



2020 General Hepatitis C Treatment Coverage Determination Request

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(You must complete all 4 pages.)

Fax completed form to: 1-800-408-2386

For urgent requests, please call: 1-800-414-2386

Patient information		Prescriber information			
Patient name		Today's date	Physician specialty		
Patient insurance ID number		Physician name		NPI/DEA number	
Patient address, city, state, ZIP		Physician address, city, state, ZIP			
Patient home telephone number		M.D. office telephone number			
Gender <input type="checkbox"/> Male <input type="checkbox"/> Female	Patient date of birth	M.D. office fax number			
Requested regimen: (check all that apply)		Please indicate if this is a new start or continuation of therapy and specify the requested duration of therapy. If continuation of therapy, please indicate start date.			
Medication	Dose/Frequency	New start	Continuation	Initial start date	Duration of therapy
<input type="checkbox"/> Daklinza® (daclatasvir)		<input type="checkbox"/>	<input type="checkbox"/>		
<input type="checkbox"/> Epclusa® (sofosbuvir/velpatasvir)-requesting brand		<input type="checkbox"/>	<input type="checkbox"/>		
<input type="checkbox"/> sofosbuvir/velpatasvir 400mg;100 mg tablet-requesting generic		<input type="checkbox"/>	<input type="checkbox"/>		
<input type="checkbox"/> Harvoni® (ledipasvir/sofosbuvir)-requesting brand		<input type="checkbox"/>	<input type="checkbox"/>		
<input type="checkbox"/> ledipasvir/sofosbuvir 90mg;400mg tablet-requesting generic		<input type="checkbox"/>	<input type="checkbox"/>		
<input type="checkbox"/> Intron-A® (interferon alfa-2b)		<input type="checkbox"/>	<input type="checkbox"/>		
<input type="checkbox"/> Mavyret® (glecaprevir/pibrentasvir)					
<input type="checkbox"/> Pegasys® (pegylated interferon alfa-2a)		<input type="checkbox"/>	<input type="checkbox"/>		
<input type="checkbox"/> Peg-Intron® (pegylated interferon alfa-2a)		<input type="checkbox"/>	<input type="checkbox"/>		
<input type="checkbox"/> ribavirin *prior authorization (PA) not required		<input type="checkbox"/>	<input type="checkbox"/>		
<input type="checkbox"/> Sovaldi® (sofosbuvir)		<input type="checkbox"/>	<input type="checkbox"/>		
<input type="checkbox"/> Sylatron® (pegylated interferon alfa-2b)		<input type="checkbox"/>	<input type="checkbox"/>		
<input type="checkbox"/> Technivie® (ombitasvir/paritaprevir/ritonavir)		<input type="checkbox"/>	<input type="checkbox"/>		
<input type="checkbox"/> Viekira Pak® (ombitasvir/paritaprevir/ritonavir/dasabuvir)		<input type="checkbox"/>	<input type="checkbox"/>		
<input type="checkbox"/> Vosevi® (sofosbuvir/velpatasvir/voxilaprevir)		<input type="checkbox"/>	<input type="checkbox"/>		
<input type="checkbox"/> Zepatier® (elbasvir/grazoprevir)		<input type="checkbox"/>	<input type="checkbox"/>		
<input type="checkbox"/> Other: _____		<input type="checkbox"/>	<input type="checkbox"/>		

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Diagnosis and medical information

Diagnosis (If treatment experienced, list all components of previous treatment regimens and describe outcome)

Chronic, hepatitis C treatment naive Chronic, hepatitis C treatment experienced Other diagnosis/(ICD 10): _____

Yes No **Has the diagnosis of chronic hepatitis C virus (HCV) infection been confirmed by the presence of HCV RNA in the serum prior to starting treatment?**

Please provide HCV RNA viral load: _____ Date: _____

What is the patient's HCV genotype?

1a 1b 2 3 4 5 6 Other (specify): _____

Medical history (Please check all boxes that apply and include office notes)

Yes No Hepatitis B coinfection Yes No HIV coinfection Yes No End stage renal disease
 Yes No Hepatocellular carcinoma Yes No Received a liver transplant Yes No Received a kidney transplant

Which of the following TESTS were used to determine LIVER STAGE? (Please check all boxes that apply and include medical records as supporting documentation)

Liver biopsy Metavir Fibroscan APRI ARFI SWEI Other (specify): _____

What is the patient's FIBROSIS STAGE?

F1 F2 F3 F4

Which of the following best describe patient's liver disease, based on liver staging tests (liver biopsy, Metavir, Fibroscan, etc.), radiological imaging, physiologic or clinical findings? (Please check box that best describe patient's liver disease and include medical records as supporting documentation)

Fibrosis stage F1, F2, or F3: No cirrhosis **Fibrosis stage F4:** Compensated cirrhosis **OR** Decompensated cirrhosis

Was testing for the presence of virus with NS5A resistance-associated polymorphisms performed? Yes No

If testing was done, does the patient have NS5A resistance-associated polymorphism at amino acid positions 28, 30, 31, or 93? Yes No

If testing was not done, provide a reason for why testing was not done:

Has patient been screened for the presence of virus with the NS3 Q80K polymorphism at baseline? Yes No

If screening was done, does patient have Q80K polymorphism? Yes No

If screening was not done, provide a reason for why screening was not done:

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Please check all boxes that apply:

1. All covered Part D drugs on any tier of the plan's formulary would not be as effective for the enrollee as the requested formulary drug and/or would likely have adverse effects for the enrollee.

2. Yes No Is the prescriber a gastroenterologist, hepatologist, or infectious disease specialist OR did the prescriber obtain a consult from one of these specialists? If NO, complete section below:

Please complete this section below only if your patient does not meet the standard requirements listed above.

Please explain why your patient should be considered for exception although not meeting the plan's suggested PA criteria. Statement should include specifically which requirement is not met and why patient should be exempt from meeting this requirement. (Please note any information that is incomplete or illegible will delay the review process.)

3. For TREATMENT EXPERIENCED patients, please complete this section:

Yes No Has patient been treated with ribavirin and/or peginterferon alfa?

If yes, list all components of previously treated regimen below and check the box that best describe treatment outcome:

HCV regimen	Treatment duration/dates	Treatment outcome
		<input type="checkbox"/> Relapsed <input type="checkbox"/> Partial responder <input type="checkbox"/> Non-responder <input type="checkbox"/> Toxicities <input type="checkbox"/> Discontinued therapy prior to completing full course <input type="checkbox"/> Other: _____
		<input type="checkbox"/> Relapsed <input type="checkbox"/> Partial responder <input type="checkbox"/> Non-responder <input type="checkbox"/> Toxicities <input type="checkbox"/> Discontinued therapy prior to completing full course <input type="checkbox"/> Other: _____

Yes No Has patient been treated with regimens containing Eplclusa®, Incivek®, Harvoni®, Victrelis®, Olysio®, Sovaldi®, Zepatier®, Daklinza®, Technivie®, Mavyret®, Vosevi® or Viekira® Pak?

If yes, list all components of previously treated regimen below and check the box that best describe treatment outcome.

HCV regimen	Treatment duration/dates	Treatment outcome
		<input type="checkbox"/> Relapsed <input type="checkbox"/> Partial responder <input type="checkbox"/> Non-responder <input type="checkbox"/> Toxicities <input type="checkbox"/> Discontinued therapy prior to completing full course <input type="checkbox"/> Other: _____
		<input type="checkbox"/> Relapsed <input type="checkbox"/> Partial responder <input type="checkbox"/> Non-responder <input type="checkbox"/> Toxicities <input type="checkbox"/> Discontinued therapy prior to completing full course <input type="checkbox"/> Other: _____

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Please check all boxes that apply (continued):

4. Will patient be taking ribavirin with the requested regimen?

Yes (ribavirin is covered without PA) No, patient is intolerant/ineligible for ribavirin

5. Other supporting information

*NOTE: All exception requests require prescriber supporting statements. Additionally, requests that are subject to prior authorization (or any other utilization management requirement), may require supporting information. Please attach supporting information, as necessary, for your request.

I attest that the medication requested is medically necessary for this patient. I further attest that the information provided is accurate and true, and that documentation supporting this information is available for review if requested by the health plan sponsor, or, if applicable, a state or federal regulatory agency. I understand that any person who knowingly makes or causes to be made a false record or statement that is material to a claim ultimately paid by the United States government or any state government may be subject to civil penalties and treble damages under both the federal and state False Claims Acts. See, e.g., 31 U.S.C. §§ 3729-3733. By signing this form, I represent that I have obtained patient consent as required under applicable state and federal law, including but not limited to the Health Information Portability and Accountability Act (HIPAA) and state re-disclosure laws related to HIV/AIDS.

Prescriber signature

Date