

Hepatitis C Medication Request Form

Fax: 617.673.0956 Mail: CarePartners of Connecticut 705 Mount Auburn Street Watertown, MA 02472 Attn: Pharmacy Utilization Management Department Member Information Last Name: First Name: Member ID#: Member DOB: Prescriber Information Prescribing Clinician: Phone #: Specialty (required): Secure Fax #: NPI #: DEA/xDEA: Prescriber Point of Contact Name (POC) (if different than provider):	Today's date//			
Mail: CarePartners of Connecticut 705 Mount Auburn Street Watertown, MA 02472 Attn: Pharmacy Utilization Management Department Member Information Last Name: Member DOB:	Submit form to:			
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)-F2), please describe the medical necessity	for requesti	ng
Was the staging of hepatic fibrosis performed by a specialist through one of the following? ☐ Yes ☐ No	Is the medication prescribed by a gastroenterologist, infection	us disease specialist, or hepatologist?	☐ Yes	□ No
	Was the staging of hepatic fibrosis performed by a special	list through one of the following?	☐ Yes	□ No

Please check all that apply and attach documentation tests:	including medical records and re	sults of diagnostic			
☐ Liver biopsy confirming METAVIR score	☐ Transient elastography (Fib	roscan) score			
☐ Fibrotest (FibroSURE) score of greater ☐ Radiological imaging					
□ APRI score	—				
☐ Physical findings or clinical evidence consist	tent with cirrhosis as attested by t	he prescriber			
Is there documented evidence of chronic liver disease	e, or in the absence of chronic liver	disease, serologic	☐ Yes	□ No	
evidence of persistent infection for at least six month			L res	□ No	
Does the patient have HIV coinfection?			☐ Yes	□ No	
Has Hepatitis B screening been performed?			☐ Yes	□ No	
If patient has active Hepatitis B infection, has the risk of Hepatitis B reactivation been assessed? Caution: FDA has warned about the risk of Hepatitis B reactivating in some patient treated with direct acting antiviral agents for Hepatitis C. AASLD recommends treating Hepatitis B concurrently or prior to Hepatitis C treatment.			□ Yes	□ No	
Does the patient have severe renal impairment or end Confirm the patient's GFR range: \Box 0 – 14 \Box		ialysis?	□ Yes	□ No	
Has the patient been previously treated for Hepatitis	C and failed treatment?				
If yes, when?What treatment	t(s)?				
Response to treatment: Relapsed Partial re	esponse		☐ Yes	□ No	
□ Null response (<2 log reduction in HCV RNA at week 12)					
Adverse reaction? Yes No					
HCV RNA levels:					
Baseline within 6 months of beginning treatment (required):IU/mL Date of lab work:					
Post-therapy:					
12 weeks after completion of treatment: IU/mL Date of lab work:					
Has there been confirmation that the patient does no Q80K polymorphism? (Olysioonly)	ot have a genotype 1a with NS3				
Qook polymorphism: (Olystoomy)		■ Unknown	☐ Yes	□ No	
Has there been confirmation that the patient does i	not have a genotype 1a with a				
baseline NS5A polymorphism? (Zepatier only)	5 31				
		☐ Unknown	☐ Yes	□ No	
Will hepatic laboratory testing be performed prior to therapy, at treatment week 8, and as clinically					
indicated?			□ Yes	□ No	
Does the patient have a diagnosis of hepatocellular carcinoma that meets Milan criteria?		☐ Yes	□ No		
If the patient requires a dosage form other than ribavirin 200mg capsules or tablets, document clinical reason and provide do					
form.					
Dosage form:					
Clinical reason:					
Are any of the following statements true?					
☐ Patient is pregnant or is planning to become p	regnant within 6 months after co	mpletion of treatm	ient		

☐ Patient is male with a female partner who is pregnant or is planning to become pregnant within 6 of treatment	months afte	r completion			
☐ None of the above					
Is the member currently awaiting a liver transplant?	☐ Yes	□ No			
Does the member have cirrhosis? If yes, please choose one:					
☐ Compensated (Child-Turcotte-Pugh Class A; no major complication of cirrhosis)	☐ Yes	□ No			
☐ Decompensated (Child-Turcotte-Pugh Class B or C)					
Is the patient being managed in a liver transplant center?	☐ Yes	□ No			
Is the member actively participating in illicit substance abuse or alcohol abuse?	☐ Yes	□ No			
Is there documented attestation that the member has been assessed for potential nonadherence?	☐ Yes	□ No			
Is the member is receiving substance or alcohol abuse counseling services or seeing an addiction specialist as an adjunct to HCV treatment?	☐ Yes	□ No			
Has a treatment plan been developed and discussed with the patient?	☐ Yes	□ No			
Did the prescriber identify any potential issues with adherence? If yes, please describe:	☐ Yes	□ No			
Have drug interactions been reviewed and evaluated?	☐ Yes	□ No			
Does this member reside in long-term care?					
Is the member enrolled in Hospice?					
Is the drug related to the terminal illness or related conditions?					
Provide an explanation of why the drug being prescribed is unrelated to the terminal illness/related conditions:					
Is this a request for a formulary tier exception (the member's drug plan charges a higher copayment for the drug prescribed than it charges for another drug that treats the condition, and I want to pay the lower copayment – excludes nonformulary drugs and drugs on the specialty tier)? Yes* No					
*If yes, a supporting statement from the prescribing physician is required. Please specify the request: (1) formulary or preferred drugs contraindicated or tried and failed, or tried and not as effective as requested drug; (2) if therapeutic failure, length of therapy on each drug and adverse outcome; (3) if not as effective, length of therapy on each drug and outcome					
By checking the following box, I certify that applying the standard review time frame may seriously life, health, or ability to attain, maintain, or regain maximum function.		my patient's			
I certify that the information provided is accurate and complete to the best of my knowledge, and I understand that any falsification, omission, or concealment of material fact may subject me to civil or criminal liability.					
Prescriber signature (STAMP NOT ACCEPTED)	Date				