Information sheet for ledipasvir with sofosbuvir (Harvoni)

Applications will be considered by the Hepatitis C Treatments Panel (HepCTP) at its regular meetings and approved subject to eligibility according to the Access Criteria are detailed below.

Harvoni may be used in hepatitis C patients of all genotypes. Harvoni is an oral fixed dose combination tablet which contains 400mg sofosbuvir and 90mg ledipasvir. We understand that Harvoni should always be combined with ribavirin in patients with decompensated cirrhosis, where a lower starting dose (200mg mane 400mg nocte) should be used. Supply of both Harvoni and ribavirin will be via direct distribution for patients with approval from the HepCTP Panel.

The Panel may assess urgent applications in between regular meetings.

Please refer to our website https://pharmac.govt.nz/harvoni for links to additional information.

Delivery:

Delivery of Harvoni and ribavirin cannot occur unless the **original scripts** for both treatments have been received by PHARMAC.

Both Harvoni and ribavirin will be delivered from a central distributor to an appropriate address. An appropriate address constitutes: a general practice, clinic, or another suitable address where someone is available to sign for the delivery **at all times** during normal work hours and where the medicines can be appropriately looked after until required.

If approved, the delivery of Harvoni should take between 7-10 days for the treatment to arrive at the nominated address. Please ensure you have discussed this application with the nominated delivery address.

Repeat deliveries will be sent to the nominated address in time to ensure that patients will have continuity of supply noting a patient will require 3 deliveries total (or 6 deliveries total, in the unusual situation where ribavirin is contraindicated).

Access criteria:

Chronic hepatitis C - Advanced disease- ribavirin is not contraindicated

Applications from any relevant practitioner. Approvals valid for 12 weeks for applications meeting the following criteria:

All of the following:

- 1. Patient has chronic hepatitis C (any genotype); and
- 2. Ribavirin treatment is not contraindicated; and
- 3. Any of the following:
 - 3.1 Patient has decompensated cirrhosis (Child-Pugh B or C); or
 - 3.2 Patient has been accepted onto a list for a liver transplant or has received a liver transplant; or
 - 3.3 Patient has essential mixed cryoglobulinaemia with associated purpuric skin rash and; either
 - 3.3.1 Glomerulonephritis; or
 - 3.3.2 Systemic vasculitis.

Chronic hepatitis C - Advanced disease - ribavirin is contraindicated

Applications from any relevant practitioner. Approvals valid for 24 weeks for applications meeting the following criteria:

All of the following:

- 1. Patient has chronic hepatitis C (any genotype); and
- 2. Ribavirin treatment is contraindicated; and
- 3. Any of the following:
 - 3.1 Patient has decompensated cirrhosis (Child-Pugh B or C); or
 - 3.2 Patient has been accepted onto a list for a liver transplant or has received a liver transplant; or
 - 3.3 Patient has essential mixed cryoglobulinaemia with associated purpuric skin rash and; either
 - 3.3.1 Glomerulonephritis; or
 - 3.3.2 Systemic vasculitis.

Application for ledipasvir with sofosbuvir (Harvoni) for Chronic hepatitis C – Advanced disease

Contact details:

HepCTP Coordinator PO Box 10 254 Wellington

Phone: 0800 023 588 (option 4)

Fax: 04 974 4826

Email: hepcpanel@pharmac.govt.nz

-	u send your application in please ensure you have:						
Included an original pr	Included an original prescription for Harvoni (two three month scripts (with different						
dates) for Harvoni will	dates) for Harvoni will be needed, if ribavirin is contraindicated)						
Included an original prescription for ribavirin when applicable							
	I have notified and nominated an appropriate physical delivery address for delivery						
I have notified the patient's GP that I have submitted application							
I have attached an according to the function and full blood	companying clinic letter (including recent liver function test, renal count).						
Patient							
*NHI							
*Gender	*Date of birth						
*Last name							
*First name	Middle name						
*Address							
*DHB of domicile							
Applicant							
*NZMC number	*Title						
*Full name							
*Department or							
Practice address							
*DUD							
*DHB							
*Email address *Phone	Extension						
*Facsimile	Extension						
Are there any others Nurse	s who need to be informed about this application? E.g. He	p C					
Contact name							
Contact email	Contact Facsimile						
Contact address							

If you are not the pa will notify them of the		, please provide the GP's details and PHARMAC staff of this application
Please note, as the applito PHARMAC	icant we exp	pect that you will notify the GP that you submitted this application
GP Name		
GP Practice address		
GP Phone		
Please note the delivery courier will require a sign	address ne nature e.g. a t to a Rural	inic, general practice or other appropriate address) eds to be where someone is available to accept delivery as the A clinic address or GP address. Delivery address or a PO Box. this application.
Clinic or General Pract	tice Name	
Address		
Phone Number		
Additional comments		

	ss Crit		low and provide ev	idence o	f support				
1.	-				•				
	Patie	ent has chronic hepatitis C							
2.							•		
	Is rib	avirin treatment contraindicated?							
	Yes	If ribavirin is contraindicated the patient will need 24 weeks of treatment with Harvoni. You will need to send PHARMAC two 3 month prescriptions (with different dates) for Harvoni.							
	Estimated glomerular filtration rate (eGFR):								
	Please include the reasons why ribavirin is contraindicated for your patient (for example, severe vascular disease, haemoglobinopathy, documented RBV allergy or intolerance):								
	No								
3. Please tick one of the following									
	3a.	Patient	has decompensa	ted cirrh	osis (Chil	d-Pugh E	3 or C);	or	
		Patient has been accepted onto a list for a liver transplant; or							
	3b.		Date accepted waiting:	onto list	t if still				
		Patient has received a liver transplant							
			Date of transpla	ant:					
		Patient I skin rasl	nas essential mix	ed cryo	globulinae	mia with	associa	ated purpuric	
	3c.	Either	Glomeruloneph	ritis					
		Or	Systemic vascu	ulitis					

Supporting evidence				
Please include any relevant additional information, including atta	ching rele	vant clinic letters		
Declaration				
By submitting this form				
by sub-initially time form				
 I confirm that all information provided is correct to the 				
I agree to provide all additional information reasonation	bly reque	ested to PHARMAC, or		
its agent.	profossio	anala informad about		
 I will ensure I keep the patients relevant healthcare professionals informed about treatment with Harvoni. 				
I will notify the patient's GP that I have submitted the	is applica	ation.		
I will notify the nominated delivery address that I have				
Applicant's signature & date:				
	Date:			
	Date.			