MASSACHUSETTS STANDARD FORM FOR HEPATITIS C MEDICATION PRIOR AUTHORIZATION REQUESTS

*Some plans might not accept this form for Medicare or Medicaid requests.

A. Destination			
Health Plan or Prescription Plan Name: Optum Rx			
Health Plan Phone: 1-800-711-4555	Health Plan Fax: 1-844-403-1027		
B. Patient Information			
Patient Name: DOB:	Gender: Male Female Other:		
Member ID #:			
C. Prescriber Information			
Prescribing Clinician:	Phone #:		
Specialty:	Secure Fax #:		
NPI #:	DEA #:		
Prescriber Point of Contact Name (POC) (if different than prescriber):			
POC Phone #:	POC Secure Fax #:		
POC Email (not required):			
Prescribing Clinician or Authorized Representative Signature:			
Date:			
D. Medication Information			
Check if Expedited Review/Urgent Request: [In checking this box, I attest to the fact that this request meets the definition and criteria for expedited review and is an urgent request.)			
□ Daklinza □ Epclusa □ Harvoni □ Olysio □ Ribavirin Generic □ Ribavirin Branded			
Sovaldi			
Requested Duration of Treatment: weeks			
Type of Therapy: Initial Continuation — weeks remaining:			
Anticipated or actual start date:			
Is the medication prescribed by, or in consultation with, a gastroenterologist, infectious disease specialist, or hepatologist? \Begin{align*} Yes \Boxed* No			
For Zepatier only: Has there been confirmation that the patient does not have a genotype 1a with a baseline NS5A polymorphism?			
☐ Yes ☐ No ☐ Unknown			
For Ribavirin only: Does the patient require a dosage form other than generic ribavirin 200 mg capsules or tablets? Yes If yes, please specify the following:			
Dosage form requested:			
Clinical reason for use:			
Are any of the following statements true?			
Patient is pregnant or plans to become pregnant within 6 months of completing treatment			
Patient is male with a female partner who is pregnant or plans to become pregnant within 6 months of completing treatment			
Patient has contraindications or intolerance to Ribavirin			

(continued on next page)

E. Patient Clinical Information			
*Please refer to plan-specific criteria for details related to required information.			
Diagnosis: ☐ B18.2 Hepatitis C (chronic) ☐ C	Other:		
HCV Genotype: ☐ 1 ☐ 1a ☐ 1b ☐ 2 [□3 □4 □5 □6	Stage of Hepatic Fibrosis: FO F1 F2 F3 F4	
		If F4: ☐ Compensated ☐ Decompensated	
Check all methods of assessment that apply	and include result:		
Method		Result	
Liver biopsy		See above	
☐ Transient elastography (FibroScan)		kPa	
☐ Shear wave elastography		kPa	
☐ MRE		kPa	
☐ FibroSure (FibroTest)			
☐ Echosens Fibrometer			
Fibrospect			
☐ APRI			
APN			
			
☐ Hepascore ☐ Other:			
Does the patient have HIV coinfection? Yes No Unknown			
Is the patient status post liver transplant? Yes No			
Confirm the patient's GFR range: 0–14 15–29 30 or greater (Please specify.)			
HCV RNA levels:			
Week 8 of treatment (if continuation request):	IU/mL Date	of lab work: IU/mL Date of lab work:	
Week of treatment (in contamation request).	Previous Treatn		
Has the patient been previously treated for Hepatitis C and failed treatment? No			
Adverse Reaction? \(\text{Yes} \) No			
	D		
Drug Name	Date of treatment (MM/YY)	<u> </u>	
		Relapsed	
		☐ Partial response☐ Null response (<2 log reduction in HCV RNA at Week 12)	
		☐ Did not complete	
		Briefly describe details:	
		Relapsed	
		☐ Partial response	
		☐ Null response (<2 log reduction in HCV RNA at Week 12)	
		☐ Did not complete	
		☐ Briefly describe details:	
		Relapsed	
		☐ Partial response	
		☐ Null response (<2 log reduction in HCV RNA at Week 12)	
		☐ Did not complete ☐ Briefly describe details:	
		☐ briefly describe details.	
Additional information pertinent to this request:			

Providers should consult the health plan's coverage policies, member benefits, and medical necessity guidelines to complete this form.

Providers may attach any additional data relevant to medical necessity criteria.