Summary of Product Characteristics

1. NAME OF THE MEDICINAL PRODUCT

LEDVIR

(Ledipasvir/Sofosbuvir Tablets 90 mg/400 mg film-coated tablets)

2. QUALITATIVE AND QUANTITATIVE COMPOSITION

Each film coated tablets contains-: Ledipasvir.....90 mg Sofosbuvir.....400 mg

For the full list of excipients, see section 6.1..

3. PHARMACEUTICAL FORM

Film-coated tablet.

Light blue, Oval shaped, biconvex, beveled edge film-coated tablet debossed with "LSF" on one side of the tablet and "M" on the other side.

4. CLINICAL PARTICULARS

4.1 Therapeutic indications

Ledipasvir/Sofosbuvir 90 mg/400 mg film coated tablets is indicated 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

Ledipasvir/Sofosbuvir 90 mg/400 mg film coated tablets treatment should be initiated and monitored by a physician experienced in the management of patients with CHC.

Posology

The recommended dose of Ledipasvir/Sofosbuvir 90 mg/400 mg film coated tablets is one tablet once daily with or without food (see section 5.2).

Table 1: Recommended treatment duration for Ledipasvir/Sofosbuvir 90 mg/400 mg film coated tablets and the recommended use of co-administered ribavirin for certain subgroups

Patient population*	Treatment and duration
Patients with genotype 1, 4, 5 or 6 CHC	
Patients without cirrhosis	 Ledipasvir/Sofosbuvir 90 mg/400 mg for 12 weeks. Ledipasvir/Sofosbuvir 90 mg/400 mg for 8 weeks may be considered in previously untreated genotype 1-infected patients (see section 5.1, ION-3 study). Ledipasvir/Sofosbuvir 90 mg/400 mg + ribavirin for 12 weeks or Ledipasvir/Sofosbuvir 90 mg/400 mg (without ribavirin) for 24 weeks should be considered for previously treated patients with uncertain subsequent retreatment options (see section 4.4).
Patients with compensated cirrhosis	Ledipasvir/Sofosbuvir 90 mg/400 mg + ribavirin for 12 weeks or Ledipasvir/Sofosbuvir 90 mg/400 mg (without ribavirin) for 24 weeks. - Ledipasvir/Sofosbuvir 90 mg/400 mg (without ribavirin) for 12 weeks may be considered for patients deemed at low risk for clinical disease progression and who have subsequent retreatment options (see section 4.4).
Patients who are post-liver transplant without cirrhosis or with compensated cirrhosis	Ledipasvir/Sofosbuvir 90 mg/400 mg + ribavirin for 12 weeks (see section 5.1). - Ledipasvir/Sofosbuvir 90 mg/400 mg (without ribavirin) for 12 weeks (in patients without cirrhosis) or 24 weeks (in patients with cirrhosis) may be considered for patients who are ineligible for or intolerant to ribavirin.
Patients with decompensated cirrhosis, irrespective of transplant status	Ledipasvir/Sofosbuvir 90 mg/400 mg + ribavirin for 12 weeks (see section 5.1). - Ledipasvir/Sofosbuvir 90 mg/400 mg (without ribavirin) for 24 weeks may be considered in patients who are ineligible for or intolerant to ribavirin.
Patients with genotype 3 CHC	

Patients with cirrhosis and/or prior	Ledipasvir/Sofosbuvir 90 mg/400 mg +
treatment failure	ribavirin for 24 weeks (see sections 4.4
	and 5.1).

* Includes patients co-infected with human immunodeficiency virus (HIV).

When used in combination with ribavirin, refer also to the Summary of Product Characteristics of ribavirin.

In patients without decompensated cirrhosis requiring the addition of ribavirin to their treatment regimen (see Table 1), the daily dose of ribavirin is weight based (< 75 kg = 1,000 mg and $\ge 75 \text{ kg} = 1,200 \text{ mg}$) and administered orally in two divided doses with food.

In patients with decompensated cirrhosis, ribavirin should be administered at a starting dose of 600 mg given in a divided daily dose. If the starting dose is well-tolerated, the dose can be titrated up to a maximum of 1,000-1,200 mg daily (1,000 mg for patients weighing < 75 kg and 1,200 mg for patients weighing \geq 75 kg). If the starting dose is not well-tolerated, the dose should be reduced as clinically indicated based on haemoglobin levels.

Dose modification of ribavirin in patients taking 1,000-1,200 mg daily

If Ledipasvir/Sofosbuvir 90 mg/400 mg film coated tablets is used in combination with ribavirin and a patient has a serious adverse reaction potentially related to ribavirin, the ribavirin dose should be modified or discontinued, if appropriate, until the adverse reaction abates or decreases in severity. Table 2 provides guidelines for dose modifications and discontinuation based on the patient's haemoglobin concentration and cardiac status.

Table	2:	Ribavirin	dose	modification	guideline	for	co-administration	with
Ledipa	svir	/Sofosbuvi	r 90 mg	g/400 mg film c	oated table	ts		

Laboratory values	Reduce ribavirin dose to 600 mg/day if:	Discontinue ribavirin if:
Haemoglobin in patients with no cardiac disease	< 10 g/dL	< 8.5 g/dL
Haemoglobin in patients with history of stable cardiac disease		< 12 g/dL despite 4 weeks at reduced dose

Once ribavirin has been withheld due to either a laboratory abnormality or clinical manifestation, an attempt may be made to restart ribavirin at 600 mg daily and further increase the dose to 800 mg daily. However, it is not recommended that ribavirin be increased to the originally assigned dose (1,000 mg to 1,200 mg daily).

Patients should be instructed that if vomiting occurs within 5 hours of dosing an additional tablet should be taken. If vomiting occurs more than 5 hours after dosing, no further dose is needed (see section 5.1).

If a dose is missed and it is within 18 hours of the normal time, patients should be instructed to take the tablet as soon as possible and then patients should take the next dose at the usual time. If it is after 18 hours then patients should be instructed to wait and take the next dose at the usual time. Patients should be instructed not to take a double dose.

Elderly

No dose adjustment is warranted for elderly patients (see section 5.2).

Renal impairment

No dose adjustment of Ledipasvir/Sofosbuvir 90 mg/400 mg film coated tablets is required for patients with mild or moderate renal impairment. The safety of Ledipasvir/Sofosbuvir has not been assessed in patients with severe renal impairment (estimated glomerular filtration rate [eGFR] < 30 mL/min/1.73 m²) or end stage renal disease (ESRD) requiring haemodialysis (see section 5.2).

Hepatic impairment

No dose adjustment of Ledipasvir/Sofosbuvir 90 mg/400 mg film coated tablets is required for patients with mild, moderate or severe hepatic impairment (Child-Pugh-Turcotte [CPT] class A, B or C) (see section 5.2). Safety and efficacy of Ledipasvir/Sofosbuvir have been established in patients with decompensated cirrhosis (see section 5.1).

Paediatric population

The safety and efficacy of Ledipasvir/Sofosbuvir 90 mg/400 mg film coated tablets in children and adolescents aged less than 18 years have not yet been established. No data are available.

Method of administration For oral use.

Patients should be instructed to swallow the tablet whole with or without food. Due to the bitter taste, it is recommended that the film-coated tablet is not chewed or crushed (see section 5.2).

4.3 Contraindications

Hypersensitivity to the active substances or to any of the excipients listed in section 6.1. Coadministration with Rosuvastatin or St. John's wort (*Hypericum perforatum*) (see section 4.5).

4.4 Special warnings and precautions for use

Ledipasvir/Sofosbuvir 90 mg/400 mg film coated tablets should not be administered concomitantly with other medicinal products containing Sofosbuvir.

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 clinical data to support the use of Ledipasvir/Sofosbuvir 90 mg/400 mg film coated tablets in patients infected with HCV genotype 3 are limited

(see section 5.1). The relative efficacy of a 12-week regimen consisting of Ledipasvir/ Sofosbuvir + ribavirin, compared to a 24-week regimen of Sofosbuvir + ribavirin has not been investigated. A conservative 24 weeks of therapy is advised in all treatmentexperienced genotype 3 patients and those treatment-naïve genotype 3 patients with cirrhosis (see section 4.2). The clinical data to support the use of Ledipasvir/Sofosbuvir 90 mg/400 mg film coated tablets in patients infected with HCV genotype 4 are limited (see section 5.1).

The efficacy of Ledipasvir/Sofosbuvir has not been studied against HCV genotype 2, 5 and 6; therefore, Ledipasvir/Sofosbuvir 90 mg/400 mg film coated tablets should not be used in patients infected with these genotypes.

Severe bradycardia and heart block

Cases of severe bradycardia and heart block have been observed when Ledipasvir/ Sofosbuvir 90 mg/400 mg film coated tablets is used with concomitant amiodarone with or without other drugs that lower heart rate. The mechanism is not established.

The concomitant use of amiodarone was limited through the clinical development of sofosbuvir plus direct-acting antivirals (DAAs). Cases are potentially life threatening, therefore amiodarone should only be used in patients on Ledipasvir/Sofosbuvir 90 mg/400 mg film coated tablets when other alternative anti-arrhythmic treatments are not tolerated or are contraindicated.

Should concomitant use of amiodarone be considered necessary it is recommended that patients are closely monitored when initiating Ledipasvir/Sofosbuvir 90 mg/400 mg film coated tablets. Patients who are identified as being high risk of bradyarrhythmia should be continuously monitored for 48 hours in an appropriate clinical setting.

Due to the long half-life of amiodarone, appropriate monitoring should also be carried out for patients who have discontinued amiodarone within the past few months and are to be initiated on Ledipasvir/Sofosbuvir 90 mg/400 mg film coated tablets.

All patients receiving Ledipasvir/Sofosbuvir 90 mg/400 mg film coated tablets in combination with amiodarone with or without other drugs that lower heart rate should also be warned of the symptoms of bradycardia and heart block and should be advised to seek medical advice urgently should they experience them.

Treatment of patients with prior exposure to HCV direct-acting antivirals

In patients who fail treatment with Ledipasvir/Sofosbuvir, selection of NS5A resistance mutations that substantially reduce the susceptibility to Ledipasvir is seen in the majority of cases (see section 5.1). Limited data indicate that such NS5A mutations do not revert on long-term follow-up. There are presently no data to support the effectiveness of retreatment of patients who have failed Ledipasvir/Sofosbuvir with a subsequent regimen that contains an NS5A inhibitor. Similarly, there are presently no data to support the effectiveness of NS3/4A protease inhibitors in patients who previously failed prior therapy that included an NS3/4A protease inhibitor. Such patients may therefore be dependent on other drug classes for clearance of HCV infection. Consequently, consideration should be given to longer treatment for patients with uncertain subsequent retreatment options.

Renal impairment

No dose adjustment of Ledipasvir/Sofosbuvir 90 mg/400 mg film coated tablets is required for patients with mild or moderate renal impairment. The safety of Ledipasvir/Sofosbuvir 90 mg/400 mg film coated tablets has not been assessed in patients with severe renal impairment (estimated glomerular filtration rate [eGFR] < 30 mL/min/1.73 m²) or end stage renal disease (ESRD) requiring haemodialysis. When Ledipasvir/Sofosbuvir 90 mg/400 mg film coated tablets is used in combination with ribavirin refer also to the Summary of Product Characteristics for ribavirin for patients with creatinine clearance (CrCl) < 50 mL/min (see section 5.2).

Patients with decompensated cirrhosis and/or who are awaiting liver transplant or postliver transplant The relative efficacy of 12 and 24 weeks of therapy has not been established. Therefore, 24 weeks of therapy is recommended (see sections 4.2 and 5.1). Treatment with Ledipasvir/Sofosbuvir 90 mg/400 mg film coated tablets should be guided by an assessment of the potential benefits and risks for the individual patient.

Use with potent P-gp inducers

Medicinal products that are potent P-glycoprotein (P-gp) inducers (e.g. rifampicin, carbamazepine and phenytoin) may significantly decrease Ledipasvir and Sofosbuvir plasma concentration which may lead to reduced therapeutic effect of Ledipasvir/Sofosbuvir 90 mg/400 mg film coated tablets. Such medicinal products should not be used with Ledipasvir/Sofosbuvir 90 mg/400 mg film coated tablets (see section 4.5).

Use with certain HIV antiretroviral regimens

Ledipasvir/Sofosbuvir 90 mg/400 mg film coated tablets has been shown to increase tenofovir exposure, especially when used together with an HIV regimen containing Tenofovir Disoproxil Fumarate and a pharmacokinetic enhancer (Ritonavir or Cobicistat). The safety of tenofovir Disoproxil Fumarate in the setting of Ledipasvir/Sofosbuvir 90 mg/ 400 mg film coated tablets and a pharmacokinetic enhancer has not been established. The potential risks and benefits associated with co-administration of Ledipasvir/Sofosbuvir 90 mg/400 mg film coated tablets with the fixed-dose combination tablet containing Elvitegravir/Cobicistat/Emtricitabine/tenofovir Disoproxil Fumarate or tenofovir Disoproxil Fumarate given in conjunction with a boosted HIV protease inhibitor (e.g. Atazanavir or Darunavir) should be considered, particularly in patients at increased risk of renal dysfunction. Patients receiving Ledipasvir/Sofosbuvir 90 mg/400 mg film coated tablets

Elvitegravir/Cobicistat/Emtricitabine/tenofovir Disoproxil Fumarate or with tenofovir Disoproxil Fumarate and a boosted HIV protease inhibitor should be monitored for tenofovir associated adverse reactions. Refer to tenofovir Disoproxil Fumarate, Emtricitabine/tenofovir Disoproxil Fumarate, or Elvitegravir/Cobicistat/Emtricitabine/ tenofovir Disoproxil Fumarate Summary of Product Characteristics for recommendations on renal monitoring.

Use with HMG-CoA reductase inhibitors

Co-administration of Ledipasvir/Sofosbuvir 90 mg/400 mg film coated tablets and HMG-CoA reductase inhibitors (statins) can significantly increase the concentration of the statin, which increases the risk of myopathy and rhabdomyolysis (see section 4.5).

HCV/HBV (hepatitis B virus) co-infection

There are no data on the use of Ledipasvir/Sofosbuvir 90 mg/400 mg film coated tablets in patients with HCV/HBV co-infection.

Paediatric population

Ledipasvir/Sofosbuvir 90 mg/400 mg film coated tablets is not recommended for use in children and adolescents under 18 years of age because the safety and efficacy have not been established in this population.

Excipients

Ledipasvir/Sofosbuvir 90 mg/400 mg film coated tablets contains lactose. Consequently, patients with rare hereditary problems of galactose intolerance, the Lapp lactase deficiency, or glucose-galactose malabsorption should not take this medicinal product.

4.5 Interaction with other medicinal products and other forms of interaction

As Ledipasvir/Sofosbuvir 90 mg/400 mg film coated tablets contains Ledipasvir and Sofosbuvir, any interactions that have been identified with these active substances individually may occur with Ledipasvir/Sofosbuvir 90 mg/400 mg film coated tablets.

Potential for Ledipasvir/Sofosbuvir 90 mg/400 mg film coated tablets to affect other medicinal products

Ledipasvir is an *in vitro* inhibitor of drug transporter P-gp and breast cancer resistance protein (BCRP) and may increase intestinal absorption of co-administered substrates for these transporters. *In vitro* data indicate that Ledipasvir may be a weak inducer of metabolising enzymes such as CYP3A4, CYP2C and UGT1A1. Compounds that are substrates of these enzymes may have decreased plasma concentrations when co-administered with Ledipasvir/Sofosbuvir. *In vitro* Ledipasvir inhibits intestinal CYP3A4 and UGT1A1. Medicinal products that have a narrow therapeutic range and which are metabolised by these isoenzymes should be used with caution and carefully monitored.

Potential for other medicinal products to affect Ledipasvir/Sofosbuvir 90 mg/400 mg film coated tablets Ledipasvir and Sofosbuvir are substrates of drug transporter P-gp and BCRP while GS-331007 is not. Medicinal products that are potent P-gp inducers (e.g. rifampicin, St. John's wort, carbamazepine and phenytoin) may decrease Ledipasvir and Sofosbuvir plasma concentrations leading to reduced therapeutic effect of Ledipasvir/Sofosbuvir and should not be used with Ledipasvir/Sofosbuvir 90 mg/400 mg film coated tablets (see sections 4.3 and 4.4). Co-administration with medicinal products that inhibit P-gp and/or BCRP may increase Ledipasvir and Sofosbuvir plasma concentrations; Ledipasvir/Sofosbuvir 90 mg/400 mg film coated tablets may be coadministered with P-gp and/or BCRP inhibitors. Clinically significant medicinal product interactions with Ledipasvir/Sofosbuvir gord tablets may be coadministered with P-gp and/or BCRP inhibitors. Clinically significant medicinal product interactions with Ledipasvir/Sofosbuvir mediated by CYP450s or UGT1A1 enzymes are not expected.

Interactions between Ledipasvir/Sofosbuvir 90 mg/400 mg film coated tablets and other medicinal products

Table 3 provides a listing of established or potentially clinically significant medicinal product interactions (where 90% confidence interval [CI] of the geometric least-squares mean [GLSM] ratio were within " \leftrightarrow ", extended above " \uparrow ", or extended below " \downarrow " the predetermined equivalence boundaries). The medicinal product interactions described are based on studies conducted with either Ledipasvir/Sofosbuvir or Ledipasvir and Sofosbuvir as individual agents, or are predicted medicinal product interactions that may occur with Ledipasvir/Sofosbuvir. The table is not all-inclusive.

Table 3: Interactions between Ledipasvir/Sofosbuvir 90 mg/400 mg film coatedtablets and other medicinal products

Medicinal product by therapeutic areas	Effects on medicinal product levels. Mean ratio (90% confidence interval) for AUC, Cmax, Cmina, b	Recommendation concerning co- administration with Ledipasvir/ Sofosbuvir 90 mg/400 mg film coated tablets
ACID REDUCING AGENTS	;	
		Ledipasvir solubility decreases as pH increases. Medicinal products that increase gastric pH are expected to
Antacids		decrease concentration of Ledipasvir.
e.g. Aluminium or m a g n e s i u m hydroxide; calcium carbonate	Interaction not studied. <i>Expected:</i> ↓ Ledipasvir ↔ Sofosbuvir ↔ GS-331007	It is recommended to separate antacid and Ledipasvir/Sofosbuvir 90 mg/400 mg film coated tablets administration by 4 hours.
	(Increase in gastric pH)	
H ₂ -receptor antagonists	;	H ₂ -receptor antagonists may be administered simultaneously with or
Famotidine (40 mg single dose)/ Ledipasvir (90 mg s i n g l e d o s e) ^c / Sofosbuvir (400 mg single dose) ^{c, d}	Ledipasvir ↓ C_{max} 0.80 (0.69, 0.93) ↔ AUC 0.89 (0.76, 1.06)	staggered from Ledipasvir/Sofosbuvir 90 mg/400 mg film coated tablets at a dose that does not exceed doses comparable to famotidine 40 mg twice daily.
Famotidine dosed simultaneously with Ledipasvir/Sofosbuvir 90 mg/400 mg film coated tablets ^d Cimetidine ^e Nizatidine ^e Ranitidine ^e	Sofosbuvir $\uparrow C_{max}$ 1.15 (0.88, 1.50) \leftrightarrow AUC 1.11 (1.00, 1.24) GS-331007 $\leftrightarrow C_{max}$ 1.06 (0.97, 1.14) \leftrightarrow AUC 1.06 (1.02, 1.11)	
	(Increase in gastric pH)	

Omeprazole (20 mg once daily)/ Ledipasvir (90 mg single dose) ^c / Sofosbuvir (400 mg single dose) ^c	Ledipasvir ↓ C _{max} 0.89 (0.61, 1.30) ↓ AUC 0.96 (0.66, 1.39)	mg/400 mg film coated tablets. Proton pump inhibitors should not be taken before Ledipasvir/Sofosbuvir 90 mg/400 mg film coated tablets.
Omeprazole dosed simultaneously with Ledipasvir/Sofosbuvir 90 mg/400 mg film coated tablets Lansoprazole ^e Rabeprazole ^e Pantoprazole ^e Esomeprazole ^e	Sofosbuvir ↔ C_{max} 1.12 (0.88, 1.42) ↔ AUC 1.00 (0.80, 1.25) GS-331007 ↔ C_{max} 1.14 (1.01, 1.29) ↔ AUC 1.03 (0.96, 1.12)	
	(Increase in gastric pH)	
ANTIARRHYTHMICS		
Amiodarone	Interaction not studied.	Use only if no other alternative is available. Close monitoring is recommended if this medicinal product is administered with Ledipasvir/ Sofosbuvir 90 mg/400 mg film coated tablets (see sections 4.4 and 4.8).

Digoxin	Interaction not studied. Expected: \uparrow Digoxin \leftrightarrow Ledipasvir \leftrightarrow Sofosbuvir \leftrightarrow GS-331007 (Inhibition of P-gp)	Co-administration of Ledipasvir/ Sofosbuvir 90 mg/400 mg film coated tablets with digoxin may increase the concentration of digoxin. Caution is warranted and therapeutic concentration monitoring of digoxin is recommended when co-administered with Ledipasvir/ Sofosbuvir 90 mg/400 mg film coated tablets.
ANTICOAGULANTS		
Dabigatran etexilate	Interaction not studied. <i>Expected:</i> ↑ Dabigatran ↔ Ledipasvir ↔ Sofosbuvir ↔ GS-331007 (Inhibition of P-gp)	Clinical monitoring, looking for signs of bleeding and anaemia, is recommended when dabigatran etexilate is co- administered with Ledipasvir/ Sofosbuvir 90 mg/400 mg film coated tablets. A coagulation test helps to identify patients with an increased bleeding risk due to increased dabigatran exposure.
ANTICONVULSANTS		
Carbamazepine Phenytoin Phenobarbital Oxcarbazepine	Interaction not studied. <i>Expected:</i> ↓ Ledipasvir ↓ Sofosbuvir ↓ GS-331007 (Induction of P-gp)	Co-administration of Ledipasvir/ Sofosbuvir 90 mg/400 mg film coated tablets with carbamazepine, phenytoin, phenobarbital or oxcarbazepine is expected to decrease the concentration of Ledipasvir and Sofosbuvir which may lead to reduced therapeutic effect of Ledipasvir/Sofosbuvir 90 mg/400 mg film coated tablets. Ledipasvir/Sofosbuvir 90 mg/400 mg film coated tablets should not be used with carbamazepine, phenytoin, phenobarbital or oxcarbazepine.
ANTIMYCOBACTERI	ALS	

Rifampicin (600 mg once daily)/ Ledipasvir (90 mg single dose) ^d	Interaction not studied. <i>Expected:</i> Rifampicin $\leftrightarrow C_{max}$ $\leftrightarrow AUC$ $\leftrightarrow C_{min}$ <i>Observed:</i> Ledipasvir $\downarrow C_{max} 0.65 (0.56, 0.76)$ $\downarrow AUC 0.41 (0.36, 0.48)$ (Induction of P-gp)	Ledipasvir/Sofosbuvir 90 mg/400 mg film coated tablets should not be used with rifampicin, a potent P-gp inducer (see section 4.4).
Rifampicin (600 mg once daily)/ Sofosbuvir (400 mg single dose) ^d	Interaction not studied. <i>Expected:</i> Rifampicin $\leftrightarrow C_{max}$ $\leftrightarrow AUC$ $\leftrightarrow C_{min}$ <i>Observed:</i> Sofosbuvir $\downarrow C_{max} 0.23 (0.19, 0.29)$ $\downarrow AUC 0.28 (0.24, 0.32)$ GS-331007 $\leftrightarrow C_{max} 1.23 (1.14, 1.34)$ $\leftrightarrow AUC 0.95 (0.88, 1.03)$ (Induction of P-gp)	
Rifabutin Rifapentine	Interaction not studied. <i>Expected:</i> ↓ Sofosbuvir ↔ GS-331007 (Induction of P-gp)	Co-administration of Ledipasvir/ Sofosbuvir 90 mg/400 mg film coated tablets with rifabutin or rifapentine is expected to decrease the concentration of Ledipasvir and Sofosbuvir, leading to reduced therapeutic effect of Ledipasvir/ Sofosbuvir 90 mg/400 mg film coated tablets. Such coadministration is not recommended.
HCV PRODUCTS		

Simeprevir (150 mg once daily)/ Ledipasvir (30 mg once daily)	Simeprevir $\uparrow C_{max} 2.61$ (2.39, 2.86) $\uparrow AUC 2.69$ (2.44, 2.96) Ledipasvir $\uparrow C_{max} 1.81$ (1.69, 2.94) $\uparrow AUC 1.92$ (1.77, 2.07)	Concentrations of Ledipasvir, Sofosbuvir and simeprevir are increased when simeprevir is coadministered with Ledipasvir/Sofosbuvir 90 mg/400 mg film coated tablets. Co-administration is not recommended.
Simeprevir ^h	Simeprevir ↔ C_{max} 0.96 (0.71, 1.30) ↔ AUC 0.94 (0.67, 1.33) Sofosbuvir ↑ C_{max} 1.91 (1.26, 2.90) ↑ AUC 3.16 (2.25, 4.44) GS-331007 ↓ C_{max} 0.69 (0.52, 0.93) ↔ AUC 1.09 (0.87, 1.37)	
HIV ANTIVIRAL AGEN	ITS: REVERSE TRAN	SCRIPTASE INHIBITORS

E f a v i r e n z / e m t r i c i t a b i n e / tenofovir disoproxil fumarate (600 mg/ 200 mg/ 300 mg/ once daily)/ Ledipasvir (90 mg once daily) ^c / Sofosbuvir (400 mg once daily) ^{c, d}	$\begin{array}{l} \mbox{Efavirenz} \\ \leftrightarrow \ C_{max} \ 0.87 \ (0.79, \\ 0.97) \\ \leftrightarrow \ AUC \ 0.90 \ (0.84, \\ 0.96) \\ \leftrightarrow \ C_{min} \ 0.91 \ (0.83, \\ 0.99) \\ \hline \\ \mbox{Emtricitabine} \\ \leftrightarrow \ C_{max} \ 1.08 \ (0.97, \\ 1.21) \\ \leftrightarrow \ AUC \ 1.05 \ (0.98, \\ 1.11) \\ \leftrightarrow \ C_{min} \ 1.04 \ (0.98, \\ 1.11) \\ \end{array}$	No dose adjustment of Ledipasvir/ Sofosbuvir 90 mg/400 mg film coated tablets or efavirenz/ emtricitabine/ tenofovir disoproxil fumarate is required.
	Tenofovir ↑ C _{max} 1.79 (1.56, 2.04) ↑ AUC 1.98 (1.77, 2.23) ↑ C _{min} 2.63 (2.32, 2.97)	
	Ledipasvir $\downarrow C_{max} 0.66 (0.59, 0.75)$ $\downarrow AUC 0.66 (0.59, 0.75)$ $\downarrow C_{min} 0.66 (0.57, 0.76)$	
	Sofosbuvir $\leftrightarrow C_{max}$ 1.03 (0.87, 1.23) \leftrightarrow AUC 0.94 (0.81, 1.10)	
	$\begin{array}{l} \text{GS-331007} \\ \leftrightarrow \ \text{C}_{\text{max}} \ 0.86 \ (0.76, \\ 0.96) \\ \leftrightarrow \ \text{AUC} \ 0.90 \ (0.83, \\ 0.97) \\ \leftrightarrow \ \text{C}_{\text{min}} \ 1.07 \ (1.02, \\ 1.13) \end{array}$	

Emtricitabine/ rilpivirine/tenofovir	Emtricitabine \leftrightarrow C _{max} 1.02 (0.98,	No dose adjustment of Ledipasvir/ Sofosbuvir 90 mg/400 mg film coated
disoproxil fumarate (200 mg/ 25 mg/ 300	1.06) ↔ AUC 1.05 (1.02,	tablets or emtricitabine/ rilpivirine/ tenofovir disoproxil fumarate is required.
mg once daily)/ Ledipasvir	$\begin{array}{rl} 1.08) \\ \leftrightarrow & C_{min} & 1.06 & (0.97, \end{array}$	
(90 mg once daily) ^c / Sofosbuvir (400 mg	1.15)	
once	Rilpivirine $\leftrightarrow C_{max} 0.97 (0.88,$	
daily) ^{c, d}	1.07) ↔ AUC 1.02 (0.94,	
	1.11)	
	↔ C_{min} 1.12 (1.03, 1.21)	
	Tenofovir	
	$\leftrightarrow C_{max} 1.32 (1.25, 1.39)$	
	↑ AUC 1.40 (1.31, 1.50)	
	↑ C _{min} 1.91 (1.74, 2.10)	
	Ledipasvir	
	↔ C _{max} 1.01 (0.95, 1.07)	
	↔ AUC 1.08 (1.02, 1.15)	
	$ \leftrightarrow \ C_{min} \ 1.16 \ (1.08, \\ 1.25) $	
	Sofosbuvir	
	↔ C _{max} 1.05 (0.93, 1.20)	
	↔ AUC 1.10 (1.01, 1.21)	
	GS-331007	
	$\leftrightarrow C_{max} 1.06 (1.01, 1.11)$	
	↔ AUC 1.15 (1.11, 1.19)	
	↔ Ć _{min} 1.18 (1.13, 1.24)	

↔ AUC 1.05 (1.01,	A b a c a v i r / lamivudine (600 mg/ 300 mg once daily)/ Ledipasvir (90 mg once daily)°/ Sofosbuvir (400 mg once daily)°.d	Abacavir ↔ C_{max} 0.92 (0.87, 0.97) ↔ AUC 0.90 (0.85, 0.94) Lamivudine ↔ C_{max} 0.93 (0.87, 1.00) ↔ AUC 0.94 (0.90, 0.98) ↔ C_min 1.12 (1.05, 1.20) Ledipasvir ↔ C_{max} 1.10 (1.01, 1.19) ↔ AUC 1.18 (1.10, 1.28) ↔ C_min 1.26 (1.17, 1.36) Sofosbuvir ↔ C_{max} 1.08 (0.85, 1.35) ↔ AUC 1.21 (1.09, 1.35) GS-331007 ↔ C_{max} 1.00 (0.94, 1.07)	No dose adjustment of Ledipasvir/ Sofosbuvir 90 mg/400 mg film coated tablets or abacavir/ lamivudine is required.
$\begin{array}{c} 1.09) \\ \leftrightarrow C_{min} \ 1.08 \ (1.01, \\ 1.14) \end{array}$		1.07) ↔ AUC 1.05 (1.01, 1.09) ↔ C_{min} 1.08 (1.01,	

Atazanavir boosted with ritonavir (300 mg/ 100 mg once daily)/ Ledipasvir (90 mg once daily) ^c / Sofosbuvir (400 mg once daily) ^{c, d}	Atazanavir $\leftrightarrow C_{max}$ 1.07 (1.00, 1.15) \leftrightarrow AUC 1.33 (1.25, 1.42) ↑ C _{min} 1.75 (1.58, 1.93)	No dose adjustment of Ledipasvir/ Sofosbuvir 90 mg/400 mg film coated tablets or atazanavir (ritonavir boosted) is required. For the combination of tenofovir/ emtricitabine + atazanavir/ritonavir, please see below.
	Ledipasvir	
	↑ C _{max} 1.98 (1.78, 2.20)	
	↑ AUC 2.13 (1.89, 2.40)	
	↑ C _{min} 2.36 (2.08, 2.67)	
	Sofosbuvir	
	$\leftrightarrow C_{max} 0.96 (0.88, 1.05)$	
	↔ ÁUC 1.08 (1.02, 1.15)	
	GS-331007 ↔ C_{max} 1.13 (1.08, 1.19) ↔ AUC 1.23 (1.18, 1.29) ← 1.28 (1.21)	
	↔ C _{min} 1.28 (1.21, 1.36)	

Darunavir boosted with ritonavir (800 mg/ 100 mg once daily)/ Ledipasvir (90 mg once daily) ^d	Darunavir ↔ C_{max} 1.02 (0.88, 1.19) ↔ AUC 0.96 (0.84, 1.11) ↔ C_{min} 0.97 (0.86, 1.10) Ledipasvir ↑ C_{max} 1.45 (1.34, 1.56) ↑ AUC 1.39 (1.28, 1.49) ↑ C_{min} 1.39 (1.29, 1.51)	No dose adjustment of Ledipasvir/ Sofosbuvir 90 mg/400 mg film coated tablets or darunavir (ritonavir boosted) is required. For the combination of tenofovir/ emtricitabine + darunavir/ritonavir, please see below.
Darunavir boosted with ritonavir (800 mg/ 100 mg once daily)/ Sofosbuvir (400 mg once daily)	Darunavir ↔ C_{max} 0.97 (0.94, 1.01) ↔ AUC 0.97 (0.94, 1.00) ↔ C_{min} 0.86 (0.78, 0.96) Sofosbuvir ↑ C_{max} 1.45 (1.10, 1.92) ↑ AUC 1.34 (1.12, 1.59) GS-331007 ↔ C_{max} 0.97 (0.90, 1.05) ↔ AUC 1.24 (1.18, 1.30)	

<pre>with ritonavir (800 mg/ 100 mg once daily) + e m t r i c i t a b i n e / tenofovir disoproxil fumarate (200 mg/ 300 mg once daily)/ Ledipasvir (90 mg once daily)^{c/} Sofosbuvir (400 mg once daily)^{c, d} D o s e d simultaneously^f</pre>	$\begin{array}{l} \leftrightarrow \ {\rm Cmax} \ 1.01 \ (0.96, \\ 1.06) \\ \leftrightarrow \ {\rm AUC} \ 1.04 \ (0.99, \\ 1.08) \\ \leftrightarrow \ {\rm Cmin} \ 1.08 \ (0.98, \\ 1.20) \\ \hline \\ \ {\rm Ritonavir} \\ \leftrightarrow \ {\rm Cmax} \ 1.17 \ (1.01, \\ 1.35) \\ \leftrightarrow \ {\rm AUC} \ 1.25 \ (1.15, \\ 1.36) \\ \uparrow \ {\rm Cmin} \ 1.48 \ (1.34, \\ 1.63) \\ \hline \\ \ {\rm Emtricitabine} \\ \leftrightarrow \ {\rm Cmax} \ 1.02 \ (0.96, \\ 1.08) \\ \leftrightarrow \ {\rm AUC} \ 1.04 \ (1.00, \\ 1.08) \\ \leftrightarrow \ {\rm AUC} \ 1.04 \ (1.00, \\ 1.08) \\ \leftrightarrow \ {\rm Cmin} \ 1.03 \ (0.97, \\ 1.10) \\ \hline \\ \ \ {\rm Tenofovir} \\ \uparrow \ {\rm Cmax} \ 1.64 \ (1.54, \\ 1.74) \\ \uparrow \ {\rm AUC} \ 1.50 \ (1.42, \\ 1.59) \\ \uparrow \ {\rm Cmin} \ 1.59 \ (1.49, \\ 1.70) \\ \hline \\ \ \ {\rm Ledipasvir} \\ \leftrightarrow \ {\rm Cmax} \ 1.11 \ (0.99, \\ 1.24) \\ \leftrightarrow \ {\rm AUC} \ 1.12 \ (1.00, \\ 1.25) \\ \leftrightarrow \ {\rm Cmin} \ 1.17 \ (1.04, \\ 1.31) \\ \hline \\ \ \ {\rm Sofosbuvir} \\ \downarrow \ {\rm Cmax} \ 0.63 \ (0.52, \\ 0.75) \\ \downarrow \ {\rm AUC} \ 0.73 \ (0.65, \\ 0.82) \\ \hline \\ \ \ {\rm GS-331007} \\ \leftrightarrow \ {\rm AUC} \ 1.20 \ (1.16, \\ \end{array}$	used in conjunction with tenofovir disoproxil fumarate, Ledipasvir/Sofosbuvir 90 mg/400 mg film coated tablets increased the concentration of tenofovir. The safety of tenofovir disoproxil fumarate in the setting of Ledipasvir/ Sofosbuvir 90 mg/400 mg film coated tablets and a pharmacokinetic enhancer (e.g. ritonavir or cobicistat) has not been established. The combination should be used with caution with frequent renal monitoring, if other alternatives are not available (see section 4.4).
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Lopinavir boosted with ritonavir + emtricitabine/ tenofovir disoproxil fumarate	Interaction not studied. <i>Expected:</i> ↑ Lopinavir ↑ Ritonavir	When given with lopinavir/ritonavir used in conjunction with tenofovir disoproxil fumarate, Ledipasvir/Sofosbuvir 90 mg/ 400 mg film coated tablets is expected to increase the concentration of tenofovir.
	 ↔ Emtricitabine ↑ Tenofovir ↑ Ledipasvir ↔ Sofosbuvir ↔ GS-331007 	The safety of tenofovir disoproxil fumarate in the setting of Ledipasvir/ Sofosbuvir 90 mg/400 mg film coated tablets and a pharmacokinetic enhancer (e.g. ritonavir or cobicistat) has not been established.
		The combination should be used with caution with frequent renal monitoring, if other alternatives are not available (see section 4.4).
Tipranavir boosted with ritonavir	Interaction not studied. <i>Expected:</i> ↓ Ledipasvir ↓ Sofosbuvir ↔ GS-331007 (Induction of P-gp)	Co-administration of Ledipasvir/ Sofosbuvir 90 mg/400 mg film coated tablets with tipranavir (ritonavir boosted) is expected to decrease the concentration of Ledipasvir, leading to reduced therapeutic effect of Ledipasvir/ Sofosbuvir 90 mg/400 mg film coated tablets. Co-administration is not recommended.
HIV ANTIVIRAL AGENTS: INTEGRASE INHIBITORS		

Raltegravir (400 mg twice daily)/ Ledipasvir (90 mg once daily) ^d		No dose adjustment of Ledipasvir/ Sofosbuvir 90 mg/400 mg film coated tablets or raltegravir is required.
	$\begin{array}{l} \mbox{Ledipasvir} \\ \leftrightarrow \ C_{max} \ 0.92 \ (0.85, \\ 1.00) \\ \leftrightarrow \ AUC \ 0.91 \ (0.84, \\ 1.00) \\ \leftrightarrow \ C_{min} \ 0.89 \ (0.81, \\ 0.98) \end{array}$	

Raltegravir (400 mg twice daily)/ Sofosbuvir (400 mg once daily) ^d	Raltegravir ↓ C_{max} 0.57 (0.44, 0.75) ↓ AUC 0.73 (0.59, 0.91) ↔ C_{min} 0.95 (0.81, 1.12)
	Sofosbuvir \leftrightarrow C _{max} 0.87 (0.71, 1.08) \leftrightarrow AUC 0.95 (0.82, 1.09)
	$\begin{array}{l} \text{GS-331007} \\ \leftrightarrow \ \text{C}_{\text{max}} \ 1.09 \ (0.99, \\ 1.19) \\ \leftrightarrow \ \text{AUC} \ 1.02 \ (0.97, \\ 1.08) \end{array}$

	i	
Elvitegravir/ cobicistat/ emtricitabine/ tenofovir disoproxil fumarate (150 mg/ 150 mg/ 200 mg/ 300 mg once daily)/ Ledipasvir (90 mg once daily)°/ Sofosbuvir (400 mg once daily)°	Interaction not studied. <i>Expected:</i> \leftrightarrow Emtricitabine ↑ Tenofovir <i>Observed:</i> Elvitegravir \leftrightarrow C _{max} 0.88 (0.82, 0.95) \leftrightarrow AUC 1.02 (0.95, 1.09) ↑ C _{min} 1.36 (1.23, 1.49) Cobicistat \leftrightarrow C _{max} 1.25 (1.18, 1.32) ↑ AUC 1.59 (1.49, 1.70) ↑ C _{min} 4.25 (3.47, 5.22) Ledipasvir ↑ C _{max} 1.63 (1.51, 1.75) ↑ AUC 1.78 (1.64, 1.94) ↑ C _{min} 1.91 (1.76, 2.08) Sofosbuvir ↑ C _{max} 1.33 (1.14, 1.56) ↑ AUC 1.36 (1.21, 1.52) GS-331007 ↑ C _{max} 1.33 (1.22, 1.44) ↑ AUC 1.44 (1.41, 1.48) ↑ C _{min} 1.53 (1.47, 1.59)	When given with elvitegravir/ cobicistat/ emtricitabine/ tenofovir disoproxil fumarate, Ledipasvir/Sofosbuvir 90 mg/400 mg film coated tablets is expected to increase the concentration of tenofovir. The safety of tenofovir disoproxil fumarate in the setting of Ledipasvir/ Sofosbuvir 90 mg/400 mg film coated tablets and a pharmacokinetic enhancer (e.g. ritonavir or cobicistat) has not been established. The combination should be used with caution with frequent renal monitoring, if other alternatives are not available (see section 4.4).
Dolutegravir	Interaction not	No dose adjustment required.
	studied. <i>Expected:</i> ↔ Dolutegravir ↔ Ledipasvir ↔ Sofosbuvir ↔ GS-331007	
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HERBAL SUPPLEMENTS

St. John's wort	Interaction not studied. <i>Expected:</i> ↓ Ledipasvir ↓ Sofosbuvir ↓ GS-331007	Co-administration of Ledipasvir/ Sofosbuvir 90 mg/400 mg film coated tablets with St. John's wort is contraindicated (see section 4.3).
	(Induction of P-gp)	
HMG-CoA REDUCTA	SE INHIBITORS	1
Rosuvastatin ^g	↑ Rosuvastatin (Inhibition of drug transporters OATP and BCRP)	Co-administration of Ledipasvir/ Sofosbuvir 90 mg/400 mg film coated tablets with rosuvastatin may significantly increase the concentration of rosuvastatin (several fold-increase in AUC) which is associated with increased risk of myopathy, including rhabdomyolysis. Co-administration of Ledipasvir/Sofosbuvir 90 mg/400 mg film coated tablets with rosuvastatin is contraindicated (see section 4.3).
Pravastatin ^g	↑ Pravastatin	Co-administration of Ledipasvir/ Sofosbuvir 90 mg/400 mg film coated tablets with pravastatin may significantly increase the concentration of pravastatin which is associated with increased risk of myopathy. Clinical and biochemical control is recommended in these patients and a dose adjustment may be needed (see section 4.4).
Other statins	<i>Expected</i> : ↑ Statins	Interactions cannot be excluded with other HMGCoA reductase inhibitors. When co-administered with Ledipasvir/ Sofosbuvir 90 mg/400 mg film coated tablets, a reduced dose of statins should be considered and careful monitoring for statin adverse reactions should be undertaken (see section 4.4).
NARCOTIC ANALGESICS		
Methadone	Interaction not studied. <i>Expected:</i> ↔ Ledipasvir	No dose adjustment of Ledipasvir/ Sofosbuvir 90 mg/400 mg film coated tablets or methadone is required.

Methadone (M e t h a d o n e maintenance therapy [30 to 1 3 0 mg/daily])/ Sofosbuvir (400 mg once daily) ^d	R-methadone ↔ C_{max} 0.99 (0.85, 1.16) ↔ AUC 1.01 (0.85, 1.21) ↔ C_{min} 0.94 (0.77, 1.14) S-methadone ↔ C_{max} 0.95 (0.79, 1.13) ↔ AUC 0.95 (0.77, 1.17) ↔ C_{min} 0.95 (0.74, 1.22) Sofosbuvir ↓ C_{max} 0.95 (0.68, 1.33) ↑ AUC 1.30 (1.00, 1.69) GS-331007 ↓ C_{max} 0.73 (0.65, 0.83) ↔ AUC 1.04 (0.89, 1.22)	
IMMUNOSUPPRESSA	ANTS	
Ciclosporing	Interaction not studied. <i>Expected:</i> ↑ Ledipasvir ↔ Ciclosporin	No dose adjustment of Ledipasvir/ Sofosbuvir 90 mg/400 mg film coated tablets or ciclosporin is required.
Ciclosporin (600 mg single dose)/ Sofosbuvir (400 mg single dose) ^h	Ciclosporin \leftrightarrow C _{max} 1.06 (0.94, 1.18) \leftrightarrow AUC 0.98 (0.85, 1.14) Sofosbuvir ↑ C _{max} 2.54 (1.87, 3.45) ↑ AUC 4.53 (3.26, 6.30) GS-331007 ↓ C _{max} 0.60 (0.53, 0.69) \leftrightarrow AUC 1.04 (0.90, 1.20)	

Tacrolimus	Interaction not studied. <i>Expected:</i> ↔ Ledipasvir	No dose adjustment of Ledipasvir/ Sofosbuvir 90 mg/400 mg film coated tablets or tacrolimus is required.
Tacrolimus (5 mg single dose)/ Sofosbuvir (400 mg single dose) ^h	Tacrolimus ↓ C _{max} 0.73 (0.59, 0.90) ↑ AUC 1.09 (0.84, 1.40)	
	Sofosbuvir ↓ C _{max} 0.97 (0.65, 1.43) ↑ AUC 1.13 (0.81, 1.57)	
	$\begin{array}{l} \text{GS-331007} \\ \leftrightarrow \ \text{C}_{\text{max}} \ 0.97 \ (0.83, \\ 1.14) \\ \leftrightarrow \ \text{AUC} \ 1.00 \ (0.87, \\ 1.13) \end{array}$	
ORAL CONTRACEPT	IVES	
Norgestimate/ ethinyl e s t r a d i o l (norgestimate 0.180 mg/ 0.215 mg/ 0.25 mg/ ethinyl estradiol 0.025 mg)/ Ledipasvir (90 mg once daily) ^d	Norelgestromin ↔ C_{max} 1.02 (0.89, 1.16) ↔ AUC 1.03 (0.90, \leftrightarrow AUC 1.03 (0.91, 1.31) \leftrightarrow C_{max} 1.09 (0.91, 1.31) Norgestrel ↔ C_{max} 1.03 (0.87, \leftrightarrow AUC 0.99 (0.82, 1.20) ↔ C_{min} 1.00 (0.81,	No dose adjustment of oral contraceptives is required.
	1.23) Ethinyl estradiol	
	↑ C_{max} 1.40 (1.18, 1.66) ↔ AUC 1.20 (1.04,	
	1.39) ↔ C_{min} 0.98 (0.79, 1.22)	

Norgestimate/ ethinyl e s t r a d i o l (norgestimate 0.180 mg/ 0.215 mg/ 0.25 mg/ ethinyl estradiol 0.025 mg)/ Sofosbuvir (400 mg once daily) ^d	Norelgestromin \leftrightarrow C _{max} 1.07 (0.94, 1.22) \leftrightarrow AUC 1.06 (0.92, 1.21) \leftrightarrow C _{min} 1.07 (0.89, 1.28) Norgestrel \leftrightarrow C _{max} 1.18 (0.99, 1.41) \uparrow AUC 1.19 (0.98, 1.45) \uparrow C _{min} 1.23 (1.00, 1.51) Ethinyl estradiol \leftrightarrow C _{max} 1.15 (0.97, 1.36) \leftrightarrow AUC 1.09 (0.94, 1.26)
	$\leftrightarrow C_{\min} 0.99 (0.80, \\ 1.23)$

a. Mean ratio (90% CI) of co-administered drug pharmacokinetics of study medicinal products alone or in combination. No effect = 1.00.

- b. All interaction studies conducted in healthy volunteers.
- c. Administered as Ledipasvir/Sofosbuvir 90 mg/400 mg film coated tablets.
- d. Lack of pharmacokinetics interaction bounds 70-143%.
- e. These are drugs within class where similar interactions could be predicted.
- f. Staggered administration (12 hours apart) of Atazanavir/ritonavir + Emtricitabine/ tenofovir Disoproxil Fumarate or Darunavir/ritonavir + Emtricitabine/tenofovir Disoproxil Fumarate and Ledipasvir/Sofosbuvir 90 mg/400 mg film coated tablets provided similar results.
- g. This study was conducted in the presence of another two direct-acting antiviral agents.
- h. Bioequivalence/Equivalence boundary 80-125%.

4.6 Fertility, pregnancy and lactation

Women of childbearing potential / contraception in males and females

When Ledipasvir/Sofosbuvir 90 mg/400 mg film coated tablets is used in combination with ribavirin, extreme care must be taken to avoid pregnancy in female patients and in female partners of male patients. Significant teratogenic and/or embryocidal effects have been demonstrated in all animal species exposed to ribavirin. Women of childbearing potential or their male partners must use an effective form of contraception during treatment and for a period of time after the treatment has concluded as recommended in the Summary of Product Characteristics for ribavirin. Refer to the Summary of Product Characteristics for ribavirin.

Pregnancy

There are no or limited amount of data (less than 300 pregnancy outcomes) from the use of Ledipasvir, Sofosbuvir or Ledipasvir/Sofosbuvir 90 mg/400 mg film coated tablets in pregnant women.

Animal studies do not indicate direct harmful effects with respect to reproductive toxicity. No significant effects on foetal development have been observed with Ledipasvir or Sofosbuvir in rats and rabbits. However, it has not been possible to fully estimate exposure margins achieved for Sofosbuvir in the rat relative to the exposure in humans at the recommended clinical dose (see section 5.3).

As a precautionary measure, it is preferable to avoid the use of Ledipasvir/Sofosbuvir 90 mg/400 mg film coated tablets during pregnancy.

Breast-feeding

It is unknown whether Ledipasvir or Sofosbuvir and its metabolites are excreted in human milk.

Available pharmacokinetic data in animals has shown excretion of Ledipasvir and metabolites of Sofosbuvir in milk (see section 5.3).

A risk to the newborns/infants cannot be excluded. Therefore, Ledipasvir/Sofosbuvir 90 mg/400 mg film coated tablets should not be used during breast-feeding.

Fertility

No human data on the effect of Ledipasvir/Sofosbuvir 90 mg/400 mg film coated tablets on fertility are available. Animal studies do not indicate harmful effects of Ledipasvir or Sofosbuvir on fertility.

If ribavirin is co-administered with Ledipasvir/Sofosbuvir 90 mg/400 mg film coated tablets, the contraindications regarding use of ribavirin during pregnancy and breast-feeding apply (see also the Summary of Product Characteristics for ribavirin).

4.7 Effects on ability to drive and use machines

Ledipasvir/Sofosbuvir 90 mg/400 mg film coated tablets (administered alone or in combination with ribavirin) has no or negligible influence on the ability to drive and use machines. However, patients should be advised that fatigue was more common in patients treated with Ledipasvir/Sofosbuvir compared to placebo.

4.8 Undesirable effects

Summary of the safety profile

The safety assessment of Ledipasvir/Sofosbuvir is based on pooled data from three Phase 3 clinical studies including 215, 539 and 326 patients who received Ledipasvir/Sofosbuvir for 8, 12 and 24 weeks, respectively; and 216, 328 and 328 patients who received Ledipasvir/Sofosbuvir + ribavirin combination therapy for 8, 12 and 24 weeks, respectively. These studies did not include any control group not receiving Ledipasvir/Sofosbuvir. Further data include a double-blind comparison of the safety of Ledipasvir/Sofosbuvir (12 weeks) and placebo in 155 cirrhotic patients.

The proportion of patients who permanently discontinued treatment due to adverse events was 0%, < 1% and 1% for patients receiving Ledipasvir/Sofosbuvir for 8, 12 and

24 weeks, respectively; and < 1%, 0%, and 2% for patients receiving Ledipasvir/ Sofosbuvir + ribavirin combination therapy for 8, 12 and 24 weeks, respectively.

In clinical studies, fatigue and headache were more common in patients treated with Ledipasvir/ Sofosbuvir compared to placebo. When Ledipasvir/Sofosbuvir was studied with ribavirin, the most frequent adverse drug reactions to Ledipasvir/Sofosbuvir + ribavirin combination therapy were consistent with the known safety profile of ribavirin, without increasing the frequency or severity of the expected adverse drug reactions.

The following adverse drug reactions have been identified with Ledipasvir/Sofosbuvir 90 mg/400 mg film coated tablets (Table 4). The adverse reactions are listed below by body 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 < 1/100), rare (\geq 1/10,000 to < 1/10,000).

Table 4: Adverse drug reactions identified with Ledipasvir/Sofosbuvir 90 mg/400 mg film coated tablets

Frequency	Ledipasvir/Sofosbuvir 90 mg/400 mg film coated tablets
Nervous system disorders:	
Very common	headache
General disorders:	
Very common	fatigue

Paediatric population

The safety and efficacy of Ledipasvir/Sofosbuvir 90 mg/400 mg film coated tablets in children and adolescents aged less than 18 years have not yet been established. No data are available.

Description of selected adverse reactions

Cardiac arrhythmias

Cases of severe bradycardia and heart block have been observed when Ledipasvir/ Sofosbuvir 90 mg/400 mg film coated tablets is used with concomitant amiodarone and/or other drugs that lower heart rate (see sections 4.4 and 4.5).

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 TMDA ADR reporting tool; website: <u>https://imis.tmda.go.tz/arrt</u> or search for TMDA Adverse Reactions Reporting Tool in the Google Play Store";

4.9 Overdose

The highest documented doses of Ledipasvir and Sofosbuvir were 120 mg twice daily for 10 days and a single dose of 1,200 mg, respectively. In these healthy volunteer studies, there were no untoward effects observed at these dose levels, and adverse reactions were similar in frequency and severity to those reported in the placebo groups. The effects of higher doses are not known.

No specific antidote is available for overdose with Ledipasvir/Sofosbuvir 90 mg/400 mg film coated tablets. If overdose occurs the patient must be monitored for evidence of toxicity. Treatment of overdose with Ledipasvir/Sofosbuvir 90 mg/400 mg film coated tablets consists of general supportive measures including monitoring of vital signs as well as observation of the clinical status of the patient. Haemodialysis is unlikely to result in significant removal of Ledipasvir as Ledipasvir is highly bound to plasma protein. Haemodialysis can efficiently remove the predominant circulating metabolite of Sofosbuvir, GS-331007, with an extraction ratio of 53%.

5. PHARMACOLOGICAL PROPERTIES

5.1 Pharmacodynamic properties

Pharmacotherapeutic group: Direct-acting antiviral; ATC code ATC code: J05AX65 Pharmacological classification: 7.13 Antivirals

Mechanism of action

Ledipasvir is a HCV inhibitor targeting the HCV NS5A protein, which is essential for both RNA replication and the assembly of HCV virions. Biochemical confirmation of NS5A inhibition by Ledipasvir is not currently possible as NS5A has no enzymatic function. *In vitro* resistance selection and crossresistance studies indicate Ledipasvir targets NS5A as its mode of action.

Sofosbuvir is a pan-genotypic inhibitor of the HCV NS5B RNA-dependent RNA polymerase, which is essential for viral replication. Sofosbuvir is a nucleotide prodrug that undergoes intracellular metabolism to form the pharmacologically active uridine analogue triphosphate (GS-461203), which can be incorporated into HCV RNA by the NS5B polymerase and acts as a chain terminator. GS-461203 (the active metabolite of Sofosbuvir) is neither an inhibitor of human DNA and RNA polymerases nor an inhibitor of mitochondrial RNA polymerase.

Antiviral activity

The EC₅₀ values of Ledipasvir and Sofosbuvir against full-length or chimeric replicons encoding NS5A and NS5B sequences from clinical isolates are detailed in Table 5. The presence of 40% human serum had no effect on the anti-HCV activity of Sofosbuvir but reduced the anti-HCV activity of Ledipasvir by 12fold against genotype 1a HCV replicons.

Genotype	Ledipasvir activi	ity (EC₅₀, nM)	Sofosbuvir activity (EC ₅₀ , nM)				
replicons	S t a b l e replicons	N S 5 A transient replicons Median (range)ª	S t a b l e replicons	N S 5 B transient replicons Median (range)ª			
Genotype 1a	0.031	0.018 (0.009- 0.085)	40	62 (29-128)			
Genotype 1b	0.004	0.006 (0.004- 0.007)	110	102 (45-170)			
Genotype 2a	21-249	-	50	29 (14-81)			
Genotype 2b	16-530 [⊳]	-	15 ^b	-			
Genotype 3a	168	-	50	81 (24-181)			
Genotype 4a	0.39	-	40	-			
Genotype 4d	0.60	-	-	-			
Genotype 5a	0.15 ^b	-	15 ^b	-			
Genotype 6a	1.1 ^b	-	14 ^b	-			
Genotype 6e	264 ^b	-	-	-			

Table 5: Activity of Ledipasvir and Sofosbuvir against chimeric replicons

a. Transient replicons carrying NS5A or NS5B from patient isolates.

b. The chimeric replicons carrying NS5A genes from genotype 2b, 5a, 6a and 6e were used for testing Ledipasvir while the chimeric replicons carrying NS5B genes from genotype 2b, 5a or 6a were used for testing Sofosbuvir.

<u>Resistance</u>

In cell culture

HCV replicons with reduced susceptibility to Ledipasvir have been selected in cell culture for genotype 1a and 1b. Reduced susceptibility to Ledipasvir was associated with the primary NS5A substitution Y93H in both genotype 1a and 1b. Additionally a Q30E substitution developed in genotype 1a replicons. Sitedirected mutagenesis of NS5A RAVs showed that substitutions conferring a fold-change > 100 and \leq 1,000 in Ledipasvir susceptibility are Q30H/R, L31I/M/V, P32L and Y93T in genotype 1a and P58D and Y93S in genotype 1b; and substitutions conferring a fold-change > 1,000 are M28A/G, Q30E/G/ K, H58D, Y93C/H/N/S in genotype 1a and A92K and Y93H in genotype 1b.

HCV replicons with reduced susceptibility to Sofosbuvir have been selected in cell culture for multiple genotypes including 1b, 2a, 2b, 3a, 4a, 5a and 6a. Reduced susceptibility to Sofosbuvir was associated with the primary NS5B substitution S282T in all replicon genotypes examined. Site-directed mutagenesis of the S282T substitution in replicons of

8 genotypes conferred 2- to 18-fold reduced susceptibility to Sofosbuvir and reduced the viral replication capacity by 89% to 99% compared to the corresponding wildtype.

In clinical studies

In a pooled analysis of patients who received Ledipasvir/Sofosbuvir in Phase 3 studies, 37 patients (29 with genotype 1a and 8 with genotype 1b) qualified for resistance analysis due to virologic failure or early study drug discontinuation and having HCV RNA > 1,000 IU/mL. Post-baseline NS5A and NS5B deep sequencing data (assay cut off of 1%) were available for 37/37 and 36/37 patients, respectively.

 $\mathsf{NS5A}$ resistance-associated variants (RAVs) were observed in post-baseline isolates from

29/37 patients (22/29 genotype 1a and 7/8 genotype 1b) not achieving sustained virologic response (SVR). Of the 29 genotype 1a patients who qualified for resistance testing, 22/29 (76%) patients harboured one or more NS5A RAVs at positions K24, M28, Q30, L31, S38 and Y93 at failure, while the remaining 7/29 patients had no NS5A RAVs detected at failure. The most common variants were Q30R, Y93H and L31M. Of the 8 genotype 1b patients who qualified for resistance testing, 7/8 (88%) harboured one or more NS5A RAVs at positions L31 and Y93 at failure, while 1/8 patients had no NS5A RAVs at failure. The most common variant was Y93H. Among the 8 patients who had no NS5A RAVs at failure, 7 patients received 8 weeks of treatment (n = 3 with Ledipasvir/Sofosbuvir; n = 4 with Ledipasvir/Sofosbuvir + ribavirin) and 1 patient received Ledipasvir/Sofosbuvir for 12 weeks. In phenotypic analyses, post-baseline isolates from patients who harboured NS5A RAVs at failure showed 20- to at least a 243-fold (the highest dose tested) reduced susceptibility to Ledipasvir. Site-directed mutagenesis of the Y93H substitution in both genotype

1a and 1b as well as the Q30R and L31M substitution in genotype 1a conferred high levels of reduced susceptibility to Ledipasvir (fold-

change in EC_{50} ranging from 544-fold to 1,677-fold).

The Sofosbuvir resistance-associated substitution S282T in NS5B was not detected in any virologic failure isolate from the Phase 3 studies. However, the NS5B S282T substitution in combination with NS5A substitutions L31M, Y93H and Q30L were detected in one patient at failure following 8 weeks of treatment with Ledipasvir/Sofosbuvir from a Phase 2 study (LONESTAR). This patient was subsequently retreated with Ledipasvir/Sofosbuvir + ribavirin for 24 weeks and achieved SVR following retreatment.

Effect of baseline HCV resistance-associated variants on treatment outcome Analyses were conducted to explore the association between pre-existing baseline

NS5A RAVs and treatment outcome. In the pooled analysis of the Phase 3 studies, 16% of patients had baseline NS5A RAVs identified by population or deep sequencing irrespective of subtype. Baseline NS5A RAVs were overrepresented in patients who experienced relapse in the Phase 3 studies (see "Clinical efficacy and safety").

Following 12 weeks of treatment with Ledipasvir/Sofosbuvir (without ribavirin) in treatmentexperienced patients (arm 1 of ION-2 study) 4/4 patients with baseline NS5A RAVs conferring a Ledipasvir fold-change of \leq 100 achieved SVR. For the same treatment arm, patients with baseline NS5A RAVs conferring a foldchange of > 100, relapse occurred in 4/13 (31%), as compared to 3/95 (3%) in those without any baseline RAVs or RAVs conferring a fold-change of \leq 100.

The group of NS5A RAVs that conferred > 100-fold shift and was observed in patients were the following substitutions in genotype 1a (M28A, Q30H/R/E, L31M/V/I, H58D, Y93H/N/C) or in genotype 1b (Y93H). The proportion of such baseline NS5A RAVs seen

with deep sequencing varied from very low (cut off for assay = 1%) to high (main part of the plasma population).

The Sofosbuvir resistance-associated substitution S282T was not detected in the baseline NS5B sequence of any patient in Phase 3 studies by population or deep sequencing. SVR was achieved in all 24 patients (n = 20 with L159F+C316N; n = 1 with L159F; and n = 3 with N142T) who had baseline variants associated with resistance to NS5B nucleoside inhibitors.

Cross-resistance

Ledipasvir was fully active against the Sofosbuvir resistance-associated substitution S282T in NS5B while all Ledipasvir resistance-associated substitutions in NS5A were fully susceptible to Sofosbuvir. Both Sofosbuvir and Ledipasvir were fully active against substitutions associated with resistance to other classes of direct-acting antivirals with different mechanisms of actions, such as NS5B non-nucleoside inhibitors and NS3 protease inhibitors. NS5A substitutions conferring resistance to Ledipasvir may reduce the antiviral activity of other NS5A inhibitors.

Clinical efficacy and safety

The efficacy of Ledipasvir/Sofosbuvir 90 mg/400 mg film coated tablets (Ledipasvir [LDV]/ Sofosbuvir [SOF]) was evaluated in three open-label

Phase 3 studies with data available for a total of 1,950 patients with genotype 1 CHC. The three Phase 3 studies included one study conducted in non-cirrhotic treatment-naïve patients (ION-3); one study in cirrhotic and non-cirrhotic treatment-naïve patients (ION-1); and one study in cirrhotic and non-cirrhotic patients who failed prior therapy with an interferon-based regimen, including regimens containing an HCV protease inhibitor (ION-2). Patients in these studies had compensated liver disease. All three Phase 3 studies evaluated the efficacy of Ledipasvir/Sofosbuvir with or without ribavirin.

Treatment duration was fixed in each study. Serum HCV RNA values were measured during the clinical studies using the COBAS TaqMan HCV test (version 2.0), for use with the High Pure System. The assay had a lower limit of quantification (LLOQ) of 25 IU/mL. SVR was the primary endpoint to determine the HCV cure rate which was defined as HCV RNA less than LLOQ at 12 weeks after the cessation of treatment.

Treatment-naïve adults without cirrhosis – ION-3 (study 0108) – Genotype 1

ION-3 evaluated 8 weeks of treatment with Ledipasvir/Sofosbuvir with or without ribavirin and 12 weeks of treatment with Ledipasvir/Sofosbuvir in treatment-naïve noncirrhotic patients with genotype 1 CHC. Patients were randomised in a 1:1:1 ratio to one of the three treatment groups and stratified by HCV genotype (1a *versus* 1b).

Patient disposition	LDV/SOF 8 weeks (n = 215)	L D V / SOF+RBV 8 weeks	LDV/SOF 12 weeks (n = 216)	TOTAL (n = 647)
		(n = 216)		
Age (years): median (range)	53 (22-75)	51 (21-71)	53 (20-71)	52 (20-75)
Male gender	60% (130)	54% (117)	59% (128)	58% (375)
Race: Black/ African American	21% (45)	17% (36)	19% (42)	19% (123)
White	76% (164)	81% (176)	77% (167)	78% (507)
Genotype 1a	80% (171)	80% (172)	80% (172)	8 0 % (515) ^a
IL28CC genotype	26% (56)	28% (60)	26% (56)	27% (172)
FibroTest-Determined Me	etavir score ^b	·		
F0-F1	33% (72)	38% (81)	33% (72)	35% (225)
F2	30% (65)	28% (61)	30% (65)	30% (191)
F3-F4	36% (77)	33% (71)	37% (79)	35% (227)
Not interpretable	< 1% (1)	1% (3)	0% (0)	< 1% (4)

Table 6: Demographics and baseline characteristics in study ION-3

a. One patient in the LDV/SOF 8-week treatment arm did not have a confirmed genotype 1 subtype.

 b. Non-missing FibroTest results are mapped to Metavir scores according to: 0-0.31 = F0-F1; 0.32-0.58 = F2; 0.59-1.00 = F3F4.

Table 7: Response rates in study ION-3

	LDV/SOF 8 weeks (n = 215)	LDV/SOF+RBV 8 weeks (n = 216)	LDV/SOF 12 weeks (n = 216)
SVR	94% (202/215)	93% (201/216)	96% (208/216)
Outcome for patients without SVR			
On-treatment virologic failure	0/215	0/216	0/216
Relapse ^a	5% (11/215)	4% (9/214)	1% (3/216)
Other ^b	< 1% (2/215)	3% (6/216)	2% (5/216)
Genotype	·	·	·
Genotype 1a	93% (159/171)	92% (159/172)	96% (165/172)
Genotype 1b	98% (42/43)	95% (42/44)	98% (43/44)

a. The denominator for relapse is the number of patients with HCV RNA < LLOQ at their last on-treatment assessment.

b. Other includes patients who did not achieve SVR and did not meet virologic failure criteria (e.g. lost to follow-up).

The 8-week treatment of Ledipasvir/Sofosbuvir without ribavirin was non-inferior to the 8-week treatment of Ledipasvir/Sofosbuvir with ribavirin (treatment difference 0.9%; 95% confidence interval: 3.9% to 5.7%) and the 12-week treatment of Ledipasvir/Sofosbuvir (treatment difference -2.3%; 97.5% confidence interval: -7.2% to 3.6%). Among patients with a baseline HCV RNA < 6 million IU/mL, the SVR was 97% (119/123) with 8-week treatment of Ledipasvir/Sofosbuvir and 96% (126/131) with 12week treatment of Ledipasvir/Sofosbuvir.

Table 8: Relapse rates	by base	eline	characteristics	in	the	ION-3	study,	virological
failure population*								

	LDV/SOF 8 weeks (n = 213)	L D V SOF+RBV 8 weeks (n = 210)	LDV/SOF 12 weeks (n = 211)
Gender			
Male	8% (10/129)	7% (8/114)	2% (3/127)
Female	1% (1/84)	1% (1/96)	0% (0/84)
IL28 genotype			
CC	4% (2/56)	0% (0/57)	0% (0/54)
Non-CC	6% (9/157)	6% (9/153)	2% (3/157)
Baseline HCV RNAª			
HCV RNA < 6 million IU/mL	2% (2/121)	2% (3/136)	2% (2/128)
HCV RNA ≥ 6 million IU/mL	10% (9/92)	8% (6/74)	1% (1/83)

* Patients lost to follow-up or who withdrew consent excluded.

a. HCV RNA values were determined using the Roche TaqMan Assay; a patient's HCV RNA may vary from visit to visit.

Treatment-naïve adults with or without cirrhosis – *ION-1 (study 0102)* – *Genotype 1* ION-1 was a randomised, open-label study that evaluated 12 and 24 weeks of treatment with Ledipasvir/Sofosbuvir with or without ribavirin in 865 treatment-naïve patients with genotype 1 CHC including those with cirrhosis (randomised 1:1:1:1). Randomisation was stratified by the presence or absence of cirrhosis and HCV genotype (1a *versus* 1b).

Patient disposition	LDV/SOF 12 weeks (n = 214)	L D V / S O F + R B V 12 weeks (n = 217)	L D V / SOF 2 4 weeks (n = 217)	L D V / S O F + R B V 24 weeks (n = 217)	TOTAL (n = 865)
Age (years): median (range)	52 (18-75)	5 2 (18-78)	5 3 (22-80)	5 3 (24-77)	52 (18-80)
Male gender	59% (127)	59% (128)	6 4 % (139)	55% (119)	59% (513)
Race: Black/ African American	11% (24)	12% (26)	1 5 % (32)	12% (26)	12% (108)
White	87% (187)	87% (188)	8 2 % (177)	84% (183)	85% (735)
Genotype 1a ^a	68% (145)	68% (148)	6 7 % (146)	66% (143)	67% (582)
IL28CC genotype	26% (55)	35% (76)	2 4 % (52)	34% (73)	30% (256)
FibroTest-Determine score ^b	ed Metavir				
F0-F1	27% (57)	26% (56)	2 9 % (62)	30% (66)	28% (241)
F2	26% (56)	25% (55)	2 2 % (47)	28% (60)	25% (218)
F3-F4	47% (100)	48% (104)	4 9 % (107)	42% (91)	46% (402)
Not interpretable	< 1% (1)	1% (2)	< 1 % (1)	0% (0)	< 1% (4)

Table 9: Demographics and baseline characteristics in study ION-1

a. Two patients in the LDV/SOF 12-week treatment arm, one patient in the LDV/ SOF+RBV 12-week treatment arm, two

patients in the LDV/SOF 24-week treatment arm, and two patients in the LDV/SOF+RBV 24-week treatment arm did not have a confirmed genotype 1 subtype.

 Non-missing FibroTest results are mapped to Metavir scores according to: 0-0.31 = F0-F1; 0.32-0.58 = F2;

0.59-1.00 = F3-F4.

Table 10: Response rates in study ION-1

	LDV/SOF 12 weeks (n = 214)	L D V / SOF+RBV 12 weeks (n = 217)	LDV/SOF 24 weeks (n = 217)	L D V / SOF+RBV 24 weeks (n = 217)
SVR	9 9 % (210/213)	97% (211/217)	9 8 % (213/217)	99% (215/217)
Outcome for patients	without SVR			
On-treatment virologic failure	0/213ª	0/217	< 1 % (1/217)	0/216
Relapse ^b	< 1% (1/212)	0/217	< 1 % (1/215)	0/216
Other ^c	< 1% (2/213)	3% (6/217)	< 1 % (2/217)	< 1% (2/217)
SVR rates for selecte	ed subgroups			
Genotype				
Genotype 1a	9 8 % (142/145)	97% (143/148)	9 9 % (144/146)	99% (141/143)
Genotype 1b	100% (67/67)	99% (67/68)	9 7 % (67/69)	100% (72/72)
Cirrhosis ^d	!		·	
No	9 9 % (176/177)	97% (177/183)	9 8 % (181/184)	99% (178/180)
Yes	94% (32/34)	100% (33/33)	9 7 % (32/33)	100% (36/36)

a. One patient was excluded from the LDV/SOF 12-week treatment arm and one patient was excluded from the LDV/SOF+RBV 24-week treatment arm as both patients were infected with genotype 4 CHC.

b. The denominator for relapse is the number of patients with HCV RNA < LLOQ at their last on-treatment assessment.

c. Other includes patients who did not achieve SVR and did not meet virologic failure criteria (e.g. lost to follow-up).

d. Patients with missing cirrhosis status were excluded from this subgroup analysis.

Previously treated adults with or without cirrhosis – ION-2 (study 0109) – Genotype 1 ION-2 was a randomised, open-label study that evaluated 12 and 24 weeks of treatment with Ledipasvir/Sofosbuvir with or without ribavirin (randomised 1:1:1:1) in genotype 1 HCV-infected patients with or without cirrhosis who failed prior therapy with an interferonbased regimen, including regimens containing an HCV protease inhibitor. Randomisation was stratified by the presence or absence of cirrhosis, HCV genotype (1a *versus* 1b) and response to prior HCV therapy (relapse/breakthrough *versus* non-response).

Patient disposition	LDV/SOF 12 weeks (n = 109)	LDV/SOF+ RBV 12 weeks (n = 111)	LDV/SOF 24 weeks (n = 109)	LDV/SOF+ RBV 24 weeks (n = 111)	TOTAL (n = 440)	
Age (years): median (range)	56 (24-67)	57 (27-75)	56 (25-68)	55 (28-70)	56 (24-75)	
Male gender	68% (74)	64% (71)	68% (74)	61% (68)	65% (287)	
Race: Black/ African American	22% (24)	14% (16)	16% (17)	18% (20)	18% (77)	
White	77% (84)	85% (94)	83% (91)	80% (89)	81% (358)	
Genotype 1a	79% (86)	79% (88)	78% (85)	79% (88)	79% (347)	
Prior HCV therap	У					
PEG-IFN+RBV	39% (43)	42% (47)	53% (58)	53% (59)	47% (207) ^a	
HCV protease inhibitor + PEG- IFN+RBV	61% (66)	58% (64)	46% (50)	46% (51)	53% (231)ª	
I L 2 8 C C genotype	9% (10)	10% (11)	14% (16)	16% (18)	13% (55)	
FibroTest-Determined Metavir score ^b						
F0-F1	14% (15)	10% (11)	12% (13)	16% (18)	13% (57)	
F2	28% (31)	26% (29)	28% (31)	30% (33)	28% (124)	
F3-F4	58% (63)	64% (71)	58% (63)	54% (60)	58% (257)	
Not interpretable	0% (0)	0% (0)	2% (2)	0% (0)	< 1% (2)	

 Table 11: Demographics and baseline characteristics in study ION-2

a. One patient in the LDV/SOF 24-week treatment arms and one patient in the LDV/ SOF+RBV 24-week treatment arm

were prior treatment failures of a non-pegylated interferon based regimen.

 Non-missing FibroTest results are mapped to Metavir scores according to: 0-0.31 = F0-F1; 0.32-0.58 = F2;

0.59-1.00 = F3-F4.

Table 12: Response rates in study ION-2

	LDV/SOF 12 weeks (n = 109)	L D V / SOF+RBV 12 weeks (n = 111)	LDV/SOF 24 weeks (n = 109)	L D V / SOF+RBV 24 weeks (n = 111)
SVR	9 4 % (102/109)	96% (107/111)	9 9 % (108/109)	99% (110/111)
Outcome for patients	without SVR	·	•	
On-treatment virologic failure	0/109	0/111	0/109	< 1% (1/111)
Relapse ^a	6% (7/108)	4% (4/111)	0/109	0/110
Other ^b	0/109	0/111	< 1% (1/109)	0/111
SVR rates for selecte	d subgroups		·	
Genotype				
Genotype 1a	95% (82/86)	95% (84/88)	99% (84/85)	99% (87/88)
Genotype 1b	87% (20/23)	100% (23/23)	1 0 0 % (24/24)	100% (23/23)
Cirrhosis		·	•	
No	95% (83/87)	100% (88/88) ^c	99% (85/86) ^c	99% (88/89)
Yes ^d	86% (19/22)	82% (18/22)	1 0 0 % (22/22)	100% (22/22)
Prior HCV therapy		•	1	
PEG-IFN+RBV	93% (40/43)	96% (45/47)	1 0 0 % (58/58)	98% (58/59)
H C V protease inhibitor + PEG- IFN+RBV	94% (62/66)	97% (62/64)	98% (49/50)	100% (51/51)

a. The denominator for relapse is the number of patients with HCV RNA < LLOQ at their last on-treatment assessment.

b. Other includes patients who did not achieve SVR and did not meet virologic failure criteria (e.g. lost to follow-up).

c. Patients with missing cirrhosis status were excluded from this subgroup analysis.

d. Metavir score = 4 or Ishak score \geq 5 by liver biopsy, or FibroTest score of > 0.75 and (APRI) of > 2.

Table 13 presents relapse rates with the 12-week regimens (with or without ribavirin) for selected subgroups (see also previous section "Effect of baseline HCV resistance-associated variants on treatment outcome"). In non-cirrhotic patients relapses only occurred in the presence of baseline NS5A RAVs, and during therapy with Ledipasvir/Sofosbuvir without ribavirin. In cirrhotic patients relapses occurred with both regimens, and in the absence and presence of baseline NS5A RAVs.

 Table 13: Relapse rates for selected subgroups in study ION-2

Tuble To: Relapse fales for selected subgroups in study for-2					
	L D V / SOF 12 weeks (n = 109)	L D V / SOF+RBV 12 weeks (n = 111)	LDV/SOF 24 weeks (n = 109)	L D V SOF+RBV 24 weeks (n = 111)	
Number of responders at end of treatment	108	111	109	110	
Cirrhosis	•				
No	5 % (4/86) ^a	0% (0/88) ^b	0% (0/86) ^b	0% (0/88)	
Yes	1 4 % (3/22)	18% (4/22)	0% (0/22)	0% (0/22)	
Presence of baseline NS	5A resistance	-associated subst	titutions ^c		
No	3 % (3/91) ^d	2% (2/94)	0% (0/96)	0% (0/95) ^f	
Yes	2 4 % (4/17) ^e	12% (2/17)	0% (0/13)	0% (0/14)	

a. These 4 non-cirrhotic relapsers all had baseline NS5A resistance-associated polymorphisms.

b. Patients with missing cirrhosis status were excluded from this subgroup analysis.

- c. Analysis (by deep sequencing) included NS5A resistance-associated polymorphisms that conferred > 2.5-fold change in EC₅₀ (K24G/N/R, M28A/G/T, Q30E/G/H/L/K/R/T, L31I/F/M/V, P32L, S38F, H58D, A92K/T, and Y93C/F/H/N/S for genotype 1a and L31I/F/M/V, P32L, P58D, A92K, and Y93C/H/N/S for genotype 1b HCV infection). d. 3/3 of these patients had cirrhosis. e. 0/4 of these patients had cirrhosis.
- f. One patient who achieved a viral load < LLOQ at end of treatment had missing baseline NS5A data and was excluded from the analysis.

HCV/HIV co-infected adults – ERADICATE

ERADICATE was an open-label study to evaluate 12 weeks of treatment with Ledipasvir/ Sofosbuvir in 50 patients with genotype 1 CHC co-infected with HIV. All patients were treatment-naïve to HCV therapy without cirrhosis, 26% (13/50) of patients were HIV antiretroviral naïve and 74% (37/50) of patients were receiving concomitant HIV antiretroviral therapy. At the time of the interim analysis 40 patients have reached 12 weeks post treatment and SVR12 was 98% (39/40).

Patients awaiting liver transplantation and post-liver transplant – SOLAR-1 (see also section 4.4)

SOLAR-1 is an open-label, multicentre study evaluating 12 and 24 weeks of treatment with Ledipasvir/Sofosbuvir + ribavirin in patients with genotype 1 or 4 CHC who have advanced liver disease and/or who have undergone liver transplantation. Seven patient populations are being evaluated (patients with decompensated cirrhosis [CPT B and C] pre-transplant; post-transplant, no cirrhosis; post-transplant CPT A; post-transplant CPT B; post-transplant CPT C; post-transplant fibrosing cholestatic hepatitis).

Preliminary data from the SOLAR-1 study include interim SVR data from a total of 302 genotype 1 patients from across the treatment groups including limited SVR data from 4 patients with fibrosing cholestatic hepatitis. An SVR4 rate of approximately 90% was achieved with Ledipasvir/Sofosbuvir + ribavirin in patients with decompensated cirrhosis

(CPT B or C) for both treatment durations studied (12 or 24 weeks). In post-liver transplant patients without decompensated liver disease, SVR4 rates were > 95%. Among patients with fibrosing cholestatic hepatitis, all 4 have achieved SVR4.

Clinical efficacy and safety in genotype 3 (see also section 4.4)

In a Phase 2 open-label study, the safety and efficacy of Ledipasvir/Sofosbuvir were evaluated with or without ribavirin in 51 treatment-naïve patients with genotype 3 HCV infection, with or without cirrhosis. Patients were treated with Ledipasvir/Sofosbuvir (n = 25) or Ledipasvir/Sofosbuvir + ribavirin (n = 26) for 12 weeks. SVR12 rates were 64% (16/25) and 100% (26/26) in the Ledipasvir/Sofosbuvir and Ledipasvir/Sofosbuvir + ribavirin treatment groups, respectively.

Clinical efficacy and safety in genotype 4 (see also section 4.4)

Two patients with genotype 4d HCV infection were enrolled into the ION-1 study. One patient received Ledipasvir/Sofosbuvir for 12 weeks; another patient received Ledipasvir/Sofosbuvir + ribavirin for 24 weeks. Both achieved SVR12. In a Phase 2 study evaluating Ledipasvir/Sofosbuvir for

12 weeks, 21 genotype 4 patients are being treated. Post-treatment week 12 data is available for 5 patients: all 5 have achieved SVR12. Ledipasvir and Sofosbuvir have demonstrated *in vitro* antiviral activity in genotype 4 viral replicons (see above "Antiviral activity").

Clinical efficacy and safety in other genotypes

Data are currently not available on the safety and efficacy of Ledipasvir/Sofosbuvir in patients infected with HCV genotype 2, 5 or 6.

Paediatric population

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

5.2 Pharmacokinetic properties

Absorption

Following oral administration of Ledipasvir/Sofosbuvir to HCV-infected patients, Ledipasvir median peak plasma concentration was observed at 4.0 hours post-dose. Sofosbuvir was absorbed quickly and the median peak plasma concentrations were observed ~ 1 hour post-dose. Median peak plasma concentration of GS-331007 was observed at 4 hours post-dose.

Based on the population pharmacokinetic analysis in HCV-infected patients, geometric mean steady-state AUC₀₋₂₄ for Ledipasvir (n = 2,113), Sofosbuvir (n = 1,542), and GS-331007 (n = 2,113) were 7,290, 1,320 and 12,000 ng•h/mL, respectively. Steady-state C_{max} for Ledipasvir, Sofosbuvir and GS-331007 were 323, 618 and 707 ng/mL, respectively. Sofosbuvir and GS-331007 AUC₀₋₂₄ and C_{max} were similar in healthy adult subjects and patients with HCV infection. Relative to healthy subjects (n = 191), Ledipasvir AUC₀24 and C_{max} were 24% lower and 32% lower, respectively, in HCV-infected patients. Ledipasvir AUC is dose proportional over the dose range of 3 to 100 mg. Sofosbuvir and GS-331007 AUCs are near dose proportional over the dose range of 200 mg to 400 mg.

Effects of food

Relative to fasting conditions, the administration of a single dose of Ledipasvir/Sofosbuvir with a moderate fat or high fat meal increased the Sofosbuvir AUC_{0-inf} by approximately 2-fold, but did not significantly affect the Sofosbuvir C_{max} . The exposures to GS-331007 and Ledipasvir were not altered in the presence of either meal type. Ledipasvir/Sofosbuvir 90 mg/400 mg film coated tablets can be administered without regard to food.

Distribution

Ledipasvir is > 99.8% bound to human plasma proteins. After a single 90 mg dose of $[^{14}C]$ -Ledipasvir in healthy subjects, the blood to plasma ratio of $[^{14}C]$ -radioactivity ranged between 0.51 and 0.66.

Sofosbuvir is approximately 61-65% bound to human plasma proteins and the binding is independent of drug concentration over the range of 1 μ g/mL to 20 μ g/mL. Protein binding of GS-331007 was minimal in human plasma. After a single 400 mg dose of [¹⁴C]-Sofosbuvir in healthy subjects, the blood to plasma ratio of [¹⁴C]-radioactivity was approximately 0.7.

Biotransformation

In vitro, no detectable metabolism of Ledipasvir was observed by human CYP1A2, CYP2C8, CYP2C9, CYP2C19, CYP2D6 and CYP3A4. Evidence of slow oxidative metabolism via an unknown mechanism has been observed. Following a single dose of 90 mg [¹⁴C]-Ledipasvir, systemic exposure was almost exclusively due to the parent drug (> 98%). Unchanged Ledipasvir is also the major species present in faeces.

Sofosbuvir is extensively metabolised in the liver to form the pharmacologically active nucleoside analogue triphosphate GS-461203. The active metabolite is not observed. The metabolic activation pathway involves sequential hydrolysis of the carboxyl ester moiety catalysed by human cathepsin A or carboxylesterase 1 and phosphoramidate cleavage by histidine triad nucleotide-binding protein 1 followed by phosphorylation by the pyrimidine nucleotide biosynthesis pathway. Dephosphorylation results in the formation of nucleoside metabolite GS-331007 that cannot be efficiently rephosphorylated and lacks antiHCV activity *in vitro*. Within Ledipasvir/Sofosbuvir, GS-331007 accounts for approximately 85% of total systemic exposure.

Elimination

Following a single 90 mg oral dose of [¹⁴C]-Ledipasvir, mean total recovery of the [¹⁴C]radioactivity in faeces and urine was 87%, with most of the radioactive dose recovered from faeces (86%). Unchanged Ledipasvir excreted in faeces accounted for a mean of 70% of the administered dose and the oxidative metabolite M19 accounted for 2.2% of the dose. These data suggest that biliary excretion of unchanged

Ledipasvir is a major route of elimination with renal excretion being a minor pathway (approximately 1%).

The median terminal half-life of Ledipasvir in healthy volunteers following administration of Ledipasvir/Sofosbuvir in the fasted state was 47 hours.

Following a single 400 mg oral dose of [¹⁴C]-Sofosbuvir, mean total recovery of the dose was greater than 92%, consisting of approximately 80%, 14%, and 2.5% recovered in urine, faeces, and expired air, respectively. The majority of the Sofosbuvir dose recovered in urine was GS-331007 (78%) while 3.5% was recovered as Sofosbuvir. This data indicate that renal clearance is the major elimination pathway for GS-331007 with a large part actively secreted. The median terminal half-lives of Sofosbuvir and GS331007 following administration of Ledipasvir/Sofosbuvir were 0.5 and 27 hours, respectively.

Neither Ledipasvir nor Sofosbuvir are substrates for hepatic uptake transporters, organic cation transporter (OCT) 1, organic anion-transporting polypeptide (OATP) 1B1 or OATP1B3. GS-331007 is not a substrate for renal transporters including organic anion transporter (OAT) 1 or OAT3, or OCT2.

In vitro potential for Ledipasvir/Sofosbuvir to affect other medicinal products

At concentrations achieved in the clinic, Ledipasvir is not an inhibitor of hepatic transporters including the OATP 1B1 or 1B3, BSEP, OCT1, OCT2, OAT1, OAT3, multidrug and toxic compound extrusion (MATE) 1 transporter, multidrug resistance protein (MRP) 2 or MRP4. Sofosbuvir and GS-331007 are not inhibitors of drug transporters P-gp, BCRP, MRP2, BSEP, OATP1B1, OATP1B3, OCT1 and GS331007 is not an inhibitor of OAT1, OCT2 and MATE1.

Sofosbuvir and GS-331007 are not inhibitors or inducers of CYP or uridine diphosphate glucuronosyltransferase (UGT) 1A1 enzymes.

Pharmacokinetics in special populations

Race and gender

No clinically relevant pharmacokinetic differences due to race have been identified for Ledipasvir, Sofosbuvir or GS-331007. No clinically relevant pharmacokinetic differences due to gender have been identified for Sofosbuvir or GS-331007. AUC and C_{max} of Ledipasvir were 77% and 58% higher, respectively, in females than males; however, the relationship between gender and Ledipasvir exposures was not considered clinically relevant.

Elderly

Population pharmacokinetic analysis in HCV-infected patients showed that within the age range (18 to 80 years) analysed, age did not have a clinically relevant effect on the exposure to Ledipasvir, Sofosbuvir or GS-331007. Clinical studies of Ledipasvir/ Sofosbuvir included 117 patients aged 65 years and over.

Renal impairment

The pharmacokinetics of Ledipasvir were studied with a single dose of 90 mg Ledipasvir in HCV negative patients with severe renal impairment (eGFR < 30 mL/min by Cockcroft-Gault, median [range] CrCl 22 [17-29] mL/min). No clinically relevant differences in Ledipasvir pharmacokinetics were observed between healthy subjects and patients with severe renal impairment.

The pharmacokinetics of Sofosbuvir were studied in HCV negative patients with mild (eGFR \ge 50 and

< 80 mL/min/1.73 m²), moderate (eGFR \geq 30 and < 50 mL/min/1.73 m²), severe renal impairment

(eGFR < 30 mL/min/1.73 m²) and patients with ESRD requiring haemodialysis following a single

400 mg dose of Sofosbuvir. Relative to patients with normal renal function (eGFR

> 80 mL/min/1.73 m²), the Sofosbuvir AUC_{0-inf} was 61%, 107% and 171% higher in mild, moderate and severe renal impairment, while the GS-331007 AUC_{0-inf} was 55%, 88% and 451% higher, respectively. In patients with ESRD, relative to patients with normal renal function, Sofosbuvir AUC_{0-inf} was 28% higher when Sofosbuvir was dosed 1 hour before haemodialysis compared with 60% higher when Sofosbuvir was dosed 1 hour after haemodialysis. The AUC_{0-inf} of GS-331007 in patients with ESRD administered with Sofosbuvir 1 hour before or 1 hour after haemodialysis was at least 10-fold and 20-fold

higher, respectively. GS-331007 is efficiently removed by haemodialysis with an extraction coefficient of approximately 53%. Following a single 400 mg dose of Sofosbuvir, a 4 hour haemodialysis removed 18% of administered Sofosbuvir dose. The safety and efficacy of Sofosbuvir have not been established in patients with severe renal impairment or ESRD.

Hepatic impairment

The pharmacokinetics of Ledipasvir were studied with a single dose of 90 mg Ledipasvir in HCV negative patients with severe hepatic impairment (CPT class C). Ledipasvir plasma exposure (AUC_{inf}) was similar in patients with severe hepatic impairment and control patients with normal hepatic function. Population pharmacokinetics analysis in HCV-infected patients indicated that cirrhosis had no clinically relevant effect on the exposure to Ledipasvir.

The pharmacokinetics of Sofosbuvir were studied following 7-day dosing of 400 mg Sofosbuvir in HCVinfected patients with moderate and severe hepatic impairment (CPT class B and C). Relative to patients with normal hepatic function, the Sofosbuvir AUC_{0-24} was 126% and 143% higher in moderate and severe hepatic impairment, while the GS-331007 AUC_{0-24} was 18% and 9% higher, respectively. Population pharmacokinetics analysis in HCV-infected patients indicated that cirrhosis had no clinically relevant effect on the exposure to Sofosbuvir and GS-331007.

Body weight

Body weight did not have a significant effect on Sofosbuvir exposure according to a population pharmacokinetic analysis. Exposure to Ledipasvir decreases with increasing body weight but the effect is not considered to be clinically relevant.

Paediatric population

The pharmacokinetics of Ledipasvir, Sofosbuvir and GS-331007 in paediatric patients have not been established (see section 4.2).

5.3 Preclinical safety data

<u>Ledipasvir</u>

No target organs of toxicity were identified in rat and dog studies with Ledipasvir at AUC exposures approximately 7 times the human exposure at the recommended clinical dose.

Ledipasvir was 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* rat micronucleus assays.

Ledipasvir carcinogenicity studies are ongoing.

Ledipasvir had no adverse effects on mating and fertility. In female rats, the mean number of corpora lutea and implantation sites were slightly reduced at maternal exposures 6-fold the exposure in humans at the recommended clinical dose. At the no observed effect level, AUC exposure to Ledipasvir was approximately 7- and 3-fold, in males and females, respectively, the human exposure at the recommended clinical dose.

No teratogenic effects were observed in rat and rabbit developmental toxicity studies with Ledipasvir.

In a rat pre- and postnatal study, at a maternally toxic dose, the developing rat offspring exhibited mean decreased body weight and body weight gain when exposed *in utero* (via maternal dosing) and during lactation (via maternal milk) at a maternal exposure 4 times the exposure in humans at the recommended clinical dose. There were no effects on survival, physical and behavioural development and reproductive performance in the offspring at maternal exposures similar to the exposure in humans at the recommended clinical dose.

When administered to lactating rats, Ledipasvir was detected in plasma of suckling rats likely due to excretion of Ledipasvir via milk.

<u>Sofosbuvir</u>

In repeat dose toxicology studies in rat and dog, high doses of the 1:1 diastereomeric mixture caused adverse liver (dog) and heart (rat) effects and gastrointestinal reactions (dog). Exposure to Sofosbuvir in rodent studies could not be detected likely due to high esterase activity; however, exposure to the major metabolite GS-331007 at doses which cause adverse effects was 16 times (rat) and 71 times (dog) higher than the clinical exposure at 400 mg Sofosbuvir. No liver or heart findings were observed in chronic toxicity studies at exposures 5 times (rat) and 16 times (dog) higher than the clinical exposure. No liver or heart findings were observed in the 2-year carcinogenicity studies at exposures 17 times (mouse) and 9 times (rat) higher than the clinical exposure.

Sofosbuvir was 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.

Carcinogenicity studies in mice and rats do not indicate any carcinogenicity potential of Sofosbuvir administered at doses up to 600 mg/kg/day in mouse and 750 mg/kg/day in rat. Exposure to GS-331007 in these studies was up to 17 times (mouse) and 9 times (rat) higher than the clinical exposure at 400 mg Sofosbuvir.

Sofosbuvir had no effects on embryo-foetal viability or on fertility in rat and was not teratogenic in rat and rabbit development studies. No adverse effects on behaviour, reproduction or development of offspring in rat were reported. In rabbit studies exposure to Sofosbuvir was 6 times the expected clinical exposure. In the rat studies, exposure to Sofosbuvir could not be determined but exposure margins based on the major human metabolite was approximately 5 times higher than the clinical exposure at 400 mg Sofosbuvir.

Sofosbuvir-derived material was transferred through the placenta in pregnant rats and into the milk of lactating rats.

6. PHARMACEUTICAL PARTICULARS

6.1 List of excipients

<u>Tablet core</u> Lactose monohydrate, Microcrystalline cellulose PH 101, Microcrystalline cellulose PH 102, Croscarmellose sodium, Colloidal silicon dioxide, Magnesium stearate

Film-coating

Polyvinyl alcohol - Part.hydrolyzed, Titanium dioxide, Macrogol/PEG, Talc, FD&C Blue #2/Indigo carmine AL 3% - 5%, FD&C Blue #2/Indigo carmine Aluminium lake

6.2 Incompatibilities

Not applicable.

6.3 Shelf life

24 months.

6.4 Special precautions for storage

Do not store above 30°C. Store in the original container.

6.5 Nature and contents of container

Ledipasvir and Sofosbuvir Tablets 90 mg/400 mg are supplied in blue high density polyethylene bottles with a blue polypropylene screw closure containing 28 tablets with a silica gel desiccant.

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

Mylan Laboratories Limited Plot No. 564/A/22, Road No.92, Jubilee Hills Hyderabad - 500034, Telangana, INDIA

8. MARKETING AUTHORISATION NUMBER

TAN 21 HM 0442

9. DATE OF FIRST AUTHORISATION/RENEWAL OF THE AUTHORISATION

2021-11-26

10. DATE OF REVISION OF THE TEXT