Harvoni® (Ledipasvir/Sofosbuvir) Treatment Agreement

If you are considering hepatitis C treatment, please read this treatment agreement carefully and be sure to ask any questions you may have before you sign the form.

In October 2014, the FDA approved ledipasvir combined with sofosbuvir in one tablet (Harvoni®) for the treatment of hepatitis C genotype 1.

Treatment with Harvoni® requires 4 scheduled visits over a 5 month period if your treatment course is 8 weeks, 5 scheduled visits over 6 months if your treatment course is 12 weeks, and 9 scheduled visits over 9 months if your treatment course is 6 months.

PREGNANCY & BREASTFEEDING WARNING

It is not known if Harvoni® will harm an unborn or breastfeeding baby, so it is recommended that women do not get pregnant or breastfeed while taking this medicine.

HOW THE TREATMENT PROCESS WORKS

You will have blood and urine tests.

- These tests will include a pregnancy test for female patients. A urine pregnancy test will be done monthly during a clinic visit.
- Random drug and alcohol tests may be requested.
- At each visit, about 2-3 tubes of blood will be collected. Other examinations and tests may be done during the treatment if your provider feels there is a need.

Provider, select the appropriate treatment regimen and reason:

Harvoni® will be given for 12 weeks if:	
 You do not have cirrhosis and have never been treated before; 	
 You have cirrhosis and have never been treated before; 	
$\hfill\Box$ You do not have cirrhosis and prior treatment failed.	
—— Harvoni® can be given for a shortened course of 8 weeks if you do not cirrhosis, have never been treated before, and have a viral load of <6 million.	have
Harvoni® will be given for 24 weeks if you have cirrhosis and prior treatment faile	d.

Your first three visits will be at the start of treatment (week 0) and weeks 2 and 4 after you begin taking the medication. Week 2 visit will be at the discretion of your provider. After that, the visits will be once each month until you stop taking the medications.

You may need to see your primary care provider more frequently if you are having side effects or problems related to the treatment.

You will have follow-up 3 months after treatment completion. If you have cirrhosis you should continue to have a liver ultrasound every six months and regular clinic visits.

TREATMENT MEDICATIONS AND SIDE EFFECTS

<u>Harvoni</u>[®] is a fixed-dose combination tablet containing ledipasvir 90mg and sofosbuvir 400mg. You will take Harvoni[®] once daily by mouth with or without food. Store the medication at room temperature. If you miss a dose, take the missed dose as soon as you remember the same day. Do not take more than 1 tablet of Harvoni[®] in a day. Take your next dose at your regular time the next day.

• The most common side effects are tiredness and headache.

Tell your healthcare provider if you are taking any of the following medicines, as they are <u>not</u> recommended to be used with Harvoni:

- Amiodarone (Cordarone®, Nexterone®, Pacerone®)
- Carbamazepine (Carbatrol®, Epitol®, Equetro®, Tegretol®)
- Oxycarbazepine (Trileptal®, Oxtellar XR®); Phenytoin (Dilantin®, Phenytek®); Phenobarbitol (Luminal®)
- Rifabutin (Mycobutin®); Rifampin (Rifadin®, Rifamate®, Rifater®, Rimactane®); Rifapentine (Priftin®)
- Rosuvastatin (Crestor®)
- Simeprevir (Olysio®)
- St. John's wort (Hypericum perforatum) or a product that contains St. John's wort
- Tipranavir (Aptivus®) used in combination with ritonavir (Norvir®)

Tell your healthcare provider if you are taking any of the following medicines, as they require dose adjustment and/or monitoring:

- An antacid that contains aluminum or magnesium hydroxide. If you take an antacid during treatment with Harvoni®, take the antacid 4 hours before or 4 hours after you take Harvoni®.
- Twice daily medicine for indigestion, heartburn, or stomach ulcers, such as famotidine (Pepcid AC®) no more than 40 mg twice daily, must be taken at the same time or 12 hours apart from Harvoni®. Nizatidine (Axid®), cimetidine (Tagamet®), and ranitidine (Zantac®) have not been studied with Harvoni® and so no dosing recommendations can be given.
- Once daily medications for indigestion, heartburn, or stomach ulcers, such as omeprazole (Prilosec®) no more than 20 mg daily, must be taken at the same time as Harvoni®. Esomeprazole (Nexium®), lansoprazole (Prevacid®), rabeprazole (Aciphex®), and pantoprazole (Protonix®) have not been studied with Harvoni® and so no dosing recommendations can be given.
- Digoxin (Lanoxin®)
- Efavirenz, emtricitabine, tenofovir disoproxil fumarate (ATRIPLA®)
- Elvitegravir, cobicistat, emtricitabine, tenofovir disoproxil fumarate (STRIBILD®)
- Tenofovir disproxil fumarate (VIREAD®, TRUVADA®) used in combination with atazanavir (Reyataz®) and ritonavir (Norvir®), darunavir (Prezista®) and ritonavir (Norvir®), or used in combination with lopinavir and ritonavir (Kaletra®)

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PLEASE NOTE:

You must let your medical, mental health, dental providers, and pharmacist(s) know that you are taking Harvoni® prior to starting any new medications. You must let your healthcare providers know about any new medications you are prescribed before starting them. This includes vitamins and other supplements.

Hepatitis C treatment should not cause pain that requires narcotic pain medication.

BENEFITS OF TREATMENT

In most cases, hepatitis C will respond to treatment as determined by a blood test that measures the presence and amount of hepatitis C in the blood. If you have no hepatitis C in your blood 12 weeks **after** the end of treatment, this is considered a "sustained virologic response" and in 99% of persons is a cure. Your chance of achieving a sustained virologic response depends on the hepatitis C genotype, how much hepatitis C virus you have in your blood at the beginning of treatment, any past treatment response, and how much liver damage you have had prior to treatment.

It is possible that you may develop some serious side effects, which will require you to stop the treatment. You may still benefit from treatment even if it does not get rid of your hepatitis C, as it may slow down the disease. You may choose to stop treatment at any time.

In Clinical Trials:

Persons with genotype 1 who were treatment-naïve (never treated before), did not have cirrhosis and were treated with Harvoni® for 12 weeks had a 99% response (cure) rate. Those who had a baseline viral load of less than 6 million and were treated for 8 weeks had a 97% response rate.

Persons with cirrhosis who were treatment-naïve had a 94% response rate.

Persons without cirrhosis in whom prior treatment failed and were treated for 12 weeks had a 95% response rate.

Persons with cirrhosis in whom prior treatment failed and were treated for 24 weeks had a 100% response rate.

WHOM TO CALL

If you have any questions about treatment, contact your primary care provider at . .

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TREATMENT AGREEMENT

To receive treatment, please review the following statements and initial beside the responses:

Provider's Name (PLEASE PRINT)	Provider's Signature/Title	 Date
Patient's Name (PLEASE PRINT)	Patient's Signature	Date
the information has been explained to	_	ia, or the meaning of
My signature below means that I hav	e read this treatment agreement a	nd/or the mouning of
razors or nail clippers and covering cut	-	, 1000
do so, I will contact my provider. I will protect myself and others	from hepatitis C by not sharing need	lles, toothbrushes.
I will do my best to take my me	dications as prescribed by my provi	der. If I am unable to
it is in the best interest of my health a	nd welfare.	
I understand that my hepatitis (I understand that my provider (er feels that stopping
provider or nurse know right away.	may not recoond to treatment	
<i>-</i> -	medications or side effects that bo	ther me, I will let my
•	cally sterile or post-menopausal.	
understand that my treatment will be		me on treatment.
treatment. As a female taking Harvoni® Lw	ill not get pregnant or breastfeed wh	nile on treatment
required to evaluate my health and we	ell-being during treatment and the e	ffectiveness of
	will be stopped if I cannot attend ap	opointments as
this ahead of time and I will reschedule	• • • • • • • • • • • • • • • • • • • •	THE PROVIDER KNOW
I am willing to visit the clinic and length of the treatment. If I am unable	d see a provider on a regular schedu	
conditions (depression, history of suici		•
blood pressure, diabetes, high cholest		• • •
I will tell my provider if I have		ch as heart disease, high
I have not abused alcohol or oth pain medications) within the last 6 mo		ocame, prescription
I agree <u>not</u> to drink alcohol or u		

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