## 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 Prescript	ion Plan Name:	MedImpact						
Health Plan Phone:	800-788-2949			Health Plan F	ax:	858-790-7100		
B. Patient Information	n							
Patient Name			DOB		Gender	$\square$ Male $\square$ Female $\square$ 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:						
🗌 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						
<i>For Zepatier only:</i> Has there been confirmation that the patient does not have a genotype 1a with a baseline NS5A polymorphism? Yes No Unknown						
<i>For Ribavirin only:</i> Does the patient require a dosage form other than generic ribavirin 200 mg capsules or tablets? Yes No 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 related to required information.   Diagnosis: B18.2 Hepatitis C (chronic)   Other:						
HCV Genotype: 1 1 1a 1b 2		Stage of Hepatic Fibrosis:         F0         F1         F2         F3         F4				
		If F 4: Compensated Decompensated				
Check all methods of assessment that apply						
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?						
Is the patient status post liver transplant? $\Box$ Yes						
<b>Confirm the patient's GFR range:</b> 0–14	15–29 [] 30 or greater ( <i>Pied</i>	se speciny.)				
HCV RNA levels: Baseline (most recent):	IU/mL Date	of lab work:				
Week 8 of treatment (if continuation request):		IU/mL Date of lab work:				
	Previous Treatr	nents				
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				
		$\Box$ Null response ( <2 log reduction in HCV RNA at Week 12)				
		Did not complete				
		Relapsed				
		Partial response				
		<ul> <li>Null response ( &lt;2 log reduction in HCV RNA at Week 12)</li> <li>Did not complete</li> </ul>				
		Briefly describe details:				
		☐ Relapsed ☐ Partial response				
		$\square$ Null response ( <2 log reduction in HCV RNA at Week 12)				
		Did not complete				
		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.