

STANDARDIZED ONE PAGE PHARMACY PRIOR AUTHORIZATION FORM

Mississippi Division of Medicaid, Pharmacy Prior Authorization Unit, 550 High St., Suite 1000, Jackson, MS 39201

□ Medicaid Fee for Service/Change Healthcare

Fax to: 1-877-537-0720 Ph: 1-877-537-0722

https://medicaid.ms.gov/providers/pharmacy/pharmacy-prior-authorization/

☐ Magnolia Health/Envolve Pharmacy Solutions Fax to: 1-877-386-4695 Ph: 1-866-399-0928 https://www.magnoliahealthplan.com/providers/pharmacy.html

□ **UnitedHealthcare**/OptumRx

Fax to: 1-866-940-7328 Ph: 1-800-310-6826 http://www.uhccommunityplan.com/health-professionals/ms/pharmacy-program.html

□ Molina Healthcare/CVS Caremark

Fax to: 1-844-312-6371 Ph: 1-844-826-4335 http://www.molinahealthcare.com/providers/ms/medicaid/pages/home.aspx

BENEFICIARY INFORMATION						
Beneficiary ID:	DOB://					
Beneficiary Full Name:						
PRESCRIBER INFORMATION						
Prescriber's NPI:						
Prescriber's Full Name:	Phone:					
Prescriber's Address:	FAX:					
PHARMACY INFORMATION						
Pharmacy NPI:						
Pharmacy Name:						
Pharmacy Phone:	Pharmacy FAX:					
CLINICAL INFORMATION						
Requested PA Start Date: Requested PA En	d Date:					
Drug/Product Requested:	Quantity:					
Days Supply: RX Refills: Diagnosis or ICI	D-10 Code(s):					
Hospital Discharge	ditional Medical Justification Attached					
Medications received through coupons and/or samples are not acceptable as justification.						
PLEASE COMPLETE AND FAX DRUG SPECIFIC CRITERIA/ADDITIONAL DOCUMENTATION FORM FOUND BELOW						
Prescribing provider's signature (signature and date stamps, or the signature of anyone other than the provider, are not acceptable)						
I certify that all information provided is accurate and appropriately documented in the patient's medical chart.						
Signature required:	Date:					
Printed name of prescribing provider:						

FAX THIS PAGE

• Note annotations and respective definitions as described on "Hepatitis C Therapy PA Request Guidelines for Regimen Selection and Definitions" page



Hepatitis C Therapy PA Request

Guidelines for Regimen Selection and Definitions

The Mississippi Division of Medicaid (DOM) will approve Hepatitis C treatment PA requests for members who meet the following criteria:

- Requested regimen is compliant with latest American Association for The Study of Liver Diseases (AASLD)/Infectious Diseases Society of America (ISDA) Recommendations for Testing, Managing, and Treating Hepatitis C <u>http://www.hcvguidelines.org/full-report-view</u> AND
- The regimens listed for each clinical scenario below are the preferred regimens for MS Medicaid beneficiaries, based on clinical and cost considerations and are consistent with AASLD/IDSA guidelines.
 OR
- Pediatric formulations of direct acting antivirals (DAA) with FDA approval will be approved for those patients
 under the age of eighteen when used according to the table below for treatment naïve children as well as in
 accordance with current AASLD guidelines including for indication and age <u>Prior authorization is still required
 prior to the first dose.</u>

GT	Age (years)	Weight (kg)	Drug/Dose	Weeks
1,4,5,6	3-5	<17	Harvoni 33.75/150 mg pellet pack	12
		17 to <35	Harvoni 45/200 mg pellet pack or tablet	12
		<u>></u> 35	Harvoni 90/400 mg tablet	12
Any	<u>></u> 6	<u>></u> 17 to < 30	Epclusa 200/50 mg tablet	12
		<u>></u> 30	sofosbuvir/velpatasvir 400/100 mg tablet	12
Any	<u>></u> 12	<u>></u> 45	Mavyret 100/40 mg tablets -OR-	8
			sofosbuvir/velpatasvir 400/100 mg tablet	12

<u>OR</u>

• Clinical rationale is provided for using a regimen that is beyond those within the current guidelines, or for selecting regimens using non-preferred drugs on the Mississippi Division of Medicaid Universal Preferred Drug List

On the PA Request Form, which must be approved prior to the 1st dose, document the clinical condition supporting the requested regimen. Unless otherwise specified, you do not need to attach documentation to the form. You are attesting to the fact that documentation is available in the patient chart for all information you provide.

DEFINITIONS/ANNOTATIONS USED ON PA FORM:

∇ Low Dose Ribavirin = 600 mg/day and increase as tolerated

- **Ribavirin-Ineligible** (documentation exists in the patient's chart for at least one of the following):
 - Hypersensitivity to RBV
 - History of severe or unstable cardiac disease
 - Pregnant women and men with pregnant partners
 - Diagnosis of hemoglobinopathy (e.g., thalassemia major, sickle cell anemia)
 - Baseline platelet count < 70,000 cells/mm3
 - ANC < 1500 cells/mm3
 - Hb < 12 gm/ml in women or <13 g/dl in men

RENAL DYSFUNCTION Patients with CrCl <50 ml/mm should not be treated with ribavirin.

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Preferred Direct Acting Antivirals

Mavyret (glecaprevir/pibrentasvir) 300/120 mg sofosbuvir/velpatasvir 400/100 mg

Non-Preferred Direct Acting Antivirals

Harvoni (ledipasvir/sofosbuvir) 90/400 mg Sovaldi (sofosbuvir) 400 mg ledipasvir/sofosbuvir 90/400 mg Vosevi (sofosbuvir, velpatasvir, and voxilaprevir) Epclusa (sofosbuvir/velpatasvir) 400/100 mg

Pediatric Indicated Direct Acting Antiretrovirals (FDA approved age ranges and

indications ONLY) (see dosing limit table)

Mavyret (glecaprevir/pibrentasvir) 400/100 mg Mavyret Oral Pellets (glecaprevir/pibrentasvir) 50/20mg Epclusa 200/50 mg tablet Harvoni (ledipasvir/sofosbuvir) 45/200 mg tablet Harvoni (ledipasvir/sofosbuvir) 90/400 mg tablet Harvoni 33.75/150 mg pellet pak Harvoni 45/200 mg pellet pak Sovaldi (sofosbuvir) 200 mg sofosbuvir/velpatasvir 400/100 mg

Preferred Regimens Listed Below (not all regimens available are listed; most <u>cost-effective</u> regimens listed below) <u>NOTE: Adult Guidelines have changed substantially; most recommendations are largely</u>

genotype non-specific; exceptions are noted in red

PLEASE CHECK REQUESTED REGIMEN

PLEASE CHECK REQUESTED REGIMEN				
ADULT:	Treatment naïve			
No cirrh	osis			
	Mavyret 100/40 mg, three (3) tablets daily for 8 weeks (for GT5/6 and/or HIV/HCV co-infection, 12 weeks is			
	recommended)			
	sofosbuvir/velpatasvir 400/100 mg, one tablet daily for 12 weeks			
Comper	nsated cirrhosis, HIV negative			
	Mavyret 100/40 mg, three (3) tablets daily for 8 weeks			
	sofosbuvir/velpatasvir 400/100, one tablet daily for 12 weeks <mark>(for GT3, add weight based RBV if Y93H positive)</mark>			
Comper	nsated cirrhosis, HIV positive			
	Mavyret 100/40 mg, three (3) tablets daily for 12 weeks			
	sofosbuvir/velpatasvir 400/100 mg, one tablet daily for 12 weeks (for GT3, add weight based RBV if Y93H positive)			
ADULT:	Treatment experienced (with or without compensated cirrhosis)			
Sofosbu	vir-based regimen			
	Mavyret 100/40 mg, three (3) tablets daily for 16 weeks			
NS3/4 p	rotease inhibitor inclusive regimen (e.g. Zepatier)			
	Vosevi 400/100/100 mg, one tablet daily for 12 weeks			
Mavyre	t			
	Vosevi 400/100/100 mg, one tablet daily for 12 weeks (if compensated cirrhosis, add weight based RBV)			
Vosevi o	or sofosbuvir + Mavyret			
	ND/OR APPROVAL OF A DRUG PRIOR AUTHORIZATION REQUEST DOES NOT GUARANTEE MEDICAID PAYMENT FOR PHARMACY PRODUCTS OR THE AMOUNT OF PAYMENT. R AND PAYMENT OF MEDICAID SERVICES ARE SUBJECT TO ALL TERMS AND CONDITIONS AND LIMITATIONS OF THE MEDICAID PROGRAM.			

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Vosevi 400/100/100 mg, one tablet daily + weight based RBV for 24 weeks				
GT 3 only: sofosbuvir/NS5A (e.g. Harvoni)				
Vosevi 400/100/100 mg, one tablet daily + weight based RBV for 12 weeks				
ADULT: Re-infection of Allograft Liver after Transplant				
DAA-treatment naïve, no decompensated cirrhosis				
Mavyret 100/40 mg, three (3) tablets daily for 12 weeks				
sofosbuvir/velpatasvir 400/100 mg, one tablet daily for 12 weeks				
DAA-treatment experienced, no decompensated cirrhosis				
Vosevi 400/100/100 mg, one tablet daily for 12 weeks				
IF multiple negative baseline characteristics, consider				
Vosevi 400/100/100 mg, one tablet daily + low dose RBV for 12 weeks				
Treatment naïve, decompensated cirrhosis				
sofosbuvir/velpatasvir 400/100 mg, one tablet daily + low dose RBV for 12 weeks				
Treatment experienced, decompensated cirrhosis (Child-Pugh B or C ONLY)				
sofosbuvir/velpatasvir 400/100 mg, one tablet daily + low dose RBV for 24 weeks				
ADULT: Decompensated Cirrhosis				
No prior sofosbuvir or NS5A failure				
sofosbuvir/velpatasvir 400/100 mg + weight-based RBV daily for 12 weeks (low dose RBV recommended for				
Child-Pugh class C cirrhosis)				
sofosbuvir/velpatasvir 400/100 mg daily for 24 weeks (will be approved only for patients with documented ineligibility for RBV)				
Prior sofosbuvir or NS5A failure				
sofosbuvir/velpatasvir 400/100 mg + weight-based RBV daily for 24 weeks (low dose RBV if Child-Pugh C)				
Other Treatment Regimen requested				
Genotype, treatment history, and extent of liver disease:				
Drug names, doses and				
durations:				
Clinical rationale for selecting regimens other than those outlined above:				

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For unique patient populations with renal impairment or HIV: please refer to the current AASLD Guidelines for recommended treatments. <u>http://www.hcvguidelines.org/full-report-view</u>

If HIV positive: Please refer to the current AASLD Guidelines for potential drug interactions and recommended dosing adjustments.

DRUG INTERACTIONS

Reference: <u>http://hep-druginteractions.org/</u> provides clinically useful, reliable, and current evidence-based information on relevant drug interactions with hepatitis medications.

CRITERIA/ADDITIONAL DOCUMENTATION HEPATITIS C FAX THIS PAGE



BENEFICIARY INFORMATION								
Beneficiary ID:		DOB:	/	/				
Beneficiary Full Name:								
Hepatitis C Therapy PA Request								
Diagnosis / Treatment Status (check all that apply) *See Hepatitis-C PA description sheet for approval criteria and intolerance definitions.								
 Prescriber is, or has consulted with a gastroenterologist, hepatologist, ID specialist or other hepatic specialist. Requires consult within the past year with documentation of recommended regimen. Active HCV infection verified by viral load within the last year: HCV RNA: million IU/mL Date: Genotype verified by lab: □ 1a □ 1b □ 2 □ 3 □ 4 □ 5 □ 6 								
HIV status: □ positive □ negative (required)	🗆 Compe	npensated cirrhosi ensated cirrhosis ver transplant		ore and Date:				
 Patient has not taken amiodarone within 535 days (required if regimen includes Harvoni or Sovaldi) 	Hepate Transpla	 Hepatocellular carcinoma and awaiting a liver transplant: Transplant date: Not yet scheduled 						
□ RBV-Ineligible reason◊:	□ CrCl _ □ Scree	sYes/No mL/min ened for HEP-B and	d HIV prior to HE	P-C treatment	start			
Hepatic fibrosis stage Last stage evaluation date:		 Date of last test: Hep B:/ HIV:/ Timing of the screening for Hep-B/HIV should be based on patient specific risk factors but lab result date must be provided and if > 1 year ago, it should be documented in the record as to why repeat testing is not clinically warranted. If the patient has had ongoing risk factors of any type, consider retesting in the month prior to HCV therapy. If positive, treatment must be considered per AASLD/IDSA and current NIH HIV guidelines. Repeat screening should be patient specific. 						
Method of cirrhosis/fibrosis stage:								
Patient is: Treatment naïve Relapser	,	Repeat bereening						
If Relapser, then prior HCV Treatment: last two regimens	s, if any							
		ates/duration of use: Res						
	Dates/duration of use: Response:							
 Prior partial responder Prior null responder Stopped prior therapy for other reason: 								
Regimen: D	Dates/duratio	n of use:	Re	sponse:				
Social History (check all that apply) □ Patient is ≥ 18 years of age OR meets current AASLD guidelines for treatment Documentation (available if requested) of: □ Counseling regarding abstinence from alcohol, IV drug use and education on how to prevent HCV transmission. □ Abstinence from drugs for at least 6 months; negative urine drug screen required if there is IV drug use history. For women of childbearing potential and male patients with female partners of childbearing potential (for RBV regimens only): □ Patient is not pregnant (or a male with a pregnant female partner) and not planning to become pregnant during treatment or within 6 months of stopping treatment. □ Agreement that partners will use 2 forms of effective contraception during treatment and for at least 6 months after stopping tx. □ Verification that monthly pregnancy tests will be performed throughout treatment.								
Other Medications (OTC, Herbal and Prescription) Inform	ation							
Drug name / strength Frequency / inst	ructions			Quantity	Refills			

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