

Clinical Policy: Glecaprevir/Pibrentasvir (Mavyret)

Reference Number: HIM.PA.SP36

Effective Date: 08.01.17 Last Review Date: 08.22 Line of Business: HIM*

Revision Log

See <u>Important Reminder</u> at the end of this policy for important regulatory and legal information.

Description

Glecaprevir and pibrentasvir (Mavyret®) are a fixed-dose combination of glecaprevir, a hepatitis C virus (HCV) NS3/4A protease inhibitor, and pibrentasvir, an HCV NS5A inhibitor.

FDA Approved Indication(s)

Mayyret is indicated for the treatment of adult and pediatric patients 3 years and older with:

- Chronic HCV genotype 1, 2, 3, 4, 5, or 6 infection without cirrhosis or with compensated cirrhosis (Child-Pugh A);
- HCV genotype 1 infection, who previously have been treated with a regimen containing an HCV NS5A inhibitor* or an NS3/4A protease inhibitor**, but not both.

Policy/Criteria

Provider must submit documentation (such as office chart notes, lab results or other clinical information) supporting that member has met all approval criteria.

It is the policy of health plans affiliated with Centene Corporation® that Mavyret is **medically necessary** when the following criteria are met:

I. Initial Approval Criteria

A. Chronic Hepatitis C Infection (must meet all):

- 1. Diagnosis of chronic HCV infection as evidenced by detectable serum HCV RNA levels by quantitative assay in the last 6 months;
- 2. Confirmed HCV genotype is one of the following (a, b, c, or d);
 - a. For treatment-naïve members: genotypes 1, 2, 3, 4, 5, or 6;
 - b. For members treatment-experienced with interferon (IFN)/pegylated-interferon (pegIFN), ribavirin (RBV), and/or sofosbuvir only: genotypes 1, 2, 3, 4, 5, or 6;
 - c. For members treatment-experienced with either an NS5A inhibitor or an NS3/4A protease inhibitor: genotype 1 (*see Appendix D*);
 - d. For Vosevi-experienced members: genotype 1, 2, 3, 4, 5, or 6;

^{*}These criteria do NOT apply to California Commercial Exchange Plans.

^{*} In clinical trials, prior NS5A inhibitor experience included ledipasvir and sofosbuvir or daclatasvir with pegylated interferon and ribavirin.

^{**} In clinical trials, prior NS3/4A protease inhibitor experience included regimens containing simeprevir and sofosbuvir, or simeprevir, boceprevir, or telaprevir with pegylated interferon and ribavirin.

^{*}Chart note documentation and copies of lab results are required



- 3. Prescribed by or in consultation with a gastroenterologist, hepatologist, infectious disease specialist, or provider who has expertise in treating HCV based on a certified training program (*see Appendix F*);
- 4. Age \geq 3 years;
- 5. If cirrhosis is present, confirmation of Child-Pugh A status;
- 6. Member is not treatment-experienced with both NS3/4A protease inhibitor AND NS5A inhibitors, such as combination therapies including Technivie[™], Viekira[™], and Zepatier[®];
- 7. Member must use **brand Epclusa**® or **Vosevi**®, unless clinically significant adverse effects are experienced or both are contraindicated*;

 *Coadministration with omeprazole up to 20 mg is not considered an acceptable medical justification
- for inability to use Epclusa
 8. Life expectancy ≥ 12 months with HCV treatment;
- 9. Member agrees to participate in a medication adherence program including both of the following components (a and b):
 - a. Medication adherence monitored by pharmacy claims data or member report;
 - b. Member's risk for non-adherence identified by adherence program or member/prescribing physician follow-up at least every 4 weeks;
- 10. Prescribed regimen is consistent with an FDA or AASLD-IDSA recommended regimen (*see Section V for reference*);
- 11. Dose does not exceed one of the following (a, b, c, or d):
 - a. Adult and pediatric members 12 years of age and older or with body weight ≥ 45 kg: glecaprevir 300 mg and pibrentasvir 120 mg (3 tablets) per day;
 - b. Pediatric members 3 years to < 12 years of age with body weight < 20 kg: glecaprevir 150 mg and pibrentasvir 60 mg per day;
 - c. Pediatric members 3 years to < 12 years of age with body weight 20 kg to < 30 kg: glecaprevir 200 mg and pibrentasvir 80 mg per day;
 - d. Pediatric members 3 years to < 12 years of age with body weight 30 kg to < 45 kg: glecaprevir 250 mg and pibrentasvir 100 mg per day.

Approval duration: up to a total of 16 weeks*

(*Approved duration should be consistent with a regimen in Section V Dosage and Administration)

B. Other diagnoses/indications

1. Refer to the off-label use policy for the relevant line of business if diagnosis is NOT specifically listed under section III (Diagnoses/Indications for which coverage is NOT authorized): HIM.PA.154 for health insurance marketplace.

II. Continued Therapy

A. Chronic Hepatitis C Infection (must meet all):

- 1. Member meets one of the following (a or b):
 - a. Currently receiving medication via Centene benefit or member has previously met initial approval criteria;
 - b. Must meet both of the following (i and ii):
 - i. Documentation supports that member is currently receiving Mavyret for chronic HCV infection and has recently completed at least 40 days of treatment with Mavyret;
 - ii. Confirmed HCV genotype is one of the following (1, 2, 3, or 4);



- 1) For treatment-naïve members: genotypes 1, 2, 3, 4, 5, or 6;
- 2) For members treatment-experienced with IFN/pegIFN, RBV, and/or sofosbuvir only: genotypes 1, 2, 3, 4, 5, or 6;
- 3) For members treatment-experienced with either an NS5A inhibitor or an NS3/4A protease inhibitor: genotype 1 (*see Appendix E*);
- 4) For Vosevi-experienced members: genotype 1, 2, 3, 4, 5, or 6;
- 2. Member is not treatment-experienced with both NS3/4A protease inhibitor AND NS5A inhibitors, such as combination therapies including Technivie, Viekira, and Zepatier;
- 3. Member is responding positively to therapy;
- 4. Dose does not exceed one of the following (a, b, c, or d):
 - a. Adult and pediatric members 12 years of age and older or with body weight ≥ 45 kg: glecaprevir 300 mg and pibrentasvir 120 mg (3 tablets) per day;
 - b. Pediatric members 3 years to < 12 years of age with body weight < 20 kg: glecaprevir 150 mg and pibrentasvir 60 mg per day;
 - c. Pediatric members 3 years to < 12 years of age with body weight 20 kg to < 30 kg: glecaprevir 200 mg and pibrentasvir 80 mg per day;
 - d. Pediatric members 3 years to < 12 years of age with body weight 30 kg to < 45 kg: glecaprevir 250 mg and pibrentasvir 100 mg per day.

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1. Refer to the off-label use policy for the relevant line of business if diagnosis is NOT specifically listed under section III (Diagnoses/Indications for which coverage is NOT authorized): HIM.PA.154 for health insurance marketplace.

III. Diagnoses/Indications for which coverage is NOT authorized:

- **A.** Non-FDA approved indications, which are not addressed in this policy, unless there is sufficient documentation of efficacy and safety according to the off label use policy HIM.PA.154 for health insurance marketplace, or evidence of coverage documents;
- **B.** Treatment-experienced members with both NS3/4A protease inhibitor AND NS5A inhibitors, such as combination therapies including: Technivie, Viekira, and Zepatier.

IV. Appendices/General Information

Appendix A: Abbreviation/Acronym Key AASLD: American Association for the

Study of Liver Diseases

FDA: Food and Drug Administration

HBV: hepatitis B virus HCV: hepatitis C virus

HIV: human immunodeficiency virus

IDSA: Infectious Diseases Society of

America

NS3/4A, NS5A/B: nonstructural protein

PegIFN: pegylated interferon

RBV: ribavirin

RNA: ribonucleic acid



Appendix B: Therapeutic Alternatives

This table provides a listing of preferred alternative therapy recommended in the approval criteria. The drugs listed here may not be a formulary agent for all relevant

lines of business and may require prior authorization.

lines of business and may require prior authorization.				
Drug Name	Dosing Regimen	Dose Limit/		
		Maximum Dose		
	Genotypes 1 through 6	sofosbuvir 400		
(Epclusa®)	Without cirrhosis or with compensated	mg/velpatasvir 100		
	cirrhosis, treatment naïve or NS3/4A	mg (1 tablet) per		
	protease inhibitor and/or pegIFN/ RBV-	day		
	experienced:			
	One tablet PO QD for 12 weeks			
sofosbuvir/velpatasvir	Genotypes 1 through 6	sofosbuvir 400 mg/		
(Epclusa [®])	Treatment-naïve and treatment-	velpatasvir 100 mg		
	experienced patients, post-liver	(1 tablet) per day		
	transplant with compensated cirrhosis			
	or without cirrhosis:			
	One tablet PO QD for 12 weeks			
Vosevi® (sofosbuvir/	Genotype 1-6	One tablet		
velpatasvir/	Treatment-experienced with NS5A	(sofosbuvir 400 mg/		
voxilaprevir)	inhibitor* with or without	velpatasvir 100 mg/		
	compensated cirrhosis:	voxilaprevir 100		
	One tablet PO QD for 12 weeks	mg) per day		
Vosevi® (sofosbuvir/	Genotype 1a or 3	One tablet		
velpatasvir/	Treatment-experienced with a	(sofosbuvir 400 mg/		
voxilaprevir)	sofosbuvir-containing regimen	velpatasvir 100 mg/		
	without NS5A inhibitor with or	voxilaprevir 100		
	without compensated cirrhosis: One	mg) per day		
	tablet PO QD for 12 weeks			
Vosevi® (sofosbuvir/	Genotype 1-6	One tablet		
velpatasvir/	Treatment-experienced with Vosevi	(sofosbuvir 400 mg/		
voxilaprevir)	with or without compensated	velpatasvir 100 mg/		
	cirrhosis:	voxilaprevir 100		
	Vosevi one tablet PO QD with	mg) per day		
	weight-based RBV for 24 weeks [†]			

Therapeutic alternatives are listed as Brand name® (generic) when the drug is available by brand name only and generic (Brand name®) when the drug is available by both brand and generic.

Treatment-experienced refers to previous treatment with NS3 protease inhibitor (telaprevir, boceprevir, or simeprevir) and/or peginterferon/RBV unless otherwise stated.

^{*} In clinical trials, prior NS5A inhibitor experience included daclatasvir, elbasvir, ledipasvir, ombitasvir, or velpatasvir

[†] Off-label, AASLD-IDSA guideline-supported dosing regimen



Appendix C: Contraindications/Boxed Warnings

- Contraindication(s):
 - o Patients with severe hepatic impairment (Child-Pugh C) or those with any history of prior hepatic decompensation.
 - o Co-administration with atazanavir or rifampin.
- Boxed warning(s): risk of hepatitis B virus (HBV) reactivation in patients coinfected with HCV and HBV.

Appendix D: Direct-Acting Antivirals for Treatment of HCV Infection

		Drug Class			
Name	NS5A Inhibitor	Nucleotide Analog NS5B Polymerase Inhibitor	Non- Nucleoside NS5B Palm Polymerase Inhibitor	NS3/4A Protease Inhibitor (PI)	CYP3A Inhibitor
Epclusa*	Velpatasvir	Sofosbuvir			
Harvoni*	Ledipasvir	Sofosbuvir			
Mavyret*	Pibrentasvir			Glecaprevir	
Sovaldi		Sofosbuvir			
Viekira Pak*	Ombitasvir		Dasabuvir	Paritaprevir	Ritonavir
Vosevi*	Velpatasvir	Sofosbuvir		Voxilaprevir	
Zepatier*	Elbasvir			Grazoprevir	

^{*}Combination drugs

Appendix E: General Information

- Acceptable medical justification for inability to use Epclusa (preferred product):
 - o In patients indicated for co-administration of Epclusa with ribavirin: contraindications to ribavirin.
- Unacceptable medical justification for inability to use Epclusa (preferred product):
 - o Coadministration with omeprazole up to 20 mg is not considered an acceptable medical justification for inability to use Epclusa.
 - Per the Epclusa Prescribing Information: "If it is considered medically necessary to coadminister, Epclusa should be administered with food and taken 4 hours before omeprazole 20 mg."
- Acceptable medical justification for inability to use Vosevi (preferred product):
 - o In patients indicated for co-administration with amiodarone: serious symptomatic bradycardia in patients taking amiodarone, with cardiac monitoring recommended.
- HBV reactivation is a Black Box Warning for all direct-acting antiviral drugs for the
 treatment of HCV. HBV reactivation has been reported when treating HCV for patients
 co-infected with HBV, leading to fulminant hepatitis, hepatic failure, and death, in some
 cases. Patients should be monitored for HBV reactivation and hepatitis flare during HCV
 treatment and post-treatment follow-up, with treatment of HBV infection as clinically
 indicated.



• Due to higher rates of virologic failure and treatment-emergent drug resistance, the data do not support labeling for treatment of HCV genotype 1 infected patients who are both NS3/4A PI and NS5A inhibitor-experienced.

• Child-Pugh Score:

	1 Point	2 Points	3 Points
Bilirubin	Less than 2 mg/dL	2-3 mg/dL	Over 3 mg/dL
	Less than 34 umol/L	34-50 umol/L	Over 50 umol/L
Albumin	Over 3.5 g/dL	2.8-3.5 g/dL	Less than 2.8 g/dL
	Over 35 g/L	28-35 g/L	Less than 28 g/L
INR	Less than 1.7	1.7 - 2.2	Over 2.2
Ascites	None	Mild / medically	Moderate-severe /
		controlled	poorly controlled
Encephalopathy	None	Mild / medically	Moderate-severe /
		controlled	poorly controlled.
		Grade I-II	Grade III-IV

Child-Pugh class is determined by the total number of points: A = 5-6 points; B = 7-9 points; C = 10-15 points

Appendix F: Healthcare Provider HCV Training

Acceptable HCV training programs and/or online courses include, but are not limited to the following:

- Hepatitis C online course (https://www.hepatitisc.uw.edu/): University of Washington is funded by the Division of Viral Hepatitis to develop a comprehensive, online self-study course for medical providers on diagnosis, monitoring, and management of hepatitis C virus infection. Free CME and CNE credit available.
- Fundamentals of Liver Disease (https://liverlearning.aasld.org/fundamentals-of-liver-disease): The AASLD, in collaboration with ECHO, the American College of Physicians (ACP), CDC, and the Department of Veterans Affairs, has developed Fundamentals of Liver Disease, a free, online CME course to improve providers' knowledge and clinical skills in hepatology.
- Clinical Care Options: http://www.clinicaloptions.com/hepatitis.aspx
- CDC training resources: https://www.cdc.gov/hepatitis/resources/professionals/trainingresources.htm

V. Dosage and Administration

Indication	Dosing Regimen	Maximum Dose	Reference
Genotypes 1-6:	Without cirrhosis or with	Adults/Peds age ≥	FDA-
Treatment-naive	compensated cirrhosis:	12 years or with	approved
	Three tablets PO QD for	body weight ≥ 45	labeling
	8 weeks	kg: glecaprevir 300	
Genotypes 1, 2, 4, 5,	Without cirrhosis:	mg/pibrentasvir 120	
or 6:	Three tablets PO QD for	mg (3 tablets) per	
Treatment-	8 weeks	day;	
experienced with			
IFN/pegIFN, RBV	With compensated	Peds age 3 years to	
and/or sofosbuvir	cirrhosis:	< 12 years of age	



Indication	Dosing Regimen	Maximum Dose	Reference
	Three tablets PO QD for	with body weight <	
	12 weeks	20 kg: glecaprevir	
Genotype 3:	Without cirrhosis or with	150 mg/pibrentasvir	
Treatment-	compensated cirrhosis:	60 mg per day;	
experienced with	Three tablets PO QD for		
IFN/pegIFN, RBV	16 weeks	Peds age 3 years to	
and/or sofosbuvir		< 12 years of age	
Genotype 1:	Without cirrhosis or with	with body weight 20	
Treatment-	compensated cirrhosis:	kg to < 30 kg:	
experienced with	Three tablets PO QD for	glecaprevir 200	
NS5A inhibitor*	16 weeks	mg/pibrentasvir 80	
without prior NS3/4A		mg per day;	
protease inhibitor [†]			
Genotype 1:	Without cirrhosis or with	Peds age 3 years to	
Treatment-	compensated cirrhosis:	< 12 years of age	
experienced with	Three tablets PO QD for	with body weight 30	
NS3/4A protease	12 weeks	kg to < 45 kg:	
inhibitor† without		glecaprevir 250	
prior NS5A inhibitor*	TI 11 PO OD C	mg/pibrentasvir 100	
Genotype 1-6:	Three tablets PO QD for	mg per day	
Treatment-naïve or	12 weeks		
treatment-	(A 16		
experienced, post-liver or kidney	(A 16-week treatment duration is recommended		
transplantation	in genotype 1-infected		
without cirrhosis or	patients who are NS5A		
with compensated	inhibitor* experienced		
cirrhosis	without prior treatment		
Cirriosis	with an NS3/4A protease		
	inhibitor [†] or in genotype		
	3-infected patients who		
	are IFN/pegIFN, RBV		
	and/or sofosbuvir		
	treatment-experienced)		
Genotypes 1-6:	With or without	Three tablets	AASLD-
Patients with prior	compensated cirrhosis:	(glecaprevir 300	IDSA
sofosbuvir/velpatasvir/		mg/pibrentasvir 120	(updated
voxilaprevir treatment	Mavyret 3 tablets PO QD	mg) per day	March 2021)
failure	+ Sovaldi 400 mg +		
	weight-based RBV for 16		
AASI D/IDSA treatment avide	weeks	ion ava undated at ivregular	

AASLD/IDSA treatment guidelines for chronic hepatitis C infection are updated at irregular intervals; refer to the most updated AASLD/IDSA guideline for most accurate treatment regimen.

^{*} In Mavyret clinical trials, subjects were treated with prior regimens containing ledipasvir and sofosbuvir or daclatasvir with (peg)interferon and RBV



† In Mavyret clinical trials, subjects were treated with prior regimens containing simeprevir and sofosbuvir, or simeprevir, boceprevir, or telaprevir with (peg)interferon and RBV.

VI. Product Availability

- Tablet: glecaprevir 100 mg and pibrentasvir 40 mg
- Oral pellet: glecaprevir 50 mg and pibrentasvir 20 mg

VII. References

- 1. Mavyret Prescribing Information. North Chicago, IL: AbbVie Inc.; June 2021. Available at: www.mavyret.com. Accessed May 5, 2022.
- 2. American Association for the Study of Liver Diseases/Infectious Disease Society of America (AASLD-IDSA). HCV guidance: recommendations for testing, managing, and treating hepatitis C. Last updated September 29, 2021. Available at: https://www.hcvguidelines.org/. Accessed May 5, 2022.
- 3. CDC. Hepatitis C Q&As for health professionals. Last updated August 7, 2020. Available at: https://www.cdc.gov/hepatitis/hcv/hcvfaq.htm. Accessed May 5, 2022.

Reviews, Revisions, and Approvals	Date	P&T
Keviews, Kevisions, and Approvais	Date	Approval
		Date
Requirement for Hep B screening was not yet approved by P & T and	09.14.17	11.19
it was therefore removed as this is under the purview of the specialist	***************************************	22129
3Q18 annual review: repeated in initial and continued approval	05.22.18	06.18
criteria the requirement against treatment-experience with both		
NS3/4A protease inhibitor AND NS5A inhibitors, as previously only		
listed in section III. diagnoses/ indications NOT allowed; expanded		
duration of tx required for COC from 30 days to 40 days; required		
verification of genotype for COC; removed requirement for advanced		
liver disease; references reviewed and updated.		
No significant change: added financial redirection to Epclusa if	07.13.18	
contraindicated to Mavyret.	10.1-10	
No significant changes: deleted an error around redirection to	10.17.18	
Epclusa.	0000000	0.7.10
2Q 2019 annual review: no significant changes; references reviewed	02.05.19	05.19
and updated.	0.7.00.10	00.10
$3Q\ 2019$ annual review: updated age ≥ 12 years or weight ≥ 45 kg to	07.02.19	08.19
be consistent with updated FDA approved indication; removed		
documented sobriety from alcohol and illicit IV drugs for ≥ 6 months		
prior to starting therapy; references reviewed and updated. Via CP.PCH.18: HIM.PA.SP36 retired and combined with HIM to	12.03.19	02.20
CP.PCH.18; added new prescriber requirement to include a "provider	12.03.19	02.20
who has expertise in treating HCV based on a certified training		
program"; Appendix F (Healthcare Provider HCV Training) added.		
RT4: updated dosing recommendations to 8 weeks total duration of		
therapy for treatment naive HCV with compensated cirrhosis across		
all genotypes (1-6).		



Reviews, Revisions, and Approvals	Date	P&T Approval Date
3Q 2020 annual review: CP.PCH.18 retired and HIM.PA.SP36	06.10.20	08.20
unretired per June SDC and prior clinical guidance; no significant		
changes; references reviewed and updated.		
3Q 2021 annual review: no significant changes; added clarification	07.12.21	08.21
that redirection to Eplcusa is for brand Epclusa in criteria; included		
reference to Appendix E with addition of contraindications that		
would warrant bypassing preferred agents; updated Appendix B		
therapeutic alternatives and section V dosing tables; updated		
reference for HIM off-label use to HIM.PA.154 (replaces		
HIM.PHAR.21); RT4: updated criteria for Mavyret pediatric age		
expansion to 3 years and older along with pediatric dosing and new		
oral pellet dosage formulation; references reviewed and updated.		
3Q 2022 annual review: no significant changes; added unacceptable	07.20.22	08.22
rationale for not using preferred Epclusa within criteria (also found		
within Appendix E); references reviewed and updated.		

Important Reminder

This clinical policy has been developed by appropriately experienced and licensed health care professionals based on a review and consideration of currently available generally accepted standards of medical practice; peer-reviewed medical literature; government agency/program approval status; evidence-based guidelines and positions of leading national health professional organizations; views of physicians practicing in relevant clinical areas affected by this clinical policy; and other available clinical information. The Health Plan makes no representations and accepts no liability with respect to the content of any external information used or relied upon in developing this clinical policy. This clinical policy is consistent with standards of medical practice current at the time that this clinical policy was approved. "Health Plan" means a health plan that has adopted this clinical policy and that is operated or administered, in whole or in part, by Centene Management Company, LLC, or any of such health plan's affiliates, as applicable.

The purpose of this clinical policy is to provide a guide to medical necessity, which is a component of the guidelines used to assist in making coverage decisions and administering benefits. It does not constitute a contract or guarantee regarding payment or results. Coverage decisions and the administration of benefits are subject to all terms, conditions, exclusions and limitations of the coverage documents (e.g., evidence of coverage, certificate of coverage, policy, contract of insurance, etc.), as well as to state and federal requirements and applicable Health Plan-level administrative policies and procedures.

This clinical policy is effective as of the date determined by the Health Plan. The date of posting may not be the effective date of this clinical policy. This clinical policy may be subject to applicable legal and regulatory requirements relating to provider notification. If there is a discrepancy between the effective date of this clinical policy and any applicable legal or regulatory requirement, the requirements of law and regulation shall govern. The Health Plan



retains the right to change, amend or withdraw this clinical policy, and additional clinical policies may be developed and adopted as needed, at any time.

This clinical policy does not constitute medical advice, medical treatment or medical care. It is not intended to dictate to providers how to practice medicine. Providers are expected to exercise professional medical judgment in providing the most appropriate care, and are solely responsible for the medical advice and treatment of members. This clinical policy is not intended to recommend treatment for members. Members should consult with their treating physician in connection with diagnosis and treatment decisions.

Providers referred to in this clinical policy are independent contractors who exercise independent judgment and over whom the Health Plan has no control or right of control. Providers are not agents or employees of the Health Plan.

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