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 Name: Tufts Health Plan				
Health Plan Phone: 888-884-2404 Fax: 617-673-0988 Online Prior Authorization: https://point32health.promptpa.com				
B. Patient Information				
Patient Name: DO	DB:	Member ID #:		
Sex Assigned at Birth: ☐ Male ☐ Female ☐ "X" or Intersex				
Current Gender: Male Female Transgender Male Transgender Female Other				
Plans do not discriminate based on race, color, national origin, age, stereotyping).	e, disability, religion, creed, :	sexual orientation, or sex (including gender identity and gender		
C. Prescriber Information				
Prescribing Clinician:	Phone #:			
Specialty:	Secure Fax #:	Secure Fax #:		
NPI #:	DEA #:	DEA #:		
Prescriber Point of Contact Name (POC) (if different than prescri	riber):			
POC Phone #:	POC Secure Fa	POC Secure Fax #:		
POC Email (not required):				
Prescribing Clinician or Authorized Representative Signatu	ure:			
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 ☐ Technivie ☐ Viekira Pak ☐ Viekira XR ☐ Zepatier ☐ Vosevi ☐ Mavyret ☐ Other				
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? 🗌 Yes 🔻 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				

E. Patient Clinical Information		
*Please refer to plan-specific criteria for details	s related to required informat	ion.
Diagnosis: ☐ B18.2 Hepatitis C (chronic) ☐ C)ther:	
HCV Genotype: ☐ 1 ☐ 1a ☐ 1b ☐ 2 [3	Stage of Hepatic Fibrosis: F0 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		
Fib-4		
Hepascore		
Other:		
Does the patient have HIV coinfection? 🗌 Yes	□ No □ Unknown	
Is the patient status post liver transplant?	S No	
Confirm the patient's GFR range: 0-14] 15–29 ☐ 30 or greater (<i>Plea</i>	ase specify.)
HCV RNA levels: Baseline (most recent): Week 8 of treatment (if continuation request):		of lab work: IU/mL Date of lab work:
	Previous Treat	ments
Has the patient been previously treated for Hep	atitis C and failed treatment? [☐ Yes ☐ No
Adverse Reaction? Yes No		
Drug Name	Date of treatment (MM/YY	Response to treatment
		☐ 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:
Additional information pertinent to this request:		

F. Exceptions to Step Therapy Please complete the applicable section(s).
Is the alternative drug required under the step therapy protocol contraindicated, or will likely cause an adverse reaction in, or physical or mental harm to the member? \square Yes \square No
If yes, briefly describe details of contraindication, adverse reaction, or harm:
Is the alternative drug required under the step therapy protocol expected to be ineffective based on the known clinical characteristics of the member and the known characteristics of the alternative drug regiment? Yes No
If yes, briefly describe details of known clinical characteristics of member and alternative drug regimen:
Has the member previously tried the alternative drug required under the step therapy protocol or another alternative drug in the same pharmacologic class or with the same mechanism of action, and such alternative drug was discontinued due to lack of efficacy or effectiveness, diminished effect, or an adverse event? \square Yes \square No
If yes, please provide details for the previous trial:
Drug Name: Dates/Duration of Use:
Did the member experience any of the following? Adverse Reaction Inadequate Response
Briefly describe details of adverse reaction or inadequate response:
Drug Name: Dates/Duration of Use:
Did the member experience any of the following? Adverse Reaction Inadequate Response
Briefly describe details of adverse reaction or inadequate response:
Is the member stable on the requested prescription drug prescribed by the health care provider, and switching drugs will likely cause an adverse reaction in or physical or mental harm to the member? Yes No
If yes, briefly provide details of the adverse reaction or physical or mental harm:

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.