) atazanavir and Viekirax + dasabuvir increase in paritaprevir exposures may be due to inhibition of OATP1B1/B3 and CYP3A by atazanavir. ↓ ombitasvir 0.77 0.83 0.89 (0.74-0.94) (0.78-1.02) ↑ 1.46 1.94 3.26 paritaprevir (1.06-1.99) (1.34-2.81) (2.06-5.16) atazanavir and Viekirax + dasabuvir increase bilirubin levels, in particular when ribavirin is part of the hepatitis C regimen (see sections 4.4 and 4.8).	_						
Mechanism: Increase in paritaprevir exposures may be due to inhibition of OATPIBL/B3 and CYP3A by atazanavir. Jombitasvir 0.77 0.83 0.89 (0.70-0.85) (0.74-0.94) (0.78-1.02)							
Mechanism: Increase in paritaprevir exposures may be due to inhibition of OATP1B1/B3 and CYP3A by atazanavir.							
Mechanism: Increase in paritaprevir exposures may be due to inhibition of OATP1B1/B3 and CYP3A by atazanavir. Jombitasvir 0.77 0.83 0.89 0.784.0.94) (0.78-1.02) 1.46 1.94 3.26 paritaprevir (1.06-1.99) (1.34-2.81) (2.06-5.16) 4 asabuvir increase bilirubin levels, in particular when ribavirin is part of the hepatitis C regimen (see sections 4.4 and 4.8).	time)						
Increase in paritaprevir exposures may be due to inhibition of OATP1B1/B3 and CYP3A by atazanavir. → ombitasvir 0.77 0.83 0.89 (0.74-0.94) (0.78-1.02) ↑	Mechanism:						
paritaprevir exposures may be due to inhibition of OATP1B1/B3 and CYP3A by atazanavir. ↓ ombitasvir							
exposures may be due to inhibition of OATP1B1/B3 and CYP3A by atazanavir.							
may be due to inhibition of OATP1B1/B3 and CYP3A by atazanavir.							•
inhibition of OATP1B1/B3 and CYP3A by atazanavir.	-						•
OATP1B1/B3 and CYP3A by atazanavir. No dose adjustment needed for Viekirax with dasabuvir. Treatment with atazanavir + Viekirax without dasabuvir is not recommended-(↑ paritaprevir (0.70-0.85) (0.74-0.94) (0.78-1.02) Treatment with atazanavir + Viekirax without dasabuvir is not recommended-(↑ paritaprevir). † 1.46	-						emaneement).
and CYP3A by atazanavir.							No dose adjustment
by atazanavir.							_
Treatment with atazanavir $+$ Viekirax without dasabuvir is not recommended-(\uparrow paritaprevir). \uparrow 1.46 1.94 3.26 paritaprevir (1.06-1.99) (1.34-2.81) (2.06-5.16) (2.06-5.16) (0.71-0.96) (0.71-0.94) (0.66-0.94) The combination of atazanavir and Viekirax $+$ dasabuvir increase bilirubin levels, in particular when ribavirin is part of the hepatitis C regimen (see sections 4.4 and 4.8). \leftrightarrow The magnitude of interaction was similar to that observed with Viekirax $+$ dasabuvir. \uparrow 2.74 2.87 3.71							
$ \begin{array}{ c c c c c c c c c c c c c c c c c c c$	by atazanavii.						dasabuvii.
$ \begin{array}{ c c c c c c c c c c c c c c c c c c c$							Treatment with atazanavir
dasabuvir is not recommended-(↑ paritaprevir). $ \uparrow 1.46 1.94 3.26 \\ paritaprevir (1.06-1.99) (1.34-2.81) (2.06-5.16) \\ \leftrightarrow dasabuvir 0.83 0.82 0.79 \\ (0.71-0.96) (0.71-0.94) (0.66-0.94) $ The combination of atazanavir and Viekirax + dasabuvir increase bilirubin levels, in particular when ribavirin is part of the hepatitis C regimen (see sections 4.4 and 4.8).							
→ ombitasvir 0.77 0.83 0.89 (0.70-0.85) (0.74-0.94) (0.78-1.02) ↑							
$ \begin{array}{c ccccccccccccccccccccccccccccccccccc$							
↑ 0.77 0.85 (0.78-1.02) ↑ 1.46 1.94 3.26 paritaprevir (1.06-1.99) (1.34-2.81) (2.06-5.16) ↑ dasabuvir 0.83 0.82 0.79 (0.71-0.96) (0.71-0.94) (0.66-0.94) ↑ The magnitude of interaction was similar atazanavir to that observed with Viekirax + dasabuvir. ↑ 2.74 2.87 3.71 The combination of atazanavir and Viekirax + dasabuvir increase bilirubin levels, in particular when ribavirin is part of the hepatitis C regimen (see sections 4.4 and 4.8).							· ·
$ \begin{array}{c ccccccccccccccccccccccccccccccccccc$			↓ ombitasvir				partaprevir).
paritaprevir $(1.06-1.99)$ $(1.34-2.81)$ $(2.06-5.16)$ atazanavir and Viekirax + dasabuvir increase bilirubin levels, in particular when ribavirin is part of the hepatitis C regimen (see sections 4.4 and 4.8).							The combination of
			· '	_	-		
(0.71-0.96) (0.71-0.94) (0.66-0.94) bilirubin levels, in particular when ribavirin is part of the hepatitis C regimen (see sections 4.4 and 4.8). The magnitude of interaction was similar to that observed with Viekirax + dasabuvir. 2.74 2.87 3.71							1
to that observed with Viekirax + dasabuvir. ↑ 2.74 2.87 3.71 → Shit data to Vois, in particular when ribavirin is part of the hepatitis C regimen (see sections 4.4 and 4.8).			, dasabavii				
to that observed with Viekirax + dasabuvir. The magnitude of interaction was similar atazanavir to that observed with Viekirax + dasabuvir. ↑ 2.74 2.87 3.71				(01,1 01,0)	(01,1 01,5 1,)	(0.00 0.5 1)	
The magnitude of interaction was similar atazanavir to that observed with Viekirax + dasabuvir. ↑ 2.74 2.87 3.71							•
The magnitude of interaction was similar atazanavir to that observed with Viekirax + dasabuvir. ↑ 2.74 2.87 3.71							*
← atazanavir							I =
atazanavir to that observed with Viekirax + dasabuvir. ↑ 2.74 2.87 3.71			\leftrightarrow	The magnitud	l de of interaction	was similar	una 7.0j.
↑ 2.74 2.87 3.71							
			↑				
			paritaprevir	(1.76-4.27)	(2.08-3.97)	(2.87-4.79)	

Medicinal Product/Poss ible Mechanism	GIVEN WITH	EFFECT	C _{max}	AUC	Ctrough	Clinical Comments
of						
Interaction	Viekirax without dasabuvir	↔ ombitasvir		l de of interaction ed with Viekirax		
Atazanavir/	Viekirax +	\leftrightarrow	1.02	1.19	1.68	
ritonavir	dasabuvir	atazanavir	(0.92-1.13)	(1.11-1.28)	(1.44-1.95)	
		↔ ombitasvir	0.83	0.90	1.00	
300/100 mg		†	(0.72-0.96)	(0.78-1.02)	(0.89-1.13)	
once daily (administered		paritaprevir	(1.61-2.98)	(2.40-4.17)	(8.94- 15.98)	
12 hours		↔ dasabuvir	0.81	0.81	0.80	
apart)	Viekirax		(0.73-0.91) Not st	(0.71-0.92)	(0.65-0.98)	
Mechanism:	without	Similar eff		observed with V	/iekirax +	
Increase in paritaprevir exposures may be due to inhibition of OATP1B1/B3 and CYP3A by atazanavir and CYP3A by the additional dose of ritonavir.	dasabuvir	↓ darunavir	dasab	0.76	0.52	
Darunavir	Viekirax + dasabuvir	↓ darunavn	(0.87-0.98)	(0.71-0.82)	(0.47-0.58)	The recommended dose of darunavir is 800 mg once
800 mg once		↔ ombitasvir	0.86 (0.77-0.95)	0.86 (0.79-0.94)	0.87 (0.82-0.92)	daily, without ritonavir,
daily (given at		†	1.54	1.29	1.30	when administered at the
the same		paritaprevir	(1.14-2.09)	(1.04-1.61)	(1.09-1.54)	same time as Viekirax +
time)		→ dasabuvir	1.10	0.94	0.90	dasabuvir (ritonavir dose in Viekirax will provide
Mechanism:	Viekirax	↔ darunavir	(0.88-1.37) 0.99	(0.78-1.14) 0. 92	(0.76-1.06) 0.74	darunavir pharmacokinetic
Unknown	without	- Garana II	(0.92-1.08)	(0.84-1.00)	(0.63-0.88)	enhancement). This regimen can be used in the
	dasabuvir	↔ ombitasvir	to that observe	de of interaction ed with Viekirax	+ dasabuvir. 1.85	absence of extensive PI resistance (i.e. lack of darunavir associated
		paritaprevir	(1.35-3.24)	(1.36-2.75)	(1.41-2.42)	RAMs), see also section 4.4.

Medicinal	GIVEN	EFFECT	Cmax	AUC	Ctrough	Clinical Comments
Product/Poss	WITH					
ible						
Mechanism						
of						
Interaction						27 1
 -						No dose adjustment
						needed for Viekirax with
						dasabuvir.
						Darunavir combined with
						Viekirax + dasabuvir is not
						recommended in patients
						with extensive PI
						resistance.
						T
						Treatment with darunavir + Viekirax without
						dasabuvir is not
						recommended-(†
						paritaprevir).
						paritapievii).
Darunavir/	Viekirax +	↔ darunavir	0.87	0.80	0.57	
ritonavir	dasabuvir		(0.79-0.96)	(0.74-0.86)	(0.48-0.67)	
		↓ ombitasvir	0.76	0.73	0.73	
600/100 mg		1	(0.65-0.88)	(0.66-0.80)	(0.64-0.83)	
twice daily		↓ paritaprevir	0.70 (0.43-1.12)	0.59 (0.44-0.79)	0.83 (0.69-1.01)	
 -		↓ dasabuvir	0.84	0.73	0.54	
Mechanism:		V dustie u i ii	(0.67-1.05)	(0.62-0.86)	(0.49-0.61)	
Unknown	Viekirax		Not st			
	without	Similar eff		observed with	Viekirax +	
	dasabuvir		dasat	ouvir.		
darunavir/	Viekirax +	↑ darunavir	0.79	1.34	0.54	
ritonavir	dasabuvir		(0.70-0.90)	(1.25-1.43)	(0.48-0.62)	
		↔	0.87	0.87	0.87	
800/100 mg		ombitasvir	(0.82-0.93)	(0.81-0.93)	(0.80-0.95)	
once daily		↓ paritaprevir	0.70 (0.50-0.99)	0.81 (0.60-1.09)	1.59 (1.23-2.05)	
		↓ dasabuvir	0.75	0.72	0.65	
(administered		↓ dababavii	(0.64-0.88)	(0.64-0.82)	(0.58-0.72)	
12 hours	Viekirax		Not st			
apart)	without	Similar eff		observed with	Viekirax +	
	dasabuvir		dasab	ouvir.		
Mechanism:						
Unknown						
Lopinavir /	Viekirax +	↔ lopinavir	0.87	0.94	1.15	Concomitant use is
ritonavir			(0.76 - 0.99)	(0.81-1.10)	(0.93-1.42)	contraindicated (see
	dasabuvir		`	1 17	1 0 4	,
l 	dasabuvir	↔ ombitosvir	1.14	1.17	1.24	section 4.3).
400/100 mg twice daily ¹	dasabuvir	↔ ombitasvir	`	1.17 (1.07-1.28) 2.17	1.24 (1.14-1.34) 2.36	,

Medicinal	GIVEN	EFFECT	Cmax	AUC	Ctrough	Clinical Comments
Product/Poss	WITH					
ible Mechanism						
of						
Interaction						
Interaction		\leftrightarrow	0.99	0.93	0.68	
Mechanism:		dasabuvir	(0.75-1.31)	(0.75-1.15)	(0.57 - 0.80)	
Increase in	Viekirax	↔ lopinavir		e of interaction		
paritaprevir	without			d with Viekirax		
exposures	dasabuvir	ombitasvir		e of interaction d with Viekirax		
may be due to		↑	4.76	6.10	12.33	
inhibition of		paritaprevir	(3.54-6.39)	(4.30-8.67)	(7.30-20.84)	
CYP3A/efflu						
x transporters						
by lopinavir						
and higher dose of						
ritonavir						
Indinavir	Viekirax	Not studied. E	Expected			Concomitant use is
Saquinavir	with or	110000000000000000000000000000000000000				contraindicated (see
Tipranavir	without	↑ paritaprevir				section 4.3).
1	dasabuvir					,
Mechanism:						
CYP3A4						
inhibition by						
protease						
inhibitors.						
HIV ANTIVIE			2.55	3.25	3.62	
Rilpivirine ²	Viekirax +	↑ rilpivirine	(2.08-3.12)	(2.80-3.77)	(3.12-4.21)	Co-administration of
25	dasabuvir	\leftrightarrow	1.11	1.09	1.05	Viekirax with rilpivirine once daily should only be
25 mg once daily		ombitasvir	(1.02-1.20)	(1.04-1.14)	(1.01-1.08)	considered in patients
administered		. 1	1.30	1.23	0.95	without known QT-
in the		paritaprevir	(0.94-1.81)	(0.93-1.64)	(0.84-1.07)	prolongation, and without
morning, with		↔ dasabuvir	1.18	1.17	1.10	other QT-prolongation co-
food		- Gusuouvii	(1.02-1.37)	(0.99-1.38)	(0.89-1.37)	medications. If the
\	Viekirax		Not st	udied:		combination is used,
Mechanism:	without	Similar eff	fect expected as	repeated ECG-monitoring		
CYP3A4	dasabuvir		dasal	ouvir.		should be done, see section
inhibition by						4.4. No dose adjustment
ritonavir.						needed for Viekirax with
Efavirenz/	Viekirax	Co-administ	ration of afovin	enz (enzyme ind	lucer) based	or without dasabuvir. Concomitant use with
emtricitabine/	with or			ritonavir + dasab		efavirenz is
tenofovir	without		ations and there	fore, early disco		contraindicated (see
disoproxil	dasabuvir		the s	tudy.		section 4.3).
fumarate						
600/300/200						
mg once daily						

Medicinal Product/Poss ible Mechanism of Interaction Mechanism: possible CYP3A4	GIVEN WITH	EFFECT	C _{max}	AUC	Ctrough	Clinical Comments			
induction by efavirenz. Nevirapine etravirine	Viekirax with or without dasabuvir	Not Studied. E ↓ ombitasvir ↓ paritaprevir ↓ dasabuvir				Concomitant use is contraindicated (see section 4.3).			
		GRASE STRAN			1				
Dolutegravir	Viekirax + dasabuvir	↑ dolutegravir	1.22 (1.15-1.29)	1.38 (1.30-1.47)	1.36 (1.19-1.55)	No dose adjustment needed for dolutegravir when coadministered with Viekirax with or without dasabuvir.			
50 mg once daily		↔ ombitasvir	0.96 (0.89-1.03)	0.95 (0.90-1.00)	0.92 (0.87-0.98)				
Mechanism: possibly due		↔ paritaprevir	0.89 (0.69-1.14)	0.84 (0.67-1.04)	0.66 (0.59-0.75)	ausuouvii			
to UGT1A1 inhibition by		↔ dasabuvir	1.01 (0.92-1.11)	0.98 (0.92-1.05)	0.92 (0.85-0.99)				
paritaprevir, dasabuvir and ombitasvir and CYP3A4 inhibition by ritonavir	Viekirax without dasabuvir	Similar eff	Not st ect expected as dasab	observed with V	Viekirax +				
Raltegravir	Viekirax + dasabuvir	↑ raltegravir	2.33 (1.66-3.27)	2.34 (1.70-3.24)	2.00 (1.17-3.42)	No dose adjustment is necessary for raltegravir or			
400 mg twice daily	uasaouvii	and ombita	relevant chang svir exposures (a) were observe	Viekirax with or without dasabuvir.					
Mechanism: Increase in	Viekirax without	† raltegravir	1.22 (0.78-1.89)	1.20 (0.74-1.95)	1.13 (0.51-2.51)				
raltegravir exposures may be due to UGT1A1 inhibition by paritaprevir, ombitasvir. and dasabuvir	dasabuvir	and ombita	svir exposures (es in dasabuvir, based on compaed during co-adr	rison with				

Medicinal	GIVEN	EFFECT	Cmax	AUC	Ctrough	Clinical Comments
Product/Poss	WITH					
ible						
Mechanism						
of						
Interaction						
HIV ANTIVIE	RALS: NUCL	EOSIDE INHI	BITORS		-	
Abacavir/	Viekirax +	↔ abacavir	0.87	0.94	NA	No dose adjustment
lamivudine	dasabuvir		(0.78 - 0.98)	(0.90-0.99)		needed for abacavir or
		1	0.78	0.88	1.29	lamivudine when co-
600/300 mg		↓ lamivudine	(0.72-0.84)	(0.82-0.93)	(1.05-1.58)	administered with Viekirax
once daily			` ′			with or without dasabuvir.
		↔	0.82	0.91	0.92	
		ombitasvir	(0.76-0.89)	(0.87-0.95)	(0.88-0.96)	4
		↔	0.84	0.82	0.73	
		paritaprevir	(0.69-1.02)	(0.70-0.97)	(0.63-0.85)	4
		↔ dasabuvir	0.94	0.91	0.95	
			(0.86-1.03)	(0.86-0.96)	(0.88-1.02)	
	Viekirax	G''1	Not st		7. 1-1	
	without	Similar en	ect expected as dasab	observed with V	v iekirax +	
	dasabuvir				1.00	
Em-	Viekirax +	↔ em-	1.05	1.07	1.09	No dose adjustment is
tricitabine/	dasabuvir	tricitabine	(1.00-1.12) 1.07	(1.00-1.14)	(1.01-1.17)	necessary for
tenofovir		↔ tenotovir	(0.93-1.24)	(1.07-1.20)	(1.13-1.36)	emtricitabine/tenofovir and
		\leftrightarrow	0.89	0.99	0.97	Viekirax with or without
200 mg once		ombitasvir	(0.81-0.97)	(0.93-1.05)	(0.90-1.04)	dasabuvir.
daily/300 mg		\	0.68	0.84	1.06	
once daily		paritaprevir	(0.42-1.11)	(0.59-1.17)	(0.83-1.35)	
		→ dasabuvir	0.85	0.85	0.85	
			(0.74-0.98)	(0.75-0.96)	(0.73-0.98)	
	Viekirax	↔ em-		de of interaction		
	without	tricitabine	0.80	ed with Viekirax	1.13	-
	dasabuvir	↔ tenotovir	(0.71-0.90)	(0.96-1.07)	(1.06-1.21)	
		\leftrightarrow	,	de of interaction	,	1
		ombitasvir	_	ed with Viekirax		
		\leftrightarrow	1.02	1.04	1.09	1
		paritaprevir	(0.63-1.64)	(0.74-1.47)	(0.88-1.35)	
HIV ANTIVIE	RALS: PHAR					
Cobicistat-	Viekirax	Not Studied. I				Concomitant use is
containing	with or		•			contraindicated (See section
regimens	without	↑ ombitasvir				4.3).
Mechanism:	dasabuvir	↑ paritaprevir				•
CYP3A4		↑ dasabuvir				
inhibition by						
cobicistat						
HMG CoA RE	DUCTASE I	NHIBITOR				
	Viekirax +	1	7.13	2.59	0.59	
	dasabuvir	rosuvastatin	(5.11-9.96)	(2.09-3.21)	(0.51-0.69)	
			0.02	0.00	0.00	4
		↔ ombitogvir	0.92	0.89	0.88	
	1	ombitasvir	(0.82-1.04)	(0.83-0.95)	(0.83-0.94)	

Medicinal	GIVEN	EFFECT	Cmax	AUC	Ctrough	Clinical Comments
Product/Poss ible	WITH					
Mechanism						
of						
Interaction		^	1.59	1.52	1.43	T1 1. 11 1
Rosuvastatin		paritaprevir	(1.13-2.23)	(1.23-1.90)	(1.22-1.68)	The maximum daily dose of rosuvastatin should be
5 mg once		↔ dasabuvir	1.07	1.08	1.15	5 mg (see section 4.4).
daily			(0.92-1.24)	(0.92-1.26)	(1.05-1.25)	
_						No dose adjustment
						needed for Viekirax with
Mechanism:						dasabuvir
OATP1B inhibition by						
paritaprevir	Viekirax	<u> </u>	2.61	1.33	0.65	The maximum daily dose
and BCRP	without	rosuvastatin	(2.01-3.39)	(1.14-1.56)	(0.57-0.74)	of rosuvastatin should be
inhibition by	dasabuvir		The	de of interaction		10 mg (see section 4.4).
paritaprevir,		↔ ombitasvir		de of interaction ed with Viekirax		
ritonavir or dasabuvir.						No dose adjustment
dasabuvir.		<u> </u>	1.40	1.22	1.06	needed for Viekirax.
		paritaprevir	(1.12-1.74)	(1.05-1.41)	(0.85-1.32)	
Pravastatin	Viekirax+	↑ pravastatin	1.37	1.82	NA	Reduce pravastatin dose by
	dasabuvir		(1.11-1.69)	(1.60-2.08)		50%.
10 mg once		↔ ombitasvir	0.95 (0.89-1.02)	0.89 (0.83-0.95)	0.94 (0.89-0.99)	
daily		↔ dasabuvir	1.00	0.96	1.03	No dose adjustment
			(0.87-1.14)	(0.85-1.09)	(0.91-1.15)	needed for Viekirax with or without dasabuvir.
Mechanism:		↔ paritaprevir	0.96 (0.69-1.32)	1.13 (0.92-1.38)	1.39 (1.21-1.59)	or without dasabuvir.
OATP1B1		paritapievii	(0.09-1.32)	(0.92-1.38)	(1.21-1.39)	
inhibition by	Viekirax	↑ pravastatin		de of interaction		
paritaprevir.	without			ed with Viekirax de of interaction		
	dasabuvir	ombitasvir	_	ed with Viekirax		
		1	1.44	1.33	1.28	
THE STATE OF THE S	T7' 1'	paritaprevir	(1.15-1.81)	(1.09-1.62)	(0.83-1.96)	
Fluvastatin	Viekirax with or	Not studied. E	xpected:			Concomitant use with fluvastatin and pitavastatin
Mechanism:	without	↑ fluvastatin				is not recommended (see
OATP1B/BC	dasabuvir	113,4544111				section 4.4).
RP inhibition		↑ pitavastatin				A temporary suspension of
by						fluvastatin and pitavastatin
paritaprevir						is recommended for the duration of treatment with
Pitavastatin						Viekirax. If statin
Mechanism:						treatment is required
OATP1B						during the treatment
inhibition by						period, a switch to dose
paritaprevir						reduced pravastatin or
	1					rosuvastatin is possible.

Product/Poss ible Mechanism of Interaction Lovastatin Simvastatin With or WITH Concomitan contraindica	
Mechanism of Interaction Not studied. Expected: Concomitan contraindical contraindical	
Interaction Lovastatin Viekirax Not studied. Expected: Concomitan contraindical contrai	
Lovastatin Viekirax Not studied. Expected: Concomitan contraindica	
Simvastatin with or contraindical	
	`
atorvastatin without dasabuvir 1 lovastatin, simvastatin, atorvastatin section 4.3).	•
Mechanism:	
CYP3A4/OA	
TP1B inhibition	
IMMUNOSUPPRESSANTS	
Ciclosporin Viekirax + ↑ 1.01 5.82 15.8 When starting	າດ ເດ-
dasabuvir ciclosporin (0.85-1.20) (4.73-7.14) (13.8- administration of the ciclosporin dasabuvir ciclosporin (0.85-1.20) (4.73-7.14)	
30 mg once 18.09) Viekirax, gi	ve one fifth of
doily single \leftrightarrow 0.99 1.08 1.15 the total dail	
ombitasvir (0.92-1.07) (1.03-1.11) (1.08-1.23) etclosporii (once daily with
1.44 1.72 1.85 VIEKIIAX. IVI	
	and/or dosing
Effect on (0.58-0.75) (0.65-0.76) (0.71-0.82) frequency as	
ciclosporin is Viekirax \(\) \(\) \(0.83 \) \(4.28 \) \(12.8 \)	
due to without ciclosporin (0.72-0.94) (3.66-5.01) (10.6-15.6) No dose adj	
CYP3A4 I dasabiivir	Viekirax with
inhibition by ombitasvir to that observed with Viekirax + dasabuvir. or without d	or without dasaouvii.
ritonavir and \uparrow 1.39 1.46 1.18	
increase in paritaprevir (1.10-1.75) (1.29-1.64) (1.08-1.30)	
paritaprevir	
exposures	
may be due to	
OATP/BCRP/	
P-gp	
inhibition by	
ciclosporin.	
Everolimus Viekirax + ↑ 4.74 27.1 16.1 Co-administ dasabuvir everolimus (4.29-5.25) (24.5-30.1) (14.5- Viekirax with the view of the view	
17 0)4 VEATER WI	th everolimus
0.75 mg 1.02 1.02 1.02 1.02	
single dose 0.99 1.02 1.02 because of a increase in 0.99 1.02 1.02 0.99 -1.06	_
\leftrightarrow 1.22 1.26 1.06	hich cannot be
paritapievii (1.05-1.45) (1.07-1.45) (0.97-1.10)	
Effect on everolimus is \rightarrow 1.03 1.08 1.14 properly dos dasabuvir (0.90-1.18) (0.98-1.20) (1.05-1.23) with availab	
0.00 1.10 (0.00 1.20) (1.00 1.20)	ee section 4.4).
CYP3A4 without Similar effect is expected as observed with Viekirax +	
inhibition by dasabuvir dasabuvir.	
ritonavir	
Sirolimus Viekirax + ↑ sirolimus 6.40 38.0 19.6 Concomitan	t use of
dasabuvir (5.34-7.68) (31.5-45.8) (16.7- sirolimus wi	
$(22.9)^6$ and dasabuv	

ible Mechanism of Interaction 0.5 mg single dose² Mechanism: Effect on sirolimus is due to CYP3A4 inhibition by ritonavir Tacrolimus 2 mg single dose² 2 mg single dose² 2 mg single dose² 3 mg single dose² 4 machanism: Effect on sirolimus by ritonavir Tacrolimus 5 mg single dose² 4 mg single dose² 5 mg single dose² 2 mg single dose² 2 mg single dose² 3 mg single dose² 4 mchanism: Effect on sirolimus is due to CYP3A4 inhibition by ritonavir Tacrolimus 5 mg single dose² 5 mg single dose² 7 mg single dose² 5 mg single dose² 7 mg single dose² 7 mg single dose² 7 mg single dose² 7 mg single dose² 8 mg single dose² 1 mg single dose² 1 mg single dose² 1 mg single dose² 2 mg single dose² 2 mg single dose² 3 mg single dose² 4 mg single dose² 5 mg single dose² 6 mibitavir 6 mibitavir 6 mg single dose² 7 mg single dose² 7 mg single dose² 8 mg single dose² 9 mibitavir 1 mg single dose² 2 mg single dose² 2 mg single dose² 3 mg single dose² 4 mg single dose² 1 mg single dose² 2 mg single dose² 3 mg single dose² 4 mg single dose² 1 mg single dose² 2 mg single dose² 3 mg single dose² 4 mg single dose² 4 mg single dose² 5 days after completion of Vickirax + dasabuvir rearment, the sirolimus dose and dosing frequency should be rearmed, along with routine monitoring of sirolimus should be rearmed, along with routine monitoring of sirolimus should be rearmed, along with routine monitoring of sirolimus should be rearmed, along with routine monitoring of sirolimus should should be rearmed, along with routine monitoring of sirolimus should should be rearmed, along with routine monitoring of sirolimus should should be rearmed, along with routine monitoring of sirolimus should shou	Medicinal Product/Poss	GIVEN WITH	EFFECT	C _{max}	AUC	Ctrough	Clinical Comments
Tacrolimus Cyp3Ad Cyp3A		WIII					
Tacrolimus Cyrg3Ar Concomination Conc							
0.5 mg single dose ² 0.5 mg single dose ³ 0.6 mg single dose ³ 0.7 mg single dose ³ 0.8 mg single dose ³ 0.8 mg single dose ³ 0.9	_						
dose ³				1.02	1.02	1.05	1.1.1.1
Mechanism: Effect on sirolimus is due to CYP3A4 without ritonavir Tacrolimus Tacrolimus Z mg single dose² Z mg single dose							
Mechanism: Effect on sirolimus is due to CYP3A4 or inhibition by ritonavir Tacrolimus	dose					<u> </u>	benefits outweigh the risks
Effect on sirolimus is due to CYP3A4 inhibition by ritonavir and sabuvir and sirolimus is used together with Vickirax + dasabuvir, administer sirolimus or vithout dasabuvir and vithout dos. And vithout dos. And vithout dasabuvir and vithout dos. And vithout disabuvir and vithout dos. And vithout disabuvir and vithout din	Mechanism						(see section 1.1) If
Sirolimus is due to CYP3A4 inhibition by ritonavir Vickirax Inhibitionavir Vickirax Inhibitionavir Vickirax Inhibitionavir Vickirax Inhibitionavir Vickirax Inhibitionavi				` '		` ′	1 ` '
Vickirax + dasabuvir Carolimus Caro	sirolimus is						with Viekirax + dasabuvir,
inhibition by ritonavir without dasabuvir without dasabuvir Similar effect is expected as observed with Vickirax + dasabuvir similar effect is expected as observed with Vickirax + dasabuvir Similar effect is expected as observed with Vickirax + dasabuvir without dasabuvir without dasabuvir without dasabuvir Similar effect is expected as observed with Vickirax + dasabuvir similar effect is expected as observed with Vickirax + dasabuvir similar effect is expected as observed with Vickirax + dasabuvir without dasabuvir yithout dasabuvir without dasabuvir without dasabuvir yithout dasabuvir yithout dasabuvir without dasabuvir yithout d		Viekirax	Not studied:				1
ritonavir dasabuvir			Similar effect	is expected as o	bserved with Vi	ekirax +	•
concentrations should be monitored every 4 to 7 days until 3 consecutive trough levels have shown stable concentrations of sirolimus. Sirolimus dose and/or dosing frequency should be adjusted as needed. 5 days after completion of Vickirax + dasabuvir treatment, the sirolimus dose and dosing frequency prior to receiving Vickirax should be resumed, along with routine monitoring of sirolimus dose and dosing frequency prior to receiving Vickirax should be resumed, along with routine monitoring of sirolimus blood concentrations. Tacrolimus 2 mg single dose? Wechanism: Effect on tacrolimus is dasabuvir and dasabuvir in thibition by ritonavir. Vickirax ↑ tacrolimus 0.89 5.7.1 (13.0-21.2) (13.0-		dasabuvir	dasabuvir				, ,
monitored every 4 to 7 days until 3 consecutive trough levels have shown stable concentrations of sirolimus. Sirolimus dose and/or dosing frequency should be adjusted as needed. 5 days after completion of Vickirax + dasabuvir treatment, the sirolimus dose and dosing frequency prior to receiving Vickirax + dasabuvir treatment, the sirolimus dose and dosing frequency prior to receiving Vickirax + dasabuvir treatment, the sirolimus dose and dosing frequency prior to receiving Vickirax + dasabuvir treatment, the sirolimus dose and dosing frequency prior to receiving Vickirax + dasabuvir treatment, the sirolimus dose and dosing frequency prior to receiving Vickirax + dasabuvir treatment, the sirolimus dose and dosing frequency prior to receiving Vickirax and dasabuvir is not entered to distribute the prior to receiving Vickirax and dasabuvir is not entered to dasabuvir are used to concentrations. Vickirax + dasabuvir treatment, the sirolimus dose and dosing frequency prior to receiving Vickirax and dasabuvir is not entered to distribute the prior to receiving Vickirax and dasabuvir are used to concentrations. Vickirax + dasabuvir treatment, the sirolimus dose and dosing frequency should be resumed, along with routine monitoring of sirolimus (0.61 to 2.1 to							Sirolimus blood
days until 3 consecutive trough levels have shown stable concentrations of sirolimus. Sirolimus dose and/or dosing frequency should be adjusted as needed. 5 days after completion of Viekirax + dasabuvir treatment, the sirolimus dose and dosing frequency prior to receiving Viekirax should be resumed, along with routine monitoring of sirolimus blood concentrations. Tacrolimus 2 mg single dose² 2 mg single dose² 4							
trough levels have shown stable concentrations of sirolimus. Sirolimus dose and/or dosing frequency should be adjusted as needed. 5 days after completion of Viekirax + dasabuvir treatment, the sirolimus dose and dosing frequency prior to receiving Viekirax should be resumed, along with routine monitoring of sirolimus blood concentrations. Tacrolimus Viekirax + dasabuvir 2 mg single dose ⁷ Mechanism: Effect on tacrolimus is due to CYP3A4 inhibition by ritonavir. Viekirax + dasabuvir 1 tacrolimus 3.99 57.1 16.6 0.00 (3.214.97) (45.5-71.7) (13.0-21.2) (13.0							I
tacrolimus Vickirax + dasabuvir Ababavir Ababavi							•
Sirolimus dose and/or dosing frequency should be adjusted as needed. 5 days after completion of Viekirax + dasabuvir treatment, the sirolimus dose and dosing frequency prior to receiving Viekirax should be resumed, along with routine monitoring of sirolimus blood concentrations. Tacrolimus 2 mg single dose² 2 mg single dose² 4 dasabuvir 4 dasabuvir 5 days after completion of Viekirax + dasabuvir treatment, the sirolimus dose and dosing frequency prior to receiving Viekirax should be resumed, along with routine monitoring of sirolimus blood concentrations. 4 0.93 0.94 0.94 0.94 0.94 0.94 0.9							•
$ \begin{array}{c ccccccccccccccccccccccccccccccccccc$							
should be adjusted as needed. 5 days after completion of Viekirax + dasabuvir treatment, the sirolimus dose and dosing frequency prior to receiving Viekirax should be resumed, along with routine monitoring of sirolimus blood concentrations. Tacrolimus 2 mg single dose ⁷ 2 mg single dose ⁷ Mechanism: Effect on tacrolimus is due to CYP3A4 inhibition by ritonavir. Viekirax + dasabuvir 1 tacrolimus 3.99 57.1 16.6 (3.21-4.97) (45.5-71.7) (13.0-21.2) (13							
$ \begin{array}{c ccccccccccccccccccccccccccccccccccc$							
$ \begin{array}{c ccccccccccccccccccccccccccccccccccc$							needed.
$ \begin{array}{c ccccccccccccccccccccccccccccccccccc$							
$ \begin{array}{c ccccccccccccccccccccccccccccccccccc$							5 days after completion of
treatment, the sirolimus dose and dosing frequency prior to receiving Vickirax should be resumed, along with routine monitoring of sirolimus blood concentrations. Tacrolimus 2 mg single dose ⁷ Mechanism: Effect on tacrolimus is due to CYP3A4 inhibition by ritonavir. Tacrolimus A tacrolim							
Tacrolimus Viekirax + dasabuvir $+$ $+$ $+$ $+$ $+$ $+$ $+$ $+$ $+$ $+$							
Tacrolimus Viekirax + dasabuvir 2 mg single dose 7 Mechanism: Effect on tacrolimus is due to CYP3A4 inhibition by ritonavir. Tacrolimus Viekirax + dasabuvir							-
Tacrolimus Viekirax + dasabuvir 2 mg single dose?							prior to receiving Viekirax
Tacrolimus Viekirax + dasabuvir Adsabuvir are initiated. Beginning the day after							, 0
Tacrolimus Viekirax + dasabuvir $\begin{array}{c ccccccccccccccccccccccccccccccccccc$							_
Tacrolimus Viekirax + dasabuvir $\begin{pmatrix} 1 & 1 & 1 & 1 & 1 & 1 & 1 & 1 & 1 & 1 $							
$\begin{array}{cccccccccccccccccccccccccccccccccccc$	Tagnali	Violeinen 1	↑ tacrolimus	3 00	57.1	16.6	
2 mg single $dose^7$	1 acronmus		tacronnus				
$\begin{array}{c ccccccccccccccccccccccccccccccccccc$	2 mg single	dasaouvii	\leftrightarrow	0.93		0.94	
Mechanism: Effect on tacrolimus is due to CYP3A4 inhibition by ritonavir. $ \begin{array}{c ccccccccccccccccccccccccccccccccccc$			ombitasvir				
Mechanism: Effect on tacrolimus is due to CYP3A4 inhibition by ritonavir.			paritanravir				benefits outweigh the risks
Effect on tacrolimus is due to CYP3A4 inhibition by ritonavir. Viekirax without dasabuvir Viekirax with Viekirax and dasabuvir are used concomitantly, tacrolimus should not be administered on the day Viekirax and dasabuvir are initiated. Beginning the day after	Mechanism:						
tacrolimus is due to CYP3A4 inhibition by ritonavir. Viekirax without dasabuvir Ombitasvir The magnitude of interaction was similar to that observed with Viekirax + dasabuvir. A.27 85.8 24.6 (19.7-30.8) concomitantly, tacrolimus should not be administered on the day Viekirax and dasabuvir are initiated. Paritaprevir Diekirax without (3.49-5.22) (67.9-108) (19.7-30.8) The magnitude of interaction was similar to that observed with Viekirax + dasabuvir. Diekirax without (3.49-5.22) (67.9-108) (19.7-30.8) The magnitude of interaction was similar to that observed with Viekirax + dasabuvir. Diekirax without dasabuvir are used concomitantly, tacrolimus should not be administered on the day Viekirax and dasabuvir are initiated. Diekirax without dasabuvir are used concomitantly, tacrolimus should not be administered on the day Viekirax and dasabuvir are initiated. Diekirax without dasabuvir are used concomitantly, tacrolimus should not be administered on the day Viekirax and dasabuvir are initiated. Diekirax without dasabuvir are used concomitantly, tacrolimus should not be administered on the day Viekirax and dasabuvir are initiated. Diekirax without dasabuvir are used concomitantly, tacrolimus should not be administered on the day Viekirax and dasabuvir are initiated. Diekirax without dasabuvir are used concomitantly, tacrolimus should not be administered on the day Viekirax and dasabuvir are initiated.			dusuouvii				
CYP3A4 dasabuvir dasabuvir		Viekirax	↑ tacrolimus	4.27	85.8	24.6	
inhibition by ritonavir. ombitasvir paritaprevir ombitasvir ombitasvir ombitasvir paritaprevir ombitasvir ombitasvir to that observed with Viekirax + dasabuvir. on the day Viekirax and dasabuvir are initiated. Beginning the day after							•
ritonavir. dasabuvir are initiated. paritaprevir Beginning the day after		dasabuvir					
paritaprevir Beginning the day after	•		.l.	to mai ooselve	od with vickilax	uasavuvii.	=
	manavir.		paritaprevir				

Medicinal Product/Poss ible	GIVEN WITH	EFFECT	Cmax	AUC	Ctrough	Clinical Comments
Mechanism of Interaction						
						initiated; reinitiate tacrolimus at a reduced dose based on tacrolimus blood concentrations. The recommended tacrolimus dosing is 0.5 mg every 7 days. Tacrolimus whole blood concentrations should be monitored upon initiation and throughout co- administration with Viekirax and dasabuvir and the dose and/or dosing frequency should be adjusted as needed. Upon completion of Viekirax and dasabuvir treatment, the appropriate dose and dosing frequency of tacrolimus should be guided by assessment of tacrolimus blood
INHALED BE	L TA AGONIS'	TS				concentrations.
Salmeterol	Viekirax	Not studied. Ex	spected:			Concomitant use is
Mechanism: CYP3A4 inhibition by ritonavir.	with or without dasabuvir	↑ salmeterol	•			contraindicated (see section 4.3).
INSULIN SEC	RETAGOGU	UES				
Repaglinide Mechanism: OATP1B1 inhibition by paritaprevir.	Viekirax with or without dasabuvir	Not Studied. E. ↑ repaglinide	xpected:			Caution should be used and dose decrease maybe needed for repaglinide when administered with Viekirax with or without
Name of the second						dasabuvir.
Carisoprodol 250 mg single dose	Viekirax with dasabuvir	↓ Carisoprodol ↔ ombitasvir	0.54 (0.47-0.63) 0.98	0.62 (0.55-0.70) 0.95	NA 0.96	No dose adjustment required for carisoprodol; increase dose if clinically indicated.
		↔ paritaprevir	(0.92-1.04) 0.88 (0.75-1.03)	(0.92-0.97) 0.96 (0.85-1.08)	(0.92-0.99) 1.14 (1.02-1.27)	mulculcu.

Medicinal	GIVEN	EFFECT	Cmax	AUC	Ctrough	Clinical Comments
Product/Poss	WITH					
ible						
Mechanism						
of Interaction						
Mechanism:		↔ dasabuvir	0.96	1.02	1.00	
CYP2C19		v dasaouvii	(0.91-1.01)	(0.97-1.07)	(0.92-1.10)	
induction by				,		
ritonavir						
	Viekirax		Not st		•	
	without	Similar effe		observed with	Viekirax +	
	dasabuvir		dasab	ouvir.		
Cyclobenzapr	Viekirax	↓ cycloben-	0.68	0.60	NA	No dose adjustment
ine 5 mg	with	zaprine	(0.61-0.75)	(0.53-0.68)		required for cyclobenzaprine; increase
single dose	dasabuvir	↔ ombitasvir	0.98	1.00	1.01	dose if clinically indicated.
			(0.92-1.04)	(0.97-1.03)	(0.98-1.04)	accommodity maneure
Mechanism:		→ paritaprevir	1.14	1.13	1.13	
decrease			(0.99-1.32)	(1.00-1.28)	(1.01-1.25)	
possibly due to CYP1A2		↔ dasabuvir	0.98	1.01	1.13	
induction by			(0.90-1.07)	(0.96-1.06)	(1.07-1.18)	
ritonavir						
11001100 / 11	Viekirax		Not stu	ıdied.	<u> </u>	
	without	Similar effe	ct expected as	observed with V	iekirax +	
	dasabuvir		dasab	uvir.		
NARCOTIC A	NALGESIC	CS			_	
Paracetamol	Viekirax	\leftrightarrow	1.02	1.17	NA	No dose adjustment
(as given in a	+	paracetamol	(0.89-1.18)	(1.09-1.26)		necessary for paracetamol
fixed-dose	dasabuvir	↔ ombitasvir	1.01	0.97	0.93	when administered with
hydrocodone/			(0.93-1.10)	(0.93-1.02)	(0.90-0.97)	Viekirax with or without
paracetamol)		→ paritaprevir	1.01 (0.80-1.27)	1.03 (0.89-1.18)	1.10 (0.97-1.26)	dasabuvir.
300 mg single		↔ dasabuvir	1.13	1.12	1.16	
dose			(1.01-1.26)	(1.05-1.19)	(1.08-1.25)	
	Viekirax		Not stu	` ′	,	
	without	Similar effe	ct expected as	observed with V	iekirax +	
	dasabuvir		dasab	uvir.		
Hydrocodone	Viekirax	↑ hydrocodo	1.27	1.90	NA	A reduction of
(as given in a	+	ne	(1.14-1.40)	(1.72-2.10)		hydrocodone dose by 50%
fixed-dose	dasabuvir	_		itaprevir and da		and/or clinical monitoring
hydrocodone/	Viekirax	same	as snown for p Not stu	paracetamol abo	VC	should be considered when
paracetamol)	without	Similar effe		observed with V	iekirax +	administered with Viekirax
5 m c viv 1	dasabuvir		dasab			with or without dasabuvir.
5 mg single						
dose						
Mechanism:						
L CYP3A4						
CYP3A4 inhibition by						

Medicinal Product/Poss	GIVEN WITH	EFFECT	C _{max}	AUC	Ctrough	Clinical Comments
ible Machanism						
Mechanism of						
Interaction						
OPIOIDS					ı	
Methadone	Viekirax +	↔ R-	1.04	1.05	0.94	No dose adjustment is
Wildingone	dasabuvir	Methadone	(0.98-1.11)	(0.98-1.11)	(0.87-1.01)	necessary for methadone
20-120 mg		↔ S-	0.99	0.99	0.86	and Viekirax with or
once daily ⁸		Methadone	(0.91-1.08)	(0.89-1.09)	(0.76-0.96)	without dasabuvir.
			ir /ombitasvir/da study con	asabuvir (based asabuvir)	on the cross-	
	Viekirax		-			
	without dasabuvir	The magnitud with Viekirax	e of interaction + dasabuvir.			
Buprenorphine	Viekirax +	↑ bu-	2.18	2.07	3.12	No dose adjustment is
/ naloxone	dasabuvir	prenorphine	(1.78-2.68)	(1.78-2.40)	(2.29-4.27)	necessary for buprenorphine/naloxone
4-24 mg/1-		↑ norbu-	2.07	1.84	2.10	and Viekirax with or
6 mg once		prenorphine	(1.42-3.01)	(1.30-2.60)	(1.49-2.97)	without dasabuvir.
daily ⁸		↑ naloxone	1.18	1.28	NA	1
		1	(0.81-1.73)	(0.92-1.79)		
Mechanism: CYP3A4		↔ ombitasvi	r/paritaprevir/da study con			
inhibition by	Viekirax	↑ bu-	1.19	1.51	1.65	
ritonavir and	without	prenorphine	(1.01-1.40)	(1.27-1.78)	(1.30-2.08)	
UGT	dasabuvir	↑ norbu-		de of interaction ed with Viekirax		
inhibition by		prenorphine	to that observe			
paritaprevir,			svir/paritaprevir	1		
ombitasvir			compa			
and			•	,		
dasabuvir.						
PHOSPHODI		· · · · · · · · · · · · · · · · · · ·				
Sildenafil	Viekirax	Not studied. E	xpected:			Concomitant use is
(when used	with and	↑ sildenafil		contraindicated (see section 4.3).		
for treatment	without	Sildellaili		4.3).		
of pulmonary	dasabuvir					
hypertension)						
Mechanism:						
CYP3A4						
inhibition by						
ritonavir. PROTON PUN	 MD INLIDITA	ODS				
INOTONPUN	Viekirax +		0.62	0.62	NA	If clinically indicated
	dasabuvir	omeprazole	(0.48-0.80)	(0.51-0.75)	1 1/1	higher doses of
	GasaouvII		1.02	1.05	1.04	Inglier doses of
		ombitasvir	(0.95-1.09)	(0.98-1.12)	(0.98-1.11)	
		. ↔ .	1.19	1.18	0.92	
		paritaprevir	(1.04-1.36)	(1.03-1.37)	(0.76-1.12)	-
		↔ dasabuvir	1.13 (1.03-1.25)	1.08 (0.98-1.20)	1.05 (0.93-1.19)	
	I .]	(1.03-1.23)	(0.70-1.20)	(U.73-1.19)	L

Medicinal	GIVEN	EFFECT	Cmax	AUC	Ctrough	Clinical Comments
Product/Poss	WITH					
ible Machanian						
Mechanism of						
Interaction						
Omeprazole	Viekirax	J.	0.48	0.46	NA	omeprazole should be
отпертидете	without	omeprazole	(0.29 - 0.78)	(0.27-0.77)		used.
40 mg once	dasabuvir	\leftrightarrow		de of interaction		
daily		ombitasvir	to that observe	No dose adjustment		
		↔ paritaprevir				needed for Viekirax with
		puritupievii				or without dasabuvir.
Mechanism:						
CYP2C19						
induction by						
ritonavir.	77' 1'	Not studied D				TC 1' ' 11 ' 1' 4 1
Esomeprazole Lansoprazole	Viekirax with and	Not studied. E	xpectea: le, lansoprazole			If clinically indicated, higher doses of
Mechanism:	without		.e, 14115ep142e16	esomeprazole/lansoprazole		
CYP2C19	dasabuvir					may be needed.
induction by						
ritonavir.						
SEDATIVES /	HYPNOTIC	S				1
Zolpidem	Viekirax +	↔ zolpidem	0.94	0.95	NA	No dose adjustment is
	dasabuvir		(0.76-1.16)	(0.74-1.23)	1.04	necessary for zolpidem.
5 mg single		↔ ombitasvir	1.07 (1.00-1.15)	1.03 (1.00-1.07)	1.04 (1.00-1.08)	
dose		Omortasvii	0.63	0.68	1.23	No dose adjustment
		paritaprevir	(0.46-0.86)	(0.55-0.85)	(1.10-1.38)	needed for Viekirax with
		↔ dasabuvir	0.93	0.95	0.92	or without dasabuvir.
	77' 1'		(0.84-1.03) Not st	(0.84-1.08)	(0.83-1.01)	
	Viekirax without	Similar eff	not st ect expected as			
	dasabuvir	211111W	dasab			
Alprazolam	Viekirax +	<u> </u>	1.09	1.34	NA	Clinical monitoring of
11p1#201#111	dasabuvir	alprazolam	(1.03-1.15)	(1.15-1.55)		patients is recommended.
0.5 mg single		↔	0.98	1.00	0.98	A decrease in alprazolam
dose		ombitasvir ↔	(0.93-1.04) 0.91	(0.96-1.04) 0.96	(0.93-1.04)	dose can be considered
		paritaprevir	(0.64-1.31)	(0.73-1.27)	(1.02-1.23)	based on clinical response.
		↔ dasabuvir	0.93	0.98	1.00	
Mechanism:			(0.83-1.04)	(0.87-1.11)	(0.87-1.15)	No dose adjustment
CYP3A4	Viekirax	Cimil	Not st	needed for Viekirax with or without dasabuvir.		
inhibition by ritonavir	without	Similar effect expected as observed with Viekirax + dasabuvir.				or without dasabuvir.
	dasabuvir	Not -4-3'-1 P		Componit		
Oral midazolam	Viekirax with or	Not studied. Expected:				Concomitant use is contraindicated (see
midazolam Triazolam	with or without	↑ midazolam or triazolam				section 4.3).
11142014111	dasabuvir					
Mechanism:						If parenteral midazolam is
CYP3A4						co-administered with
inhibition by						Viekirax with or without dasabuvir, close clinical
ritonavir.						uasaouvii, ciose ciiiiical

Medicinal Product/Poss ible Mechanism of Interaction	GIVEN WITH	EFFECT	Cmax	AUC	Ctrough	Clinical Comments
						monitoring for respiratory depression and/or prolonged sedation should be exercised and dosage adjustment should be considered.
Diazepam	Viekirax +	↓diazepam	1.18	0.78	NA	No dose adjustment
	dasabuvir		(1.07-1.30)	(0.73-0.82)		required for diazepam;
2 mg single		↓ ↓	1.10	0.56	NA	increase dose if clinically
dose		nordiazepam	(1.03-1.19)	(0.45-0.70)		indicated.
		↔ ombitasvir	1.00	0.98	0.93	
Mechanism:			(0.93-1.08)	(0.93-1.03)	(0.88-0.98)	
CYP2C19		↔ paritaprevir	0.95	0.91	0.92	
induction by			(0.77-1.18)	(0.78-1.07)	(0.82-1.03)	
ritonavir		↔ dasabuvir	1.05	1.01	1.05	
			(0.98-1.13) (0.94-1.08) (0.98-1.12)			-
	Viekirax	Similar offe	Not studied.			
	without	Similar effect expected as observed with Viekirax + dasabuvir.				
	dasabuvir		dusae			
THYROID HO						1
Levothyroxine	Viekirax	Not studied. Ex	pected:	Clinical monitoring and		
	with or	↑ levothyroxine	.			dose adjustment may be
Mechanism:	without	ICVOIIIyIOXIIIC		required for levothyroxine		
UGT1A1	dasabuvir					
inhibition by						
paritaprevir,						
ombitasvir						
and						

- 1. Lopinavir/ritonavir 800/200 mg once daily (administered in the evening) was also administered with Viekirax with or without dasabuvir. The effect on C_{max} and AUC of DAAs and lopinavir was similar to that observed when lopinavir/ritonavir 400/100 mg twice daily was administered with Viekirax with or without dasabuvir.
- 2. Rilpivirine was also administered in the evening with food and at night 4 hours after dinner with Viekirax + dasabuvir in other two arms in the study. The effect on rilpivirine exposures was similar to that observed when rilpivirine was administered in the morning with food with Viekirax + dasabuvir (shown in the table above).
- 3. Ciclosporin 100 mg dosed alone, 10 mg administered with Viekirax and 30 mg administered with Viekirax + dasabuvir. Dose normalized cyclosporine ratios are shown for interaction with Viekirax with or without dasabuvir.
- 4. C_{12} := concentration at 12 hours following single dose of everolimus.

dasabuvir.

Medicinal	GIVEN	EFFECT	Cmax	AUC	Ctrough	Clinical Comments
Product/Poss	WITH					
ible						
Mechanism						
of						
Interaction						

- 5. Sirolimus 2 mg was dosed alone, 0.5 mg administered with Viekirax + dasabuvir. Dose normalized sirolimus ratios are shown for interaction with Viekirax + dasabuvir.
- 6. C₂₄:= concentration at 24 hours following single dose of cyclosporine, tacrolimus or sirolimus.
- 7. Tacrolimus 2 mg was dosed alone, 0.5 mg administered with Viekirax and 2 mg was administered with Viekirax + dasabuvir. Dose normalized tacrolimus ratios are shown for interaction with Viekirax with or without dasabuvir.
- 8. Dose normalised parameters reported for methadone, buprenorphine and naloxone.

Note: Doses used for Viekirax and dasabuvir were: ombitasvir 25 mg, paritaprevir 150 mg, ritonavir 100 mg, once daily and dasabuvir 400 mg twice daily or 250 mg twice daily. The dasabuvir exposures obtained with the 400 mg formulation and the 250 mg tablet are similar. Viekirax with or without dasabuvir was administered as multiple doses in all the drug interaction studies except the drug interaction studies with carbamazepine, gemfibrozil, ketoconazole, and sulfamethoxazole/trimethoprim..

Paediatric population

Drug interaction studies have only been performed in adults.

4.6 Fertility, pregnancy and lactation

Women of childbearing potential / contraception in males and females

Extreme caution must be taken to avoid pregnancy in female patients and female partners of male patients when Viekirax is taken in combination with ribavirin. Significant teratogenic and/or embryocidal effects have been demonstrated in all animal species exposed to ribavirin; therefore, ribavirin is contraindicated in women who are pregnant and in the male partners of women who are pregnant. Refer to the Summary of Product Characteristics for ribavirin for additional information.

Female patients: Women of childbearing potential should not receive ribavirin unless they are using an effective form of contraception during treatment with ribavirin and for 4 months after treatment. Ethinyloestradiol is contraindicated in combination with Viekirax (see sections 4.3 and 4.4).

Male patients and their female partners: Either male patients or their female partners of childbearing potential must use a form of effective contraception during treatment with ribavirin and for 7 months after treatment.

Pregnancy

There are very limited data from the use of Viekirax in pregnant women. Studies with ombitasvir and paritaprevir/ritonavir in animals have shown malformations (see section 5.3). The potential risk for humans is unknown. Viekirax should not be used during pregnancy or in women of childbearing potential not using effective contraception.

If ribavirin is co-administered with Viekirax, the contraindications regarding use of ribavirin during pregnancy apply (see also the Summary of Product Characteristics of ribavirin).

Breast-feeding

It is not known whether paritaprevir /ritonavir or ombitasvir and their metabolites are excreted in human breast milk. Available pharmacokinetic data in animals have shown excretion of active substance and metabolite in milk (see section 5.3). Because of the potential for adverse reactions from the medicinal product in breastfed infants, a decision must be made whether to discontinue breast-feeding or discontinue treatment with Viekirax, taking into account the importance of the therapy to the mother. For patients co-administered ribavirin refer to the Summary of Product Characteristics of ribavirin.

Fertility

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

4.7 Effects on ability to drive and use machines

Viekirax has no or negligible influence on the ability to drive and use machines. Patients should be informed that fatigue has been reported during treatment with Viekirax in combination with dasabuvir and ribavirin (see section 4.8).

4.8 Undesirable effects

Summary of the safety profile

In subjects receiving Viekirax and dasabuvir with ribavirin, the most commonly reported adverse reactions (greater than 20% of subjects) were fatigue and nausea. The proportion of subjects who permanently discontinued treatment due to adverse reactions was 0.2% (5/2,044) and 4.8% (99/2,044) of subjects had ribavirin dose reductions due to adverse reactions.

Tabulated list of adverse reactions

The safety summary is based on pooled data from phase 2 and 3 clinical trials in subjects who received Viekirax and dasabuvir with or without ribavirin. The majority of adverse reactions presented in Table 3 were of grade 1 severity in Viekirax and dasabuvir-containing regimens.

The adverse reactions are listed below by system organ class and frequency. Frequencies are defined as follows: very common ($\geq 1/10$), common ($\geq 1/100$) to <1/10), uncommon ($\geq 1/1,000$) to <1/1,000) or very rare (<1/10,000).

Table 3. Adverse drug reactions identified with Viekirax in combination with dasabuvir with and without ribavirin

Frequency	Viekirax + dasabuvir + ribavirin* N = 2,044	Viekirax + dasabuvir N = 588
Blood and lymphatic syst	em disorders	
Common	Anaemia	
Immune system disorders		
Frequency unknown	Anaphylactic reactions	Anaphylactic reactions
Metabolism and nutrition	disorders	
Uncommon	Dehydration	
Psychiatric disorders		
Very common	Insomnia	
Gastrointestinal disorder	S	
Very common	Nausea, Diarrhoea	
Common	Vomiting	
Hepatobiliary disorders		
Frequency unknown	Hepatic decompensation and hepatic failure	Hepatic decompensation and hepatic failure
Skin and subcutaneous tis	ssue disorders	
Very common	Pruritus	
Common		Pruritus
Rare	Angioedema	Angioedema
		<u> </u>
General disorders and ad	lministration and administration s	site conditions
	Asthenia	
Very common		
	Fatigue	

^{*}Data set includes all genotype 1-infected subjects in Phase 2 and 3 trials including subjects with cirrhosis.

Note: For laboratory abnormalities, refer to Table 4

Description of selected adverse reactions

Compared to subjects without cirrhosis, in subjects with compensated cirrhosis there was an increased rate of indirect hyperbilirubinemia when ribavirin was part of the regimen.

Laboratory abnormalities

Changes in selected laboratory parameters are described in Table 4. A side-by-side tabulation is shown to simplify presentation; direct comparison across trials should not be made due to differing trial designs.

Table 4. Selected treatment emergent laboratory abnormalities

	SAPPHIRE I and II	PEARL II, III, and IV	TURQUOISE II (subjects with cirrhosis)	
Laboratory Parameters	Viekirax and dasabuvir + ribavirin	Viekirax and dasabuvir 12 weeks	Viekirax and dasabuvir + ribavirin	
	12 weeks	N=509	12 or 24 weeks	
	N = 770	n (%)	N=380	
	n (%)		n (%)	
ALT				
>5-20 × ULN* (Grade 3)	6/765 (0.8%)	1/509 (0.2%)	4/380 (1.1%)	
>20 × ULN (Grade 4)	3/765 (0.4%)	0	2/380 (0.5%)	
Haemoglobin				
<100-80 g/L (grade 2)	41/765 (5.4%)	0	30/380 (7.9%)	
<80-65 g/L (grade 3)	1/765 (0.1%)	0	3/380 (0.8%)	
<65 g/L (Grade 4)	0	0	1/380 (0.3%)	
Total bilirubin				
>3-10 × ULN (grade 3)	19/765 (2.5%)	2/509 (0.4%)	37/380 (9.7%)	
>10 × ULN (grade 4)	1/765 (0.1%)	0	0	
	1/765 (0.1%) al according to testing labora		0	

Serum ALT elevations

In a pooled analysis of clinical trials with Viekirax and dasabuvir with and without ribavirin, 1% of subjects experienced serum ALT levels greater than 5 times the upper limit of normal (ULN) after starting treatment. As the incidence of such elevations was 26% among women taking a concomitant ethinyloestradiol-containing medicinal product, such medicinal products are contraindicated with Viekirax with or without dasabuvir. No increase in incidence of ALT elevations was observed with other types of estrogens commonly used for hormone replacement therapy (e.g. oestradiol and conjugated estrogens). ALT elevations were typically asymptomatic, generally occurred during the first 4 weeks of treatment (mean time 20 days, range 8-57 days) and most resolved with ongoing therapy. Two patients discontinued Viekirax and dasabuvir due to elevated ALT, including one on ethinyloestradiol. Three interrupted Viekirax and dasabuvir for one to seven days, including one on ethinyloestradiol. The majority of these ALT elevations were transient and assessed as drug-related. Elevations in ALT were generally not associated with bilirubin elevations. Cirrhosis was not a risk factor for elevated ALT (see section 4.4).

Serum bilirubin elevations

Transient elevations in serum bilirubin (predominantly indirect) were observed in subjects receiving Viekirax and dasabuvir with ribavirin, related to the inhibition of the bilirubin transporters OATP1B1/1B3 by paritaprevir and ribavirin-induced haemolysis. Bilirubin elevations occurred after initiation of treatment, peaked by study Week 1, and generally resolved with ongoing therapy. Bilirubin elevations were not associated with aminotransferase elevations. The frequency of indirect bilirubin elevations was lower among subjects who did not receive ribavirin.

Liver transplant recipients

The overall safety profile in HCV-infected transplant recipients who were administered Viekirax and dasabuvir and ribavirin (in addition to their immunosuppressant medications) was similar to subjects

treated with Viekirax and dasabuvir and ribavirin in phase 3 clinical trials, although some adverse reactions were increased in frequency. 10 subjects (29.4%) had at least one post baseline haemoglobin value of less than 10 g/dL. 10 of 34 subjects (29.4%) dose modified ribavirin due to decrease in haemoglobin and 2.9% (1/34) had an interruption of ribavirin. Ribavirin dose modification did not impact SVR rates. 5 subjects required erythropoietin, all of whom initiated ribavirin at the starting dose of 1000 to 1200 mg daily. No subject received a blood transfusion.

HIV/HCV co-infected patients

The overall safety profile in HCV/HIV-1 co-infected subjects was similar to that observed in HCV monoinfected subjects. Transient elevations in total bilirubin >3 x ULN (mostly indirect) occurred in 17 (27.0%) subjects; 15 of these subjects were receiving atazanavir. None of the subjects with hyperbilirubinemia had concomitant elevations of aminotransferases.

GTI-infected subjects with or without cirrhosis with severe renal impairment or end-stage renal disease (ESRD)

Viekirax and dasabuvir with or without ribavirin were assessed in 68 subjects with genotype 1 infection with or without cirrhosis who have severe renal impairment or ESRD (see Section 5.1). The overall safety profile in subjects with severe renal impairment was similar to that seen in prior Phase 3 studies in subjects without severe renal impairment, except that a greater proportion of subjects required intervention due to ribavirin-associated decreases in serum haemoglobin. The mean baseline haemoglobin level was 12.1 g/dL and the mean decline in haemoglobin at the end of treatment for subjects taking RBV was 1.2 g/dL. Thirty-nine of the 50 subjects who received ribavirin required interruption of ribavirin, and 11 of these subjects were also treated with erythropoietin. Four subjects experienced a haemoglobin level < 8 g/dL. Two subjects received a blood transfusion. Adverse events of anaemia were not seen in the 18 GT1b-infected subjects who did not receive ribavirin. Viekirax with or without dasabuvir was also evaluated without ribavirin in 18 GT1a- and GT4-infected patients; no adverse events of anaemia were seen in these subjects.

Paediatric population

The safety of Viekirax in children and adolescents aged < 18 years has not yet been established. No data are available.

Reporting of suspected adverse reactions

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

4.9 Overdose

The highest documented single dose administered to healthy volunteers was 400 mg for paritaprevir (with 100 mg ritonavir), 200 mg for ritonavir (with 100 mg paritaprevir) and 350 mg for ombitasvir. No study related adverse reactions with paritaprevir, ritonavir, or ombitasvir were observed. Transient increases in indirect bilirubin were observed at the highest doses of paritaprevir/ritonavir. In case of overdose, it is recommended that the patient be monitored for any signs or symptoms of adverse reactions or effects and appropriate symptomatic treatment instituted immediately.

5. PHARMACOLOGICAL PROPERTIES

5.1 Pharmacodynamic properties

Pharmacotherapeutic group: Antivirals for systemic use; direct-acting antivirals, ATC code: J05AP53

Mechanism of action

Viekirax, when co-administered with dasabuvir, combines three direct-acting antiviral medicinal products with distinct mechanisms of action and non-overlapping resistance profiles to target HCV at multiple steps in the viral lifecycle. Refer to the Summary of Product Characteristics of dasabuvir for its pharmacological properties.

Ritonavir

Ritonavir is not active against HCV. Ritonavir is a CYP3A inhibitor that increases the systemic exposure of the CYP3A substrate paritaprevir.

Ombitasvir

Ombitasvir is an inhibitor of HCV NS5A which is essential for viral replication.

Paritaprevir

Paritaprevir is an inhibitor of HCV NS3/4A protease which is necessary for the proteolytic cleavage of the HCV encoded polyprotein (into mature forms of the NS3, NS4A, NS4B, NS5A, and NS5B proteins) and is essential for viral replication.

Activity in cell culture and/or biochemical studies

Ombitasvir

The EC₅₀ of ombitasvir against genotype 1a-H77 and 1b-Con1 strains in HCV replicon cell culture assays was 14.1 and 5 pM, respectively. The activity of ombitasvir was attenuated 11- to 13-fold in the presence of 40% human plasma. The mean EC₅₀ of ombitasvir against replicons containing NS5A from a panel of treatment-naïve genotype 1a and 1b isolates in the HCV replicon cell culture assay was 0.66 pM (range 0.35 to 0.88 pM; n=11) and 1.0 pM (range 0.74 to 1.5 pM; n=11), respectively. Ombitasvir has EC₅₀ values of 12, 4.3, 19, 1.7, 3.2, and 366 pM against replicon cell lines constructed with NS5A from single isolates representing genotypes 2a, 2b, 3a, 4a, 5a, and 6a, respectively.

Paritaprevir

The EC $_{50}$ of paritaprevir against genotype 1a-H77 and 1b-Con1 strains in the HCV replicon cell culture assay was 1.0 and 0.21 nM, respectively. The activity of paritaprevir was attenuated 24 to 27 -fold in the presence of 40% human plasma. The mean EC $_{50}$ of paritaprevir against replicons containing NS3 from a panel of treatment-naïve genotype 1a and 1b isolates in the HCV replicon cell culture assay was 0.86 nM (range 0.43 to 1.87 nM; n=11) and 0.06 nM (range 0.03 to 0.09 nM; n=9), respectively. Paritaprevir had an EC $_{50}$ value of 5.3 nM against the 2a-JFH-1 replicon cell line, and EC $_{50}$ values of 19, 0.09, and 0.68 nM against replicon cell lines containing NS3 from a single isolate each of genotype 3a, 4a, and 6a, respectively

Ritonavir did not exhibit a direct antiviral effect on the replication of HCV subgenomic replicons, and the presence of ritonavir did not affect the *in vitro* antiviral activity of paritaprevir.

Resistance

In cell culture

Genotype 1

Resistance to paritaprevir and ombitasvir conferred by variants in NS3 and NS5A respectively, selected in cell culture or identified in Phase 2b and 3 clinical trials were phenotypically characterised in the appropriate genotype 1a or 1b replicons.

In genotype 1a, substitutions F43L, R155K, A156T, and D168A/F/H/V/Y in HCV NS3 reduced susceptibility to paritaprevir. In the genotype 1a replicon, the activity of paritaprevir was reduced 20-, 37-, and 17-fold by the F43L, R155K and A156T substitutions, respectively. The activity of paritaprevir was reduced 96-fold by D168V, and 50- to 219-fold by each of the other D168 substitutions. The activity of paritaprevir in genotype 1a was not significantly affected (less than or equal to 3-fold) by single substitutions V36A/M, V55I, Y56H, Q80K or E357K. Double variants including combinations of V36LM, F43L, Y56H, Q80K or E357K with R155K or with a D168 substitution reduced the activity of paritaprevir by an additional 2 to 3-fold relative to the single R155K or D168 substitution. In the genotype 1b replicon, the activity of paritaprevir was reduced 76- and 159-and 337- fold by D168A, D168H, D168V, and D168Y respectively. Y56H alone could not be evaluated due to poor replication capacity, however, the combination of Y56H and D168A/V/Y reduced the activity of paritaprevir by 700- to 4118-fold.

In genotype 1a, substitutions M28T/V, Q30E/R, L31V, H58D, Y93C/H/N, and M28V + Q30R in HCV NS5A reduced susceptibility to ombitasvir. In the genotype 1a replicon, the activity of ombitasvir was reduced by 896-, 58- and 243-fold against the M28T/V and H58D substitutions, respectively, and 1326-, 800-, 155-foldand 1675- to 66740- fold by the Q30E/R, L31V and Y93C/H/N substitutions, respectively. Y93H, Y93N or M28V in combination with Q30R reduced the activity of ombitasvir by more than 42,802-fold. In genotype 1b, substitutions L28T, L31F/V, as well as Y93H alone or in combination with L28M, R30Q, L31F/M/V or P58S in HCV NS5A reduced susceptibility to ombitasvir. In the genotype 1b replicon, the activity of ombitasvir was reduced by less than 10-fold by variants at amino acid positions 30 and 31. The activity of ombitasvir was reduced by 661-, 77-, 284- and 142-fold against the genotype 1b substitutions L28T, Y93H, R30Q in combination with Y93H, and L31M in combination with Y93H, respectively. All other double substitutions of Y93H in combination with substitutions at positions 28, 31, or 58 reduced the activity of ombitasvir by more than 400-fold.

Genotype 4

In genotype 4a, resistance to paritaprevir or ombitasvir by variants in NS3 or NS5A, respectively, selected in cell culture were phenotypically characterised. Substitutions R155C, A156T/V, and D168H/V in HCV NS3 reduced susceptibility to paritaprevir by 40- to 323-fold. Substitution L28V in HCV NS5A reduced the susceptibility to ombitasvir by 21-fold.

Effect of baseline HCV substitutions/polymorphisms on treatment outcome
A pooled analysis of subjects with genotype 1 HCV infection, who were treated with ombitasvir, paritaprevir, and dasabuvir (a non-nucleotide NS5B inhibitor) with or without ribavirin in the Phase 2b and 3 clinical trials was conducted to explore the association between baseline NS3/4A, NS5A or NS5B substitutions/polymorphisms and treatment outcome in recommended regimens.

In the greater than 500 genotype 1a baseline samples in this analysis, the most frequently observed resistance-associated variants were M28V (7.4%) in NS5A and S556G (2.9%) in NS5B. Q80K, although a highly prevalent polymorphism in NS3 (41.2% of samples), confers minimal resistance to paritaprevir.

Resistance-associated variants at amino acid positions R155 and D168 in NS3 were rarely observed (less than 1%) at baseline. In the greater than 200 genotype 1b baseline samples in this analysis, the most frequently observed resistance-associated variants observed were Y93H (7.5%) in NS5A, and C316N (17.0%) and S556G (15%) in NS5B. Given the low virologic failure rates observed with recommended treatment regimens for HCV genotype 1a- and 1b-infected subjects, the presence of baseline variants appears to have little impact on the likelihood of achieving SVR.

In clinical studies

Of the 2,510 HCV genotype 1 infected subjects who were treated with regimens containing ombitasvir, paritaprevir, and dasabuvir with or without ribavirin (for 8, 12, or 24 weeks) in Phase 2b and 3 clinical trials, a total of 74 subjects (3%) experienced virologic failure (primarily post-treatment relapse). Treatment-emergent variants and their prevalence in these virologic failure populations are shown in Table 5. In the 67 genotype 1a infected subjects, NS3 variants were observed in 50 subjects, NS5A variants were observed in 46 subjects, NS5B variants were observed in 37 subjects, and treatment-emergent variants were seen in all 3 drug targets in 30 subjects. In the 7 genotype 1b infected subjects, treatment-emergent variants were observed in NS3 in 4 subjects, in NS5A in 2 subjects, and in both NS3 and NS5A in 1 subject. No genotype 1b infected subjects had treatment-emergent variants in all 3 drug targets.

Table 5. Treatment-emergent amino acid substitutions in the pooled analysis of Viekirax and dasabuvir with and without RBV regimens in Phase 2b and Phase 3 clinical trials (N=2510)

		Genotype 1a N=67 ^b	Genotype 1b N=7
Target	Emergent amino acid substitutions ^a	% (n)	% (n)
NS3	V55I°	6 (4)	
	Y56H ^c	9 (6)	42.9 (3) ^d
	I132V ^c	6 (4)	
	R155K	13.4 (9)	
	D168A	6 (4)	
	D168V	50.7 (34)	42.9 (3) ^d
	D168Y	7.5 (5)	
	V36A°, V36M°, F43L°, D168H, E357K°	< 5%	
NS5A	M28T	20.9 (14)	
	M28V ^e	9 (6)	
	Q30R ^e	40.3 (27)	
	Ү93Н		28.6 (2)
	H58D, H58P, Y93N	< 5%	
NS5B	A553T	6.1 (4)	
	S556G	33.3 (22)	
	C316Y, M414T, G554S, S556R, G558R, D559G, D559N, Y561H	< 5%	

- a. Observed in at least 2 subjects of the same subtype.
- b. N=66 for the NS5B target.
- c. Substitutions were observed in combination with other emergent substitutions at NS3 position R155 or D168.
- d. Observed in combination in genotype 1b-infected subjects.
- e. Observed in combination in 6% (4/67) of the subjects.

Note: The following variants were selected in cell culture but were not treatment-emergent: NS3 variants A156T in genotype 1a, and R155Q and D168H in genotype 1b; NS5A variants Y93C/H in

genotype 1a, and L31F/V or Y93H in combination with L28M, L31F/V or P58S in genotype 1b; and NS5B variants Y448H in genotype 1a, and M414T and Y448H in genotype 1b.

Persistence of resistance-associated substitutions

The persistence of paritaprevir, ombitasvir, and dasabuvir resistance-associated amino acid substitutions in NS3, NS5A, and NS5B, respectively, was assessed in genotype 1a-infected subjects in Phase 2b trials. Paritaprevir treatment-emergent variants V36A/M, R155K or D168V were observed in NS3 in 47 subjects. Ombitasvir treatment-emergent variants M28T, M28V or Q30R in NS5A were observed in 32 subjects. Dasabuvir treatment-emergent variants M414T, G554S, S556G, G558R or D559G/N in NS5B were observed in 34 subjects.

NS3 variants V36A/M and R155K and NS5B variants M414T and S556G remained detectable at post-treatment Week 48, whereas NS3 variant D168V and all other NS5B variants were not observed at post-treatment Week 48. All treatment-emergent variants in NS5A remained detectable at post-treatment Week 48. Due to high SVR rates in genotype 1b, trends in persistence of treatment-emergent variants in this genotype could not be established.

The lack of detection of virus containing a resistance-associated substitution does not indicate that the resistant virus is no longer present at clinically significant levels. The long-term clinical impact of the emergence or persistence of virus containing Viekirax- and dasabuvir-resistance-associated substitutions on future treatment is unknown.

Cross-resistance

Cross-resistance is expected among NS5A inhibitors, NS3/4A protease inhibitors, and non-nucleoside NS5B inhibitors by class. The impact of prior ombitasvir, paritaprevir or dasabuvir treatment experience on the efficacy of other NS5A inhibitors, NS3/4A protease inhibitors, or NS5B inhibitors has not been studied.

Clinical efficacy and safety

Clinical studies in subjects with genotype 1 hepatitis C infection

The efficacy and safety of Viekirax in combination with dasabuvir with and without ribavirin was evaluated in eight Phase 3 clinical trials, including two trials exclusively in subjects with cirrhosis (Child-Pugh A), in over 2,360 subjects with genotype 1 chronic hepatitis C infection as summarised in Table 6.

Table 6. Phase 3 global multicentre studies conducted with Viekirax and dasabuvir with or without ribavirin (RBV).

Trial	Number of subjects treated	HCV genotype (GT)	Summary of study design
Treatment-naïve, with	out cirrhosis		
SAPPHIRE I	631	GT1	Arm A: Viekirax and dasabuvir + RBV Arm B: Placebo
PEARL III	419	GT1b	Arm A: Viekirax and dasabuvir + RBV Arm B: Viekirax and dasabuvir
PEARL IV	305	GT1a	Arm A: Viekirax and dasabuvir + RBV Arm B: Viekirax and dasabuvir
GARNET (open-label)	166	GT1b	Viekirax and dasabuvir (8 weeks)
Peginterferon+ribaviri	n experienced -, v	vithout cirrho	osis
SAPPHIRE II	394	GT1	Arm A: Viekirax and dasabuvir + RBV Arm B: Placebo
PEARL II (open-label)	179	GT1b	Arm A: Viekirax and dasabuvir + RBV Arm B: Viekirax and dasabuvir
Treatment-naïve and J	peginterferon+rib	avirin -exper	ienced, with compensated cirrhosis
TURQUOISE II (open-label)	380	GT1	Arm A: Viekirax and dasabuvir + RBV (12 weeks) Arm B: Viekirax and dasabuvir + RBV (24 weeks)
TURQUOISE III (open-label)	60	GT1b	Viekirax and dasabuvir (12 weeks)

In all eight trials, the Viekirax dose was 25 mg/150 mg/100 mg once daily and the dasabuvir dose was 250 mg twice daily. For subjects who received ribavirin, the ribavirin dose was 1000 mg per day for subjects weighing less than 75 kg or 1200 mg per day for subjects weighing greater than or equal to 75 kg.

Sustained virologic response (SVR) was the primary endpoint to determine the HCV cure rate in the Phase 3 studies and was defined as unquantifiable or undetectable HCV RNA 12 weeks after the end of treatment (SVR12). Treatment duration was fixed in each trial and was not guided by subjects' HCV RNA levels (no response guided algorithm). Plasma HCV RNA values were measured during the clinical trials using the COBAS TaqMan HCV test (version 2.0), for use with the High Pure System (except GARNET which used COBAS AmpliPrep/COBAS TaqMan HCV Test v2.0). The High Pure system assay had a lower limit of quantification (LLOQ) of 25 IU per mL and the AmpliPrep assay had a LLOQ of 15 IU per mL.

Clinical trials in treatment-naïve adults

<u>SAPPHIRE-I – genotype 1, treatment-naïve, without cirrhosis</u>

Design: randomised, global multicentre, double-blind, placebo-controlled Viekirax and dasabuvir with weight-based ribavirin for 12 weeks

Treated subjects (N=631) had a median age of 52 years (range: 18 to 70); 54.5% were male; 5.4% were Black; 15.2% had a history of depression or bipolar disorder; 79.1% had baseline HCV RNA levels of at least 800,000 IU/mL; 15.4% had portal fibrosis (F2) and 8.7% had bridging fibrosis (F3); 67.7% had HCV genotype 1a infection; 32.3% had HCV genotype 1b infection.

Table 7. SVR12 for genotype 1-infected treatment-naïve subjects in SAPPHIRE-I

Tuesday and sudsome	Viekirax and dasabuvir with RBV for 12 weeks					
Treatment outcome	n/N	% %	95% CI			
O 11 ON 7D 14						
Overall SVR12	456/473	96.4	94.7, 98.1			
HCV genotype 1a	308/322	95.7	93.4, 97.9			
HCV genotype 1b	148/151	98.0	95.8, 100.0			
Outcome for subjects without SVR12						
On-treatment VF ^a	1/473	0.2				
Relapse	7/463	1.5				
Other ^b	9/473	1.9				

a. Confirmed HCV \geq 25 IU/mL after HCV RNA < 25 IU/mL during treatment, confirmed 1 log₁₀ IU/mL increase in HCV RNA from nadir, or HCV RNA persistently \geq 25 IU/mL with at least 6 weeks of treatment.

No subjects with HCV genotype 1b infection experienced on-treatment virologic failure and one subject with HCV genotype 1b infection experienced relapse.

<u>PEARL-III – genotype 1b, treatment-naïve, without cirrhosis</u>

Design: randomised, global multicentre, double-blind, regimen-controlled

Treatment: Viekirax and dasabuvir without ribavirin or with weight-based ribavirin for 12 weeks

Treated subjects (N=419) had a median age of 50 years (range: 19 to 70), 45.8% were male; 4.8% were Black; 9.3% had a history of depression or bipolar disorder; 73.3% had baseline HCV RNA of at least 800,000 IU/mL; 20.3% had portal fibrosis (F2) and 10.0% had bridging fibrosis (F3).

Table 8. SVR12 for genotype 1b-infected treatment-naïve subjects in PEARL III

	Viekirax and dasabuvir for 12 weeks						
Treatment outcome		With RI	3V	Without RBV			
	n/N	%	95% CI	n/N	%	95% CI	
Overall SVR12	209/210	99.5	98.6, 100.0	20 9/209	100	98.2, 100.0	
Outcome for subjects without SVR12							
On-treatment VF	1/210	0.5		0/209	0		
Relapse	0/210	0		0/209	0		
Other	0/210	0		0/209	0		

PEARL-IV – genotype 1a, treatment-naïve, without cirrhosis

Design: randomised, global multicentre, double-blind, regimen-controlled

b. Other includes early drug discontinuation not due to virologic failure missing HCV RNA values in the SVR12 window.

Treatment: Viekirax and dasabuvir without ribavirin or with weight-based ribavirin for 12 weeks

Treated subjects (N=305) had a median age of 54 years (range: 19 to 70); 65.2% were male; 11.8% were Black; 20.7% had a history of depression or bipolar disorder; 86.6% had baseline HCV RNA levels of at least 800,000 IU/mL; 18.4% had portal fibrosis (F2) and 17.7% had bridging fibrosis (F3).

Table 9. SVR12 for genotype 1a-infected treatment-naïve subjects in PEARL IV

	Viekirax and dasabuvir for 12 weeks							
Treatment outcome		With 1	RBV	Without RBV				
Treatment outcome	n/N	%	95% CI	n/N	%	95% CI		
Overall SVR12	97/100	97.0	93.7, 100.0	185/205	90.2	86.2, 94.3		
Outcome for subjects without SVR12								
On-treatment VF	1/100	1.0		6/205	2.9			
Relapse	1/98	1.0		10/194	5.2			
Other	1/100	1.0		4/205	2.0			

GARNET – Genotype 1b, Treatment-Naïve without cirrhosis.

Design: open-label, single-arm, global multicentre

Treatment: Viekirax and dasabuvir for 8 weeks

Treated subjects (N=166) had a median age of 53 years (range: 22 to 82); 56.6% were female; 3.0% were Asian; 0.6% were Black; 7.2% had baseline HCV RNA levels of at least 6,000,000 IU per mL; 9% had advanced fibrosis (F3) and 98.2% had HCV genotype 1b infection (one subject each had genotype 1a, 1d, and 6 infection).

Table 10. SVR12 for Genotype 1b-infected treatment-naïve subjects without cirrhosis

	Viekirax and dasabuvir for 8 weeks n/N (%)	
SVR ₁₂	160/163 (98.2)	
95% CI ^a	96.1, 100.0	
F0-F1	138/139 (99.3) ^b	
F2	9/9 (100)	
F3	13/15 (86.7)°	

a. Calculated using the normal approximation to the binomial distribution

Clinical trials in peginterferon+ribavirin-experienced adults

<u>SAPPHIRE-II – genotype 1, pegIFN+RBV-experienced, without cirrhosis</u>

Design: randomised, global multicentre, double-blind, placebo-controlled Viekirax and dasabuvir with weight-based ribavirin for 12 weeks

b. 1 patient discontinued due to non-compliance

c. Relapse in 2/15 patients (confirmed HCV RNA \geq 15 IU/mL post-treatment before or during SVR12 window among subjects with HCV RNA \leq 15 IU/mL at last observation with at least 51 days of treatment).

Treated subjects (N=394) had a median age of 54 years (range: 19 to 71); 49.0% were prior pegIFN/RBV null responders; 21.8/% were prior pegIFN/RBV partial responders, and 29.2% were prior pegIFN/RBV relapsers; 57.6% were male; 8.1% were Black; 20.6% had a history of depression or bipolar disorder; 87.1% had baseline HCV RNA levels of at least 800,000 IU per mL; 17.8% had portal fibrosis (F2) and 14.5% had bridging fibrosis (F3); 58.4% had HCV genotype 1a infection; 41.4% had HCV genotype 1b infection.

Table 11. SVR12 for genotype 1-infected peginterferon+ribavirin-experienced subjects in SAPPHIRE-II

	Viekirax and dasabuvir with RBV for 12 weeks					
Treatment outcome	n/N	%	95% CI			
Overall SVR12	286/297	96.3	94.1, 98.4			
HCV genotype 1a	166/173	96.0	93.0, 98.9			
Prior pegIFN/RBV null responder	83/87	95.4	91.0, 99.8			
Prior pegIFN/RBV partial responder	36/36	100	100.0, 100.0			
Prior pegIFN/RBV relapser	47/50	94.0	87.4, 100.0			
HCV genotype 1b	119/123	96.7	93.6, 99.9			
Prior pegIFN/RBV null responder	56/59	94.9	89.3, 100.0			
Prior pegIFN/RBV partial responder	28/28	100	100.0, 100.0			
Prior pegIFN/RBV relapser	35/36	97.2	91.9, 100.0			
Outcome for subjects without SVR12						
On-treatment VF	0/297	0				
Relapse	7/293	2.4				
Other	4/297	1.3				

No subjects with HCV genotype 1b infection experienced on-treatment virologic failure and 2 subjects with HCV genotype 1b infection experienced relapse.

PEARL-II – *genotype 1b*, *pegIFN+RBV-experienced*, *without cirrhosis*

Design: randomised, global multicentre, open-label

Treatment: Viekirax and dasabuvir without ribavirin or with weight-based ribavirin for 12 weeks

Treated subjects (N=179) had a median age of 57 years (range: 26 to 70); 35.2% were prior pegIFN/RBV null responders; 28.5% were prior pegIFN/RBV partial responders, and 36.3% were prior pegIFN/RBV relapsers; 54.2% were male; 3.9% were Black; 12.8% had a history of depression or bipolar disorder; 87.7% had baseline HCV RNA levels of at least 800,000 IU/mL; 17.9% had portal fibrosis (F2) and 14.0% had bridging fibrosis (F3).

Table 12. SVR12 for genotype 1b-infected peginterferon+ribavirin-experienced subjects in PEARL II

	Viekirax and dasabuvir for 12 weeks						
Treatment outcome	With RBV			Without RBV			
	n/N	%	95% CI	n/N	%	95% CI	
Overall SVR12	86/88	97.7	94.6, 100.0	91/91	100	95.9, 100.0	
Prior pegIFN/RBV null responder	30/31	96.8	90.6, 100.0	32/32	100	89.3, 100.0	
Prior pegIFN/RBV partial responder	24/25	96.0	88.3, 100.0	26/26	100	87.1, 100.0	
Prior pegIFN/RBV relapser	32/32	100	89.3, 100.0	33/33	100	89.6, 100.0	
Outcome for subjects without SVR12							
On-treatment VF	0/88	0		0/91	0		
Relapse	0/88	0		0/91	0		
Other	2/88	2.3		0/91	0		

Clinical trial in subjects with compensated cirrhosis

TURQUOISE-II – treatment-naïve or pegIFN + RBV-experienced with compensated cirrhosis

Design: randomised, global multicentre, open-label

Treatment: Viekirax and dasabuvir with weight-based ribavirin for 12 or 24 weeks

Treated subjects (N=380) had a median age of 58 years (range: 21 to 71); 42.1% were treatment-naïve, 36.1% were prior pegIFN/RBV null responders; 8.2% were prior pegIFN/RBV partial responders, 13.7% were prior pegIFN/RBV relapsers; 70.3% were male; 3.2% were Black; 14.7% had platelet counts of less than 90 x 10⁹/L; 49.7% had albumin less than 40 g/L; 86.1% had baseline HCV RNA levels of at least 800,000 IU/mL; 24.7% had a history of depression or bipolar disorder; 68.7% had HCV genotype 1a infection, 31.3% had HCV genotype 1b infection.

Table 13. SVR12 for genotype 1-infected subjects with compensated cirrhosis who were treatment-naïve or previously treated with pegIFN/RBV

Treatment outcome	Viekirax and dasabuvir with RBV						
		12 week	s	24 weeks			
-	n/N	%	CI ^a	n/N	%	CIa	
Overall SVR12	191/208	91.8	87.6, 96.1	166/172	96.5	93.4, 99.6	
HCV genotype 1a	124/140	88.6	83.3, 93.8	115/121	95.0	91.2, 98.9	
Treatment naïve	59/64	92.2		53/56	94.6		
Prior pegIFN/RBV null responders	40/50	80.0		39/42	92.9		
Prior pegIFN/RBV partial responders	11/11	100		10/10	100		
Prior pegIFN/RBV Prior	14/15	93.3		13/13	100		
relapsers							
HCV genotype 1b	67/68	98.5	95.7, 100	51/51	100	93.0, 100	
Treatment naïve	22/22	100		18/18	100		
Prior pegIFN/RBV null responders	25/25	100		20/20	100		
Prior pegIFN/RBV partial responders	6/7	85.7		3/3	100		
Prior pegIFN/RBV Prior	14/14	100		10/10	100		
relapsers							
Outcome for subjects without SVR12							
On-treatment VF	1/208	0.5		3/172	1.7		
Relapse	12/203	5.9		1/164	0.6		
Other	4/208	1.9		2/172	1.21		

a. 97.5% confidence intervals are used for the primary efficacy endpoints (overall SVR12 rate); 95% confidence intervals are used for additional efficacy endpoints (SVR12 rates in HCV genotype 1a and 1b-infected subjects).

Relapse rates in GT1a cirrhotic subjects by baseline laboratory values are presented in Table 14.

Table 14. TURQUOISE-II: Relapse Rates by Baseline Laboratory Values after 12 and 24 Weeks of Treatment in Subjects with Genotype 1a Infection and Compensated Cirrhosis

	Viekirax and dasabuvir with RBV 12-week arm	Viekirax and dasabuvir with RBV 24-week arm
Number of Responders at the End of Treatment	135	113
AFP* $< 20 \text{ ng/mL}$, platelets $\ge 90 \times 10^9 / \text{L}$, AND alb	pumin \geq 35 g/L prior to treat	atment
Yes (for all three parameters listed above)	1/87 (1%)	0/68 (0%)
No (for any parameter listed above)	10/48 (21%)	1/45 (2%)
*AFP= serum alpha fetoprotein		

In subjects with all three favourable baseline laboratory values (AFP \leq 20 ng/mL, platelets \geq 90 x 10⁹/L, and albumin \geq 35 g/L), relapse rates were similar in subjects treated for 12 or 24 weeks.

<u>TURQUOISE-III:</u> treatment-naïve or pegIFN + RBV-experienced with compensated cirrhosis

Design: global multicentre, open-label

Treatment: Viekirax and dasabuvir without ribavirin for 12 weeks

60 patients were randomized and treated, and 60/60 (100%) achieved SVR12. Main characteristics are shown below.

Table 15. Main demographics in TURQUOISE-III

Characteristics	N = 60
Age, median (range) years	60.5 (26-78)
Male gender, n (%)	37 (61)
Prior HCV Treatment:	
naïve, n (%)	27 (45)
Peg-IFN + RBV, n (%)	33 (55)
Baseline albumin, median g/L	40.0
< 35, n (%)	10 (17)
≥ 35, n (%)	50 (83)
Baseline platelet count, median (\times 10 ⁹ /L)	132.0
< 90, n (%)	13 (22)
≥ 90, n (%)	47 (78)

Pooled analyses of clinical trials

Durability of response

Overall, 660 subjects in Phase 2 and 3 clinical trials had HCV RNA results for both the SVR12 and SVR24 time points. Among these subjects, the positive predictive value of SVR12 on SVR24 was 99.8%.

Pooled efficacy analysis

In Phase 3 clinical trials, 1075 subjects (including 181 with compensated cirrhosis) with genotype 1 HCV infection received the recommended regimen (see section 4.2). Table 16 shows SVR rates for these subjects.

In subjects who received the recommended regimen, 97% achieved SVR overall (among which 181 subjects with compensated cirrhosis achieved 97% SVR), while 0.5% experienced virologic breakthrough and 1.2% experienced post-treatment relapse.

Table 16. SVR12 rates for recommended treatment regimens by patient population

		enotype 1b nd dasabuvir	HCV Genotype 1a Viekirax and dasabuvir with RBV	
	Without cirrhosis	With compensated cirrhosis	Without cirrhosis	With compensated cirrhosis
Treatment duration	12 weeks	12 weeks	12 weeks	24 weeks
Treatment-naïve	100% (210/210)	100% (27/27)	96% (403/420)	95% (53/56)
pegIFN + RBV experienced	100% (91/91)	100% (33/33)	96% (166/173)	95% (62/65)
Prior relapse	100% (33/33)	100% (3/3)	94% (47/50)	100% (13/13)
Prior partial response	100% (26/26)	100% (5/5)	100% (36/36)	100% (10/10)
Prior null response	100% (32/32)	100% (7/7)	95% (83/87)	93% (39/42)
Other pegIFN/RBV failures	0	100% (18/18)+	0	0
TOTAL	100% (301/301)	100% (60/60)	96% (569/593)	95% (115/121)

⁺Other types of pegIFN/RBV failure include less well documented non-response, relapse/breakthrough or other pegIFN failure.

Viekirax without ribavirin and without dasabuvir was also evaluated in genotype 1b infected subjects in Phase 2 studies M13-393 (PEARL-I) and M12-536. PEARL I was conducted in the US and Europe, M12-536 in Japan. The treatment-experienced subjects studied were primarily pegIFN/RBV null responders. The doses of ombitasvir, paritaprevir, ritonavir were 25 mg 150 mg, 100 mg once daily in PEARL-I, while the dose of paritaprevir was 100 mg or 150 mg in study M12-536. Treatment duration was 12 weeks for treatment naïve subjects, 12-24 weeks for treatment experienced subjects and 24 weeks for subjects with cirrhosis. Overall, 107 of 113 subjects without cirrhosis and 147 of 155 subjects with cirrhosis achieved SVR12 after 12-24 weeks of treatment.

Viekirax with ribavirin & without dasabuvir was evaluated for 12 weeks in genotype 1 treatment naive and treatment experienced non-cirrhotic subjects in a phase 2 study M11-652 (AVIATOR). The doses of paritaprevir were 100 mg and 200 mg and ombitasvir 25 mg. Ribavirin was dosed based on weight (1000 mg – 1200 mg per day). Overall, 72 of 79 treatment-naive subjects (45 of 52 GT1a and 27 of 27 GT1b) and 40 of 45 treatment-experienced subjects (21 of 26 GT1a and 19 of 19 GT1b) achieved SVR12 after 12 weeks of treatment.

Impact of ribavirin dose adjustment on probability of SVR

In Phase 3 clinical trials, 91.5% of subjects did not require ribavirin dose adjustments during therapy. In the 8.5% of subjects who had ribavirin dose adjustments during therapy, the SVR rate (98.5%) was comparable to subjects who maintained their starting ribavirin dose throughout treatment.

<u>TURQUOISE-I:</u> treatment-naïve or pegIFN + RBV-experienced with HCV GT1 or GT4/HIV-1 co-infection, without cirrhosis or with compensated cirrhosis

Design: randomised, global multicentre, open-label

Treatment: Viekirax with or without dasabuvir coadminstered with or without weight-based ribavirin for 12 or 24 weeks

See section 4.2 for dosing recommendations in HCV/HIV-1 co-infected patients. HCV GT1- or 4-infected subjects with HIV-1 coinfection were on a stable HIV-1 antiretroviral therapy (ART) regimen that included ritonavir-boosted atazanavir, raltegravir, dolutegravir (Part 2 only), or darunavir (Part 1b and Part 2 GT4 only)-, co-administered with a backbone of tenofovir plus emtricitabine or lamivudine.

Part 1 of the study was a Phase 2 pilot cohort consisting of 2 parts, Part 1a (63 subjects) and Part 1b (22 subjects). Part 2 was a Phase 3 cohort consisting of 233 subjects.

In Part 1a, all subjects received Viekirax and dasabuvir with ribavirin for 12 or 24 weeks. Treated subjects (N = 63) had a median age of 51 years (range: 31 to 69); 24% were Black; 19% had compensated cirrhosis; 67% were treatment-naïve; 33% had failed prior treatment with pegIFN/RBV; 89% had HCV genotype 1a infection.

In Part 1b, all subjects received Viekirax and dasabuvir with ribavirin for 12 weeks. Treated subjects (N = 22) had a median age of 54 years (range: 34 to 68); 41% were Black; 14% had compensated cirrhosis; 86% were HCV treatment-naïve; 14% had failed prior treatment with pegIFN/RBV; 68% had HCV genotype 1a infection.

In Part 2, subjects with HCV GT1 received Viekirax and dasabuvir with or without ribavirin for 12 or 24 weeks. Subjects with HCV GT4 received Viekirax with ribavirin for 12 or 24 Weeks. Treated subjects (N = 233) had a median age of 49 years (range: 26 to 69); 10% were Black; 12% had compensated cirrhosis; 66% were treatment-naïve; 32% had failed prior treatment with pegIFN/RBV; 2% had failed prior treatment with sofosbuvir.

Table 17 shows the primary efficacy analysis of SVR12 performed on subjects with HCV GT1/HIV-1 co-infection that received recommended regimen in Part 2 of the TURQUOISE-I study.

Table 17. Primary SVR12 Assessment for Part 2 Subjects with HCV GT1/HIV-1 co-infection in TURQUOISE-I

Endpoint	Viekirax and dasabuvir with/without ribavirin for 12 or 24 Weeks $N = 200^a$
SVR12, n/N (%) [95% CI]	194/200 (97.0) [93.6, 98.6]
Outcome for subjects not achieving SVR12	
On-treatment virologic failure	1
Post-treatment relapse	1
Other ^b	4

a. Includes all HCV GT1 subjects in Part 2 excluding Arm G subjects that did not receive recommended regimen.

b. Includes subjects who discontinued due to adverse event, loss to follow-up or subject withdrawal, and subjects with reinfection

Efficacy analyses performed on other parts of the study demonstrated similarly high SVR12 rates. In Part 1a, SVR12 was achieved by 29/31 (93.5%) subjects on the 12-week arm (95% CI: 79.3%, 98.2%) and by 29/32 (90.6%) subjects on the 24-week arm (95% CI: 75.8% – 96.8%). There was 1 relapse in the 12-week arm and 1 on-treatment virologic failure in the 24-week arm. In Part 1b, SVR12 was achieved by 22/22 (100%) subjects (95% CI: 85.1%, 100%). In Part 2, SVR12 was achieved by 27/28 (96.4%) subjects with HCV GT4/HIV-1 coinfection (95% CI: 82.3%, 99.4%) with no virologic failures.

The SVR12 rates in HCV/HIV-1 co-infected subjects were thus consistent with SVR12 rates in the phase 3 trials of HCV mono-infected subjects.

<u>CORAL-I: treatment-naïve or pegIFN + RBV-experienced, at least 3 months post liver transplant or 12 months post renal transplant</u>

Design: randomised, global multicentre, open-label

Treatment: Viekirax and dasabuvir for 12 or 24 weeks with or without ribavirin (investigator chosen dose) for GT1 and GT4 infection

In subjects with liver transplant, no cirrhosis and GT1 infection, patients were dosed with Viekirax and dasabuvir for 12-24 weeks, with and without RBV. Liver transplant subjects with cirrhosis were dosed with Viekirax and dasabuvir with RBV (GT1a for 24 weeks [n=4], GT1b for 12 weeks [n=2]). Subjects with renal transplant and no cirrhosis were dosed for 12 weeks (with RBV for GT1a [n=9], without RBV for GT1b [n=3]). Subjects with liver transplant and GT4 infection were dosed with Viekirax with RBV (non-cirrhotic for 12 weeks [n=2] and cirrhotic for 24 weeks [n=1]. The dose of ribavirin was left to the discretion of the investigator, with most subjects receiving 600 to 800 mg per day as a starting dose, and most subjects also receiving 600 to 800 mg per day at the end of treatment.

A total of 129 subjects were treated, 84 with GT1a, 41 with GT1b, 1 with GT1 other, 3 with GT4 infection. Overall, 61% had fibrosis stage F0-F1, 26% F2, 9% F3, and 4% F4. 61% had prior HCV treatment experience before transplant. For immunosuppressive medication, most subjects were taking tacrolimus (81%), with the remainder taking cyclosporine.

Among all GT1 subjects who were post liver transplant, 111/114 (97.4%) achieved SVR12; with 2 relapsing post treatment and 1 breakthrough on treatment. Among the GT1 subjects who were post renal transplant, 9/12 (75%) achieved SVR12; however, there were no virologic failures. All 3 (100%) subjects with GT4 infection who were post liver transplant achieved SVR12.

Clinical trial in patients receiving opioid substitution therapy

In a phase 2, multicentre, open-label, single arm study, 38 treatment-naïve or pegIFN/RBV treatment experienced, non-cirrhotic subjects with genotype 1 infection who were on stable doses of methadone (N=19) or buprenorphine +/- naloxone (N=19) received 12 weeks of Viekirax and dasabuvir with ribavirin. Treated subjects had a median age of 51 years (range: 26 to 64); 65.8% were male and 5.3% were Black. A majority (86.8%) had baseline HCV RNA levels of at least 800,000 IU/mL and a majority (84.2%) had genotype 1a infection; 15.8% had portal fibrosis (F2) and 5.3% had bridging fibrosis (F3); and 94.7% were naïve to prior HCV treatment.

Overall, 37 (97.4%) of 38 subjects achieved SVR12. No subjects experienced on-treatment virologic failure or relapse.

<u>RUBY-I</u>; treatment-naïve or pegIFN + RBV experienced with or without cirrhosis who have severe renal impairment or end stage renal disease (ESRD)

Design: multicentre, open-label

Treatment: Viekirax and dasabuvir with or without RBV for 12 or 24 weeks

Severe renal impairment or ESRD includes CKD Stage 4 defined as eGFR <30-15 mL/min/1.73 m² or CKD Stage 5 defined as <15 mL/min/1.73 m² or requiring haemodialysis. Treated subjects (N=68) had a median age of 58 years (range: 32-77 years); 83.8% were male; 58.8% were Black; 73.5% of subjects were infected with HCV GT1a; 75.0%% had Stage 5 CKD and 69.1% were on haemodialysis.

Sixty four of 68 (94.1%) subjects achieved SVR12. One subject experienced relapse at Post-Treatment Week 4, 2 subjects prematurely discontinued study drug and 1 subject had missing SVR12 data.

See also Section 4.8 for discussion of safety information for RUBY-I.

In another open-label phase 3b study evaluating 12 weeks of Viekirax with or without dasabuvir and without RBV in non-cirrhotic, treatment-naive GT1a and GT4 patients with CKD stage 4 or 5 (Ruby II), the SVR12 rate was 94.4% (17/18), with no subjects experiencing on-treatment virologic failure or relapse.

Clinical trials in subjects with genotype 4 chronic hepatitis C

<u>PEARL- I– genotype 4, treatment-naïve or pegIFN + RBV experienced without cirrhosis</u>

Design: randomised, global multicentre, open-label

Treatment: treatment naïve: Viekirax without ribavirin or with weight-based ribavirin for 12 weeks

pegIFN + RBV experienced: Viekirax with weight-based ribavirin for 12 weeks

Subjects (N=135) had a median age of 51 years (range: 19 to 70); 63,7% were treatment-naïve, 17.0% were prior pegIFN/RBV null responders, 6.7% were prior pegIFN/RBV partial responders, 12.6% were prior pegIFN/RBV relapsers; 65.2% were male; 8.9% were Black, 69.6% had baseline HCV RNA levels at least 800,000 IU/mL; 6.7% had bridging fibrosis (F3).

Table 18. SVR12 for genotype 4-infected, subjects who were treatment-naïve or previously treated with pegIFN/RBV in PEARL I

	Ombitasvir + paritaprevir + ritonavir* for 12 weeks					
Treatment outcome	Treatment-naïve With RBV		Treatment-naïve Without RBV		pegIFN + RBV- experienced With RBV	
Treatment outcome						
	n/N	%	n/N	%	n/N	%
Overall SVR12	42/42	100%	40/44	90.9%	49/49	100%
Outcome for subjects without SVR12						
On-treatment VF	0/42	0	1/44	2.3%	0/49	0
Relapse	0/42	0	2/44	4.5%	0/49	0
Other	0/42	0	1/44	2.3%	0/49	0

^{*} Ombitasvir tablets, paritaprevir tablets and ritonavir capsules administered separately.

AGATE-1 -treatment-naïve or pegIFN +RBV experienced patients with compensated cirrhosis

Design: randomised, global multicentre, open-label

Treatment: Viekirax with weight-based ribavirin for 12 or 16 weeks

Subjects had a median age of 56 years (range: 32 to 81); 50% were treatment-naïve, 28% were prior pegIFN/RBV null responders; 10% were prior pegIFN/RBV partial responders, 13% were prior pegIFN/RBV relapsers; 70% were male; 17% were Black; 73% had baseline HCV RNA levels of at least 800,000 IU per mL; 17% had platelet counts of less than 90 x 10⁹ per L; and 4% had albumin less than 3.5 mg per dL.

Table 19. SVR12 for HCV Genotype 4-Infected Subjects with Compensated Cirrhosis

	Ombitasvir + Paritaprevir + Ritonavir with RBV		
	12 Weeks	16 Weeks	
SVR12 % (n/N)	97% (57/59)	98% (60/61)	
Outcome for subjects without SVR12			
On-treatment virologic failure	2 (1/59)	0 (0/61)	
Post-treatment relapse	0 (0/57)	0 (0/59)	
Other	2 (1/59)	2 (1/61)	

Paediatric population

The European Medicines Agency has deferred the obligation to submit the results of studies with Viekirax in one or more subsets of the paediatric populations in the treatment of chronic hepatitis C (see section 4.2 for information on paediatric use).

5.2 Pharmacokinetic properties

The pharmacokinetic properties of the combination of Viekirax with dasabuvir have been evaluated in healthy adult subjects and in subjects with chronic hepatitis C. Table 20 shows mean C_{max} and AUC of Viekirax 25 mg/150 mg/100 mg once daily with dasabuvir 250 mg twice daily following multiple doses with food in healthy volunteers.

Table 20. Geometric mean C_{max} , AUC of multiple doses of Viekirax 150 mg/100 mg/25 mg once daily with dasabuvir 250 mg twice daily with food in healthy volunteers

	C _{max} (ng/ml) (% CV)	AUC (ng*hr/ml) (% CV)
Ombitasvir	127 (31)	1420 (36)
Paritaprevir	1470 (87)	6990 (96)
Ritonavir	1600 (40)	9470 (41)

Absorption

Ombitasvir, paritaprevir and ritonavir were absorbed after oral administration with mean T_{max} of approximately 4 to 5 hours. While ombitasvir exposures increased in a dose proportional manner, paritaprevir and ritonavir exposures increased in a more than dose proportional manner. Accumulation is

minimal for ombitasvir and approximately 1.5- to 2-fold for ritonavir and paritaprevir. Pharmacokinetic steady state for the combination is achieved after approximately 12 days of dosing.

The absolute bioavailability of ombitasvir and paritaprevir was approximately 50% when administered with food as Viekirax.

Effect of paritaprevir/ritonavir on ombitasvir and dasabuvir

In the presence of paritaprevir/ritonavir, dasabuvir exposures decreased by approximately 50% to 60% while ombitasvir exposures increased by 31-47%.

Effect of ombitasvir on paritaprevir/ritonavir and dasabuvir

In the presence of ombitasvir, paritaprevir exposures were minimally affected (5% to 27% change) while dasabuvir exposures increase by approximately 30%.

Effect of dasabuvir on paritaprevir/ritonavir and ombitasvir

In the presence of dasabuvir, paritaprevir exposures increased by 50% to 65% while there was no change in ombitasvir exposures.

Effects of food

Ombitasvir, paritaprevir and ritonavir should be administered with food. All clinical trials with ombitasvir, paritaprevir and ritonavir have been conducted following administration with food.

Food increased the exposure (AUC) of ombitasvir, paritaprevir and ritonavir by up to 82%, 211% and 49%, respectively relative to the fasting state. The increase in exposure was similar regardless of meal type (e.g., high-fat versus moderate-fat) or calorie content (approximately 600 Kcal versus approximately 1000 Kcal). To maximise absorption, Viekirax should be taken with food without regard to fat or calorie content.

Distribution

Ombitasvir, paritaprevir and ritonavir are highly bound to plasma proteins. Plasma protein binding is not meaningfully altered in subjects with renal or hepatic impairment. The blood to plasma concentration ratios in humans ranged from 0.6 to 0.8 indicating that ombitasvir and paritaprevir were preferentially distributed in the plasma compartment of whole blood. Ombitasvir was approximately 99.9% bound to human plasma proteins. Paritaprevir was approximately 97-98.6% bound to human plasma proteins. Ritonavir was greater than 99% bound to human plasma proteins.

In vitro data indicate that paritaprevir is a substrate for the human hepatic uptake transporters, OATP1B1 and OATP1B3.

Biotransformation

Ombitasvir

Ombitasvir is metabolised via amide hydrolysis followed by oxidative metabolism. Following a 25 mg single dose of ¹⁴C-ombitasvir given alone, unchanged parent drug accounted for 8.9% of total radioactivity in human plasma; a total of 13 metabolites were identified in human plasma. These metabolites are not expected to have antiviral activity or off-target pharmacologic activity.

Paritaprevir

Paritaprevir is metabolised predominantly by CYP3A4 and to a lesser extent CYP3A5. Following administration of a single 200 mg/100 mg oral dose of ¹⁴C paritaprevir /ritonavir to humans, the parent drug was the major circulating component, accounting for approximately 90% of the plasma radioactivity. At least 5 minor metabolites of paritaprevir have been identified in circulation that accounted for approximately 10% of plasma radioactivity. These metabolites are not expected to have antiviral activity.

Ritonavir

Ritonavir is predominantly metabolised by CYP3A and to a lesser extent, by CYP2D6. Nearly the entire plasma radioactivity after a single 600 mg dose of ¹⁴C-ritonavir oral solution in humans was attributed to unchanged ritonavir.

Elimination

Ombitasvir

Following dosing of ombitasvir/paritaprevir/ritonavir with or without dasabuvir, mean plasma half-life of ombitasvir was approximately 21 to 25 hours. Following a single 25 mg dose of ¹⁴C- ombitasvir approximately 90% of the radioactivity was recovered in faeces and 2% in urine. Unchanged parent drug accounted for 88% of total radioactivity recovered in faeces, indicating that biliary excretion is a major elimination pathway for ombitasvir.

Paritaprevir

Following dosing of ombitasvir/paritaprevir /ritonavir with or without dasabuvir, mean plasma half-life of paritaprevir was approximately 5.5 hours. Following a 200 mg ¹⁴C -paritaprevir dose with 100 mg ritonavir, approximately 88% of the radioactivity was recovered in faeces with limited radioactivity (8.8%) in urine. Metabolism as well as biliary excretion of parent drug contribute to the elimination of paritaprevir.

Ritonavir

Following dosing of ombitasvir/paritaprevir /ritonavir, mean plasma half-life of ritonavir was approximately 4 hours. Following a 600 mg dose of ¹⁴C -ritonavir oral solution, 86.4% of the radioactivity was recovered in the faeces and 11.3% of the dose was excreted in the urine.

In vitro interaction data

Ombitasvir and paritaprevir do not inhibit organic anion transporter (OAT1) *in vivo* and are not expected to inhibit organic cation transporters (OCT1 and OCT2), organic anion transporters (OAT3), or multidrug and toxin extrusion proteins (MATE1 and MATE2K) at clinically relevant concentrations. Ritonavir does not inhibit OAT1 and is not expected to inhibit OCT2, OAT3, MATE1 and MATE2K at clinically relevant concentrations.

Special populations

Elderly

Based on population pharmacokinetic analysis of data from Phase 3 clinical studies, a 10 year increase or decrease in age from 54 years (median age in the Phase 3 studies) would result in approximately 10% change in ombitasvir exposures, and $\leq 20\%$ change in paritaprevir exposures. There is no pharmacokinetic information in patients ≥ 75 years.

Sex or body weight

Based on population pharmacokinetic analysis of data from Phase 3 clinical studies, female subjects would have approximately 55% higher, 100% higher and 15% higher ombitasvir, paritaprevir and ritonavir exposures than male subjects. However, no dose-adjustment based on gender is warranted. A 10 kg change in body weight from 76 kg (median weight in the Phase 3 studies) would results in <10% change in ombitasvir exposures, and no change in paritaprevir exposures. Body weight is not a significant predictor of ritonavir exposures.

Race or ethnicity

Based on population pharmacokinetic analysis of data from Phase 3 clinical studies, Asian subjects had 18% to 21% higher ombitasvir exposures, and 37% to 39% higher paritaprevir exposures than non-Asian subjects. The ritonavir exposures were comparable between Asians and non-Asians.

Renal impairment

The changes in ombitasvir, paritaprevir, and ritonavir exposures in subjects with mild, moderate and severe renal impairment are not considered to be clinically significant. Limited data in patients with end-stage renal disease indicate no clinically significant changes in exposure also in this patient group. No dose adjustment of Viekirax with and without dasabuvir is required for patients with mild, moderate or severe renal impairment, or end-stage-renal disease on dialysis (see section 4.2).

Pharmacokinetics of the combination of ombitasvir 25 mg, paritaprevir 150 mg, and ritonavir 100 mg, with or without dasabuvir 400 mg were evaluated in subjects with mild (CrCl: 60 to 89 ml/min), moderate (CrCl: 30 to 59 ml/min) and severe (CrCl: 15 to 29 ml/min) renal impairment.

Following administration of Viekirax and dasabuvir

Compared to the subjects with normal renal function, ombitasvir exposures were comparable in subjects with mild, moderate and severe renal impairment. Compared to the subjects with normal renal function, paritaprevir C_{max} values were comparable, but AUC values were 19%, 33% and 45% higher in mild, moderate and severe renal impairment, respectively. Ritonavir plasma concentrations increased when renal function was reduced: C_{max} and AUC values were 26% to 42% higher, 48% to 80% higher and 66% to 114% higher in subjects with mild, moderate and severe renal impairment, respectively.

Following administration of Viekirax

Following administration of Viekirax, the changes in ombitasvir, paritaprevir, and ritonavir exposures in subjects with mild, moderate and severe renal impairment were similar to those observed when Viekirax was administered with dasabuvir, and are not considered to be clinically significant.

Hepatic impairment

Following administration of Viekirax and dasabuvir

Pharmacokinetics of the combination of ombitasvir 25 mg, paritaprevir 200 mg, and ritonavir 100 mg, with dasabuvir 400 mg were evaluated in non-HCV infected subjects with mild (Child-Pugh A), moderate (Child-Pugh B) and severe (Child-Pugh C) hepatic impairment.

In subjects with mild hepatic impairment, paritaprevir, ritonavir and ombitasvir mean C_{max} and AUC values decreased by 29% to 48%, 34% to 38% and up to 8%, respectively, compared to subjects with normal hepatic function.

In subjects with moderate hepatic impairment, ombitasvir and ritonavir mean C_{max} and AUC values decreased by 29% to 30% and 30 to 33%, respectively, while paritaprevir mean C_{max} and AUC values increased by 26% to 62% compared to subjects with normal hepatic function. (see sections 4.2, 4.4, and 4.8).

In subjects with severe hepatic impairment, paritaprevir mean C_{max} and AUC values increased by 3.2-to 9.5-fold; ritonavir mean C_{max} values were 35% lower and AUC values were 13% higher and ombitasvir mean C_{max} and AUC values decreased by 68% and 54%, respectively, compared to subjects with normal hepatic function, therefore, Viekirax must not be used in patients with severe hepatic impairment (see sections 4.2 and 4.4).

In HCV-infected subjects, in comparison to those without cirrhosis, paritaprevir AUC increased to 2.2- to 2.4-fold for those with compensated cirrhosis (Child-Pugh A) and 3- to 4-fold for those with Child-Pugh B cirrhosis.

Following administration of Viekirax

Pharmacokinetics of the combination of ombitasvir 25 mg, paritaprevir 200 mg, and ritonavir 100 mg were not evaluated in subjects with mild (Child-Pugh A), moderate (Child-Pugh B) and severe (Child-Pugh C) hepatic impairment. Results from the pharmacokinetic evaluation of the combination of ombitasvir 25 mg, paritaprevir 200 mg, and ritonavir 100 mg, with dasabuvir 400 mg can be extrapolated to the combination of ombitasvir 25 mg, paritaprevir 200 mg, and ritonavir 100 mg.

Paediatric population

The pharmacokinetics of Viekirax in paediatric patients has not been established (see section 4.2).

5.3 Preclinical safety data

Ombitasvir

Ombitasvir and its major inactive human metabolites (M29, M36) were not genotoxic in a battery of *in vitro* or *in vivo* assays, including bacterial mutagenicity, chromosome aberration using human peripheral blood lymphocytes and *in vivo* mouse micronucleus assays.

Ombitasvir was not carcinogenic in a 6-month transgenic mouse study up to the highest dosage tested (150 mg/kg/day), resulting in ombitasvir AUC exposures approximately 26-fold higher than those in humans at the recommended clinical dose of 25 mg.

Similarly, ombitasvir was not carcinogenic in a 2-year rat study up to the highest dose tested (30 mg per kg per day), resulting in ombitasvir exposures approximately 16-fold higher than those in humans at 25 mg.

Ombitasvir has shown malformations in rabbits at maximal feasible exposures 4-fold higher than the AUC exposure at recommended clinical dose. Malformations at low incidence were observed mainly in the eyes (microphthalmia) and teeth (absent incisors). In mice, an increased incidence of open eye lid was present in foetuses of dams administered ombitasvir; however, the relationship to treatment with ombitasvir is uncertain. The major, inactive human metabolites of ombitasvir were not teratogenic in mice at exposures approximately 26 times higher than in humans at the recommended clinical dose. Ombitasvir had no effect on fertility when evaluated in mice.

Unchanged ombitasvir was the predominant component observed in the milk of lactating rats, without effect on nursing pups. Ombitasvir-derived material was minimally transferred through the placenta in pregnant rats.

Paritaprevir/ritonavir

Paritaprevir was positive in an *in vitro* human chromosome aberration test. Paritaprevir was negative in a bacterial mutation assay, and in two *in vivo* genetic toxicology assays (rat bone marrow micronucleus and rat liver Comet tests).

Paritaprevir /ritonavir was not carcinogenic in a 6-month transgenic mouse study up to the highest dosage tested (300 mg/30 mg/kg/day), resulting in paritaprevir AUC exposures approximately 38-fold higher than those in humans at the recommended dose of 150 mg. Similarly, paritaprevir/ritonavir was not carcinogenic in a 2-year rat study up to the highest dosage tested (300 mg/30 mg/kg/day), resulting in paritaprevir AUC exposures approximately 8-fold higher than those in humans at 150 mg.

Paritaprevir/ritonavir has shown malformations (open eye lids) at a low incidence in mice at exposures 32/8-fold higher than the exposure in humans at the recommended clinical dose. Paritaprevir/ritonavir had no effects on embryo-foetal viability or on fertility when evaluated in rats at exposures 2- to 8-fold higher than the exposure in humans at the recommended clinical dose.

Paritaprevir and its hydrolysis product M13 were the predominant components observed in the milk of lactating rats, without effect on nursing pups. Paritaprevir -derived material was minimally transferred through the placenta in pregnant rats.

6. PHARMACEUTICAL PARTICULARS

6.1 List of excipients

Tablet core

Copovidone
Tocofersolan
Propylene glycol monolaurate
Sorbitan monolaurate
Colloidal anhydrous silica (E 551)
Sodium stearyl fumarate

Film-coating

Poly(vinyl alcohol) (E 1203) Macrogol (3350) Talc (E 553b) Titanium dioxide (E 171) Iron oxide red (E 172)

6.2 Incompatibilities

Not applicable.

6.3 Shelf life

3 years.

6.4 Special precautions for storage

This medicinal product does not require any special storage conditions.

6.5 Nature and contents of container

PVC/PE/PCTFE aluminium foil blister packs. Pack-size of 56 tablets (multipack carton containing 4 inner cartons of 14 tablets each).

6.6 Special precautions for disposal

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

7. MARKETING AUTHORISATION HOLDER

AbbVie Deutschland GmbH & Co. KG Knollstrasse 67061 Ludwigshafen Germany

8. MARKETING AUTHORISATION NUMBER(S)

EU/1/14/982/001

9. DATE OF FIRST AUTHORISATION/RENEWAL OF THE AUTHORISATION

Date of first authorisation: 15 January 2015 Date of latest renewal: 19 September 2019

10. DATE OF REVISION OF THE TEXT

Detailed information on this medicinal product is available on the website of the European Medicines Agency http://www.ema.europa.eu.

ANNEX II

- A. MANUFACTURER(S) RESPONSIBLE FOR BATCH RELEASE
- B. CONDITIONS OR RESTRICTIONS REGARDING SUPPLY AND USE
- C. OTHER CONDITIONS AND REQUIREMENTS OF THE MARKETING AUTHORISATION
- D. CONDITIONS OR RESTRICTIONS WITH REGARD TO THE SAFE AND EFFECTIVE USE OF THE MEDICINAL PRODUCT

A. MANUFACTURER(S) RESPONSIBLE FOR BATCH RELEASE

AbbVie Deutschland GmbH & Co. KG Knollstrasse 67061 Ludwigshafen GERMANY

B. CONDITIONS OR RESTRICTIONS REGARDING SUPPLY AND USE

Medicinal product subject to restricted medical prescription (see Annex I: Summary of Product Characteristics, section 4.2).

C. OTHER CONDITIONS AND REQUIREMENTS OF THE MARKETING AUTHORISATION

• Periodic safety update reports (PSURs)

The requirements for submission of PSURs for this medicinal product are set out in the list of Union reference dates (EURD list) provided for under Article 107c(7) of Directive 2001/83/EC and any subsequent updates published on the European medicines web-portal.

D. CONDITIONS OR RESTRICTIONS WITH REGARD TO THE SAFE AND EFFECTIVE USE OF THE MEDICINAL PRODUCT

• Risk management plan (RMP)

The marketing authorisation holder (MAH) shall perform the required pharmacovigilance activities and interventions detailed in the agreed RMP presented in Module 1.8.2 of the marketing authorisation and any agreed subsequent updates of the RMP.

An updated RMP should be submitted:

- At the request of the European Medicines Agency;
- Whenever the risk management system is modified, especially as the result of new information being received that may lead to a significant change to the benefit/risk profile or as the result of an important (pharmacovigilance or risk minimisation) milestone being reached.

ANNEX III LABELLING AND PACKAGE LEAFLET

A. LABELLING

PARTICULARS TO APPEAR ON THE OUTER PACKAGING
Outer carton of multipack containing 56 (4 packs of 14) film-coated tablets - including blue box
, , , , , , , , , , , , , , , , , , , ,
1. NAME OF THE MEDICINAL PRODUCT
Viekirax 12.5 mg / 75 mg / 50 mg film-coated tablets ombitasvir / paritaprevir / ritonavir
emerus (II / parampre (II / Ittelia (II
2. STATEMENT OF ACTIVE SUBSTANCE(S)
Each film control tablet contains 12.5 mg of ambitagyin 75 mg of paritannovin and 50 mg of ritangyin
Each film coated tablet contains 12.5 mg of ombitasvir, 75 mg of paritaprevir and 50 mg of ritonavir.
3. LIST OF EXCIPIENTS
J. EIST OF EACHTENTS
4. PHARMACEUTICAL FORM AND CONTENTS
4. THARMACEUTICAL FORM AND CONTENTS
Multipack: 56 (4 packs of 14) film-coated tablets
5. METHOD AND ROUTE(S) OF ADMINISTRATION
Read the package leaflet before use.
Oral use
Take two tablets in the morning
C SDECIAL WARNING THAT THE MEDICINAL BRODUCT MUST BE STORED OUT OF
6. SPECIAL WARNING THAT THE MEDICINAL PRODUCT MUST BE STORED OUT OF THE SIGHT AND REACH OF CHILDREN
Keep out of the sight and reach of children.
Reep out of the sight and reach of children.
7 OTHER SPECIAL WARNING(S) IF NECESSARV
7. OTHER SPECIAL WARNING(S), IF NECESSARY
8. EXPIRY DATE
8. EXPIRY DATE

10. SPECIAL PRECAUTIONS FOR DISPOSAL OF UNUSED MEDICINAL PRODUCTS OR WASTE MATERIALS DERIVED FROM SUCH MEDICINAL PRODUCTS, IF APPROPRIATE

11.	NAME AND ADDRESS OF THE MARKETING AUTHORISATION HOLDER
Knol	Vie Deutschland GmbH & Co. KG Istrasse 1 Ludwigshafen nany
12.	MARKETING AUTHORISATION NUMBER(S)
EU/1	/14/982/001
13.	BATCH NUMBER
Lot	
14.	GENERAL CLASSIFICATION FOR SUPPLY
15.	INSTRUCTIONS ON USE
16.	INFORMATION IN BRAILLE
vieki	rax
17.	UNIQUE IDENTIFIER – 2D BARCODE
2D ba	arcode carrying the unique identifier included.
18.	UNIQUE IDENTIFIER - HUMAN READABLE DATA
PC: SN: NN:	

PARTICULARS TO APPEAR ON THE OUTER PACKAGING
Inner carton of multipack of 14 film-coated tablets – without blue box
1. NAME OF THE MEDICINAL PRODUCT
Viekirax 12.5 mg / 75 mg / 50 mg film-coated tablets ombitasvir / paritaprevir / ritonavir
2. STATEMENT OF ACTIVE SUBSTANCE(S)
Each film coated tablet contains 12.5 mg of ombitasvir, 75 mg of paritaprevir and 50 mg of ritonavir.
3. LIST OF EXCIPIENTS
4. PHARMACEUTICAL FORM AND CONTENTS
14 film-coated tablets Component of a multipack, can't be sold separately.
5. METHOD AND ROUTE(S) OF ADMINISTRATION
Read the package leaflet before use. Oral use Take two tablets in the morning.
6. SPECIAL WARNING THAT THE MEDICINAL PRODUCT MUST BE STORED OUT OF THE SIGHT AND REACH OF CHILDREN
Keep out of the sight and reach of children.
7. OTHER SPECIAL WARNING(S), IF NECESSARY
8. EXPIRY DATE
EXP
9. SPECIAL STORAGE CONDITIONS

10. SPECIAL PRECAUTIONS FOR DISPOSAL OF UNUSED MEDICINAL PRODUCTS OR WASTE MATERIALS DERIVED FROM SUCH MEDICINAL PRODUCTS, IF APPROPRIATE

11. NAME AND ADDRESS OF THE MARKETING AUTHORISATION HOLDER
AbbVie Deutschland GmbH & Co. KG Knollstrasse 67061 Ludwigshafen Germany
12. MARKETING AUTHORISATION NUMBER(S)
EU/1/14/982/001
13. BATCH NUMBER
Lot
14. GENERAL CLASSIFICATION FOR SUPPLY
15. INSTRUCTIONS ON USE
16. INFORMATION IN BRAILLE
viekirax
17. UNIQUE IDENTIFIER – 2D BARCODE
18. UNIQUE IDENTIFIER - HUMAN READABLE DATA

MINIMUM PARTICULARS TO APPEAR ON BLISTERS OR STRIPS
BLISTER FOIL
1 NAME OF THE MEDICINAL PRODUCT
1. NAME OF THE MEDICINAL PRODUCT
Viekirax 12.5 mg / 75 mg / 50 mg tablets
ombitasvir / paritaprevir / ritonavir
2. NAME OF THE MARKETING AUTHORISATION HOLDER
AbbVie (as logo)
3. EXPIRY DATE
EXP
4. BATCH NUMBER
Lot
5. OTHER

B. PACKAGE LEAFLET

Package leaflet: Information for the patient

Viekirax 12.5 mg/75 mg/50 mg film-coated tablets

ombitasvir/paritaprevir/ritonavir

Read all of this leaflet carefully before you start taking this medicine because it contains important information for you.

- Keep this leaflet. You may need to read it again.
- If you have any further questions, ask your doctor or pharmacist.
- This medicine has been prescribed for you only. Do not pass it on to others. It may harm them, even if their signs of illness are the same as yours.
- If you get any side effects, talk to your doctor or pharmacist. This includes any possible side effects not listed in this leaflet. See section 4.

What is in this leaflet

- 1. What Viekirax is and what it is used for
- 2. What you need to know before you take Viekirax
- 3. How to take Viekirax
- 4. Possible side effects
- 5. How to store Viekirax
- 6. Contents of the pack and other information

1. What Viekirax is and what it is used for

Viekirax contains the active substances ombitasvir, paritaprevir and ritonavir. It is an antiviral medicine used to treat adults with chronic (long-term) hepatitis C (an infectious disease that affects the liver, caused by the hepatitis C virus).

The combined action of the three active substances stops the hepatitis C virus from multiplying and infecting new cells, thus clearing the virus from your blood over a period of time. Ombitasvir and paritaprevir block two proteins essential for the virus to multiply. Ritonavir acts as a 'booster' to prolong the action of paritaprevir in the body.

Viekirax tablets are taken with other antiviral medicines such as dasabuvir and ribavirin. Your doctor will talk with you about which of these medicines to take with Viekirax.

It is very important that you also read the package leaflets for the other antiviral medicines that you take with Viekirax. If you have any questions about your medicines, please ask your doctor or pharmacist.

2. What you need to know before you take Viekirax

Do not take Viekirax

- If you are allergic to ombitasvir, paritaprevir, ritonavir, or any of the other ingredients of this medicine (listed in section 6).
- If you have moderate to severe liver problems other than hepatitis C.
- If you are taking any of the medicines listed in the following table. This is because serious or life-threatening effects can occur when Viekirax is taken with these medicines. These medicines can affect the way Viekirax works and Viekirax can affect the way these other medicines work.

Medicines you must not take with Viekirax	
Medicine or active substance	Purpose of the medicine
alfuzosin	for an enlarged prostate
amiodarone, disopyramide, dronedarone	used to correct irregular heartbeats
astemizole, terfenadine	for allergy symptoms. These medicines may
,	be available without a prescription
atorvastatin, lovastatin, simvastatin,	to lower blood cholesterol
lomitapide	
carbamazepine, phenytoin, phenobarbital	for epilepsy
cisapride	for relieving certain stomach problems
clarithromycin, fusidic acid, rifampicin,	for bacterial infections
telithromycin	
colchicine in patients who have severe	for treating gout attacks
problems with their liver or kidneys	
conivaptan	for making the sodium levels in the blood
•	normal
efavirenz, etravirine, lopinavir/ritonavir,	for HIV infection
saquinavir, tipranavir, nevirapine, indinavir,	
cobicistat	
apalutamide,enzalutamide	for prostate cancer
ergotamine, dihydroergotamine	for migraine headaches
ergonovine, methylergometrine	used in childbirth
ethinyloestradiol-containing medicines such	for contraception
as those contained in most contraceptive pills	•
and vaginal rings used for contraception	
itraconozole, ketoconozole, posaconazole,	for fungal infections
voriconazole	
midazolam, triazolam (when taken by mouth)	for anxiety or trouble sleeping
mitotane	for symptoms of malignant tumours of the
	adrenal glands
pimozide, lurasidone	for schizophrenia
quetiapine	for schizophrenia, bipolar disorder and major
	depressive disorder
quinidine	for abnormal heart rhythms or malaria
ranolazine	for chronic angina (chest pain)
salmeterol	for asthma
sildenafil	when used to treat a heart and lung disorder
	called "pulmonary arterial hypertension"
St. John's Wort (hypericum perforatum)	a herbal medicine for anxiety and mild
	depression. This medicine is available
	without a prescription
ticagrelor	stops blood from clotting
	-

Do not take Viekirax if any of the above apply to you. If you are not sure, talk to your doctor or pharmacist before taking Viekirax.

Warnings and precautions

Talk to your doctor or pharmacist before taking Viekirax if you:

- have liver disease other than hepatitis C;
- -have a current or previous infection with the hepatitis B virus, since your doctor may want to monitor you more closely.

-have diabetes. You may need closer monitoring of your blood glucose levels and/or adjustment of your diabetes medicines after starting Viekirax. Some diabetic patients have experienced low sugar levels in the blood (hypoglycaemia) after starting treatment with medicines like Viekirax.

When taking Viekirax with dasabuvir, tell your doctor if you have the following symptoms as they may be a sign of worsening liver problems:

- Feel sick (nauseous), are sick (vomit) or lose your appetite
- Notice yellowing of your skin or eyes
- Your urine is darker than normal
- Confusion
- Notice swelling of your stomach area

If any of the above apply to you (or you are not sure), talk to your doctor or pharmacist before taking Viekirax.

Tell your doctor if you have a history of depression or psychiatric illness. Depression, including suicidal thoughts and behaviours, has been reported in some patients taking this medicine, particularly in patients with a prior history of depression or psychiatric illness or in patients taking ribavirin with this medicine. You or your caregiver should also immediately inform your doctor of any changes in behaviour or mood and of any suicidal thoughts you may have.

Blood tests

Your doctor will test your blood before, during and after your treatment with Viekirax. This is so that your doctor can:

- Decide what other medicines you should take with Viekirax and for how long.
- Confirm if your treatment has worked and if you are free of the hepatitis C virus.
- Check for side effects of Viekirax or other antiviral medicines your doctor has prescribed for you to use with Viekirax (such as "dasabuvir" and "ribavirin").

Children and adolescents

Do not give Viekirax to children and adolescents under 18 years of age. The use of Viekirax in children and adolescents has not yet been studied.

Other medicines and Viekirax

Tell your doctor or pharmacist if you are taking, have recently taken and before starting any other medicines.

There are some medicines you **must not take** with Viekirax see the previous table "Medicines you must not take with Viekirax".

Tell your doctor or pharmacist before taking Viekirax, if you are taking any of the medicines in the table below. The doctor may need to change your dose of these medicines. Tell your doctor or pharmacist before taking Viekirax also if you are using hormonal contraceptives. See the section on contraception below.

Medicines you must tell your doctor about before taking Viekirax	
Medicine or active substance	Purpose of the medicine
alprazolam, diazepam	for anxiety, panic attacks and trouble sleeping
ciclosporin, everolimus, sirolimus, tacrolimus	to suppress the immune system
cyclobenzaprine, carisoprodol	for muscle spasms
colchicine for patients whose kidney and liver	for treating gout attacks or familial
function test are normal	Mediterranean fever
digoxin, amlodipine, nifedipine, valsartan,	for heart problems or high blood
diltiazem, verapamil, candesartan, losartan	pressure
encorafenib	for skin cancer
furosemide	for the build-up of too much fluid in the body
fostamatinib	for low platelet counts
hydrocodone	for pain
levothyroxine	for thyroid problems
rilpivirine, darunavir, atazanavir	for HIV infection
omeprazole, lansoprazole, esomeprazole	for stomach ulcers and other stomach
	problems
ibrutinib, imatinib	for the treatment of some cancers of the blood
fluvastatin, pitavastatin, pravastatin, rosuvastatin	to lower blood cholesterol
dabigatran	to thin the blood
fexofenadine	for hay fever
s-mephenytoin	for epilepsy
sulfasalazine	for inflammatory bowel disease
repaglinide	for lowering blood sugar
erythromycin	for bacterial infections
steroid or corticosteroid medicines (such as	for many different conditions including
fluticasone)	serious illnesses and allergies
trazodone	for anxiety and depression
warfarin and other similar medicines called vitamin K antagonists*	to thin the blood

^{*}Your doctor may need to increase the frequency of your blood tests to check how well your blood can clot.

If any of the above apply to you (or you are not sure), talk to your doctor or pharmacist before taking Viekirax.

Pregnancy and contraception

The effects of Viekirax during pregnancy are not known. Viekirax should not be used during pregnancy or in women of childbearing potential not using effective contraception.

• You or your partner must use an effective method of contraception during treatment. Contraceptive medicines that contain ethinyloestradiol cannot be used in combination with Viekirax. Ask your doctor about the best contraception for you.

Extra precautions are needed if Viekirax is taken together with ribavirin. Ribavirin may cause severe birth defects. Ribavirin stays for a long time in the body after treatment is stopped, and effective contraception is therefore needed both during treatment and for some time afterwards.

- There is a risk for birth defects when ribavirin is given to a female patient that becomes pregnant.
- There may also be a risk for birth defects if ribavirin is taken by a male patient, whose female partner becomes pregnant.
- Read the "Contraception" section of the package leaflet for ribavirin very carefully. It is important that both men and women read the information.
- If you or your partner becomes pregnant during treatment with Viekirax and ribavirin or in the months that follow, you must contact your doctor immediately.

Breastfeeding

You should not breast-feed during treatment with Viekirax. It is not known whether the active substances in Viekirax (ombitasvir, paritaprevir and ritonavir) pass into breast milk.

Driving and using machines

Some patients have reported feeling very tired when taking Viekirax with other medicines for their hepatitis C infection. If you feel tired, do not drive or use machines.

3. How to take Viekirax

Always take this medicine exactly as your doctor has told you. Check with your doctor or pharmacist if you are not sure.

Viekirax tablets are usually taken with other anti-viral medicines such as "dasabuvir" and "ribavirin".

How much to take

The recommended dose is two tablets taken together in the morning.

How to take

- Take the tablets in the morning with food. The type of food is not important.
- Swallow the tablets whole with water.
- Do not chew, crush or break the tablets as they may have a bitter taste.

How long to take Viekirax for

You will take Viekirax for 8, 12 or 24 weeks. Your doctor will tell you how long your treatment will last. Do not stop taking Viekirax unless your doctor tells you to. It is very important that you complete the full course of treatment. This will give the medicines the best chance to clear the hepatitis C virus infection.

If you take more Viekirax than you should

If you accidentally take more than the recommended dose, you should contact your doctor or go to the nearest hospital straight away. Keep the medicine pack with you so that you can easily describe what you have taken.

If you forget to take Viekirax

It is important not to miss a dose of this medicine. If you do miss a dose and it is:

• More than 12 hours until your next dose - take the missed dose with food as soon as possible.

• Less than 12 hours until your next dose - do not take the missed dose, take your next dose as usual with food.

Do not take a double dose to make up for a forgotten dose.

If you have any further questions on the use of this medicine, ask your doctor or pharmacist.

4. Possible side effects

Stop taking Viekirax and talk to your doctor or get medical help immediately if any of the following occur:

Side effects when taking Viekirax with or without dasabuvir and with or without ribavirin:

Frequency not known: cannot be estimated based on available data

- Serious allergic reactions, signs may include:
 - o Difficulty breathing or swallowing
 - o Dizziness or light-headedness, which may be due to low blood pressure
 - O Swelling of the face, lips, tongue or throat
 - o Rash and itching of the skin
- Worsening liver problems. Symptoms include:
 - o Feel sick (nauseous), are sick (vomit) or lose your appetite
 - o Notice yellowing of your skin or eyes
 - Your urine is darker than normal
 - o Confusion
 - Notice swelling of your stomach area

Like all medicines, this medicine can cause side effects, although not everybody gets them.

Tell your doctor or pharmacist if you notice any of the following side effects.

Side effects when taking Viekirax with dasabuvir:

Common: may affect up to 1 in 10 people

• Itching.

Rare: may affect up to 1 in 1,000 people

• Swelling of the layers of skin which can affect any part of the body including the face, tongue or throat and may cause difficulty swallowing or breathing (angioedema)

Side effects when taking Viekirax with dasabuvir and ribavirin:

Very common: may affect more than 1 in 10 people

- Feeling very tired (fatigue)
- Feeling sick (nausea)
- Itching
- Trouble sleeping (insomnia)
- Feeling weak or lack of energy (asthenia)
- Diarrhoea

Common: may affect up to 1 in 10 people

- Anaemia (low number of red blood cells)
- Vomiting

Uncommon: may affect up to 1 in 100 people

Dehydration

Rare: may affect up to 1 in 1,000 people

• Swelling of the layers of skin which can affect any part of the body including the face, tongue or throat and may cause difficulty swallowing or breathing (angioedema)

Reporting of side effects

If you get any side effects, talk to your doctor or pharmacist. This includes any possible side effects not listed in this leaflet. You can also report side effects directly via the national reporting system listed in Appendix V. By reporting side effects you can help provide more information on the safety of this medicine.

5. How to store Viekirax

Keep this medicine out of the sight and reach of children.

Do not use the medicine after the expiry date which is stated on the carton after 'EXP'. The expiry date refers to the last day of that month.

This medicine does not require any special storage conditions.

Do not throw away any medicines via wastewater or household waste. Ask your pharmacist how to throw away medicines you no longer use. These measures will help to protect the environment.

6. Contents of the pack and other information

What Viekirax contains

- Each tablet contains 12.5 mg of ombitasvir, 75 mg of paritaprevir and 50 mg of ritonavir.
- The other ingredients are:
 - Tablet core: copovidone, tocofersolan, propylene glycol monolaurate, sorbitan monolaurate, colloidal anhydrous silica (E 551), sodium stearyl fumarate.
 - Tablet film-coating: poly(vinyl alcohol) (E 1203), macrogol (3350), talc (E 553b), titanium dioxide (E 171) and red iron oxide (E 172).

What Viekirax looks like and contents of the pack

Viekirax tablets are pink, oblong film-coated tablets of dimmensions 18.8 mm x 10.0 mm, marked with 'AV1'. Viekirax tablets are packed into foil blisters containing 2 tablets. Each carton contains 56 tablets (multipack carton containing 4 inner cartons of 14 tablets each).

Marketing Authorisation Holder and Manufacturer

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This leaflet was last revised in

Other sources of information

Detailed information on this medicine is available on the European Medicines Agency web site: http://www.ema.europa.eu.