



Rational Drug Therapy Program WVU School of Pharmacy PO Box 9511 HSCN Morgantown, WV 26506 Fax: 1-800-531-7787 Phone: 1-800-847-3859

Office of Pharmacy Services Prior Authorization Criteria for Chronic Hepatitis C Virus (HCV) Therapy

Effective 01/01/2018

Patient Consent Form
Prior Authorization Request Form
Prior Authorization Continuation Request Form
HepC Treatment Algorithm (Attachment A) and Preferred Regimens

Criteria for Approval

- All documentation must be fully completed, including the patient consent form. The viral genotype and a fibrosis score substantiated by a validated evidence-based method <u>must</u> be reported when requesting prior authorization; **AND**
- 2) Prescriber must submit laboratory evidence <u>confirming</u> that the patient is more likely than not to have a Metavir fibrosis score of F2 or greater; **AND**
- 3) Patient must meet the minimum FDA approved age requirement as specified in the package label; AND
- 4) Selected treatment regimen must be prescribed by, or in conjunction with, a board-certified gastroenterologist, hepatologist or infectious disease physician; **AND**
- 5) Patient has abstained from the use of illicit drugs and alcohol for a minimum of three (3) months, as indicated by their signature on the Patient Consent form; **AND**
- 6) Patient must agree to complete the full regimen and the patient and the provider must agree that an SVR12 will be collected and made available to WV Medicaid to verify therapy success.

Duration of Approval

- A list of accepted regimens and treatment duration for chronic Hepatitis C therapy may be found in
 <u>Attachment A</u> located at the end of this document. Initial approvals will be for a maximum of 12 weeks
 and require submission of the starting HCV RNA level.
- Additional therapy beyond 12 weeks may be requested by completing the <u>Prior Authorization</u>
 <u>Continuation Request Form</u> and is approvable only after receipt of a viral load indicating treatment efficacy as suggested by AASLD guidelines*.
- Emergency fills will NOT be granted under any circumstance.
- * AASLD guidelines recommend that quantitative HCV viral load testing be done after 4 weeks of therapy (TW4). If HCV RNA is detectable at week 4 of treatment, repeat quantitative HCV RNA viral load testing is recommended after 2 additional weeks of treatment. If the quantitative HCV viral load has <u>increased</u> by greater than 10-fold (>1 log₁₀ IU/mL) on repeat testing at week 6 (or thereafter), then discontinuation of HCV treatment is recommended.





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ADDITIONAL CONSIDERATIONS

- 1) It is highly recommended that the patient be vaccinated against Hepatitis A and Hepatitis B.
- 2) Cirrhosis should be substantiated either through biopsy or the presence of **at least two** of the following clinical features:
 - a. Cirrhotic features on imaging (MRI, ultrasound, or CT)
 - b. Ascites
 - c. Esophageal varices
 - d. Reversed AST:ALT ratio (> 1), thrombocytopenia (< 130,000 platelets/μL), and coagulopathy (INR > 2)
- 3) For HCV/HIV co-infections all requests must be reviewed for drug-drug interactions prior to approval. Please submit a list of the patient's current HIV regimen along with your request for coverage of the selected HCV regimen.

PRIOR AUTHORIZATION MAY BE DENIED FOR THE FOLLOWING REASONS

- 1) Failure to report a genotype, fibrosis score or other significant omission from required documentation.
- 2) Any request falling outside the manufacturer guidelines for safe use.
- 3) Evidence exists that the patient has abused any illicit substance or alcohol in the past three (3) months.
- 4) Patient is taking a concomitant medication that has significant clinical interactions with the requested regimen.
- 5) Requests for continuation of coverage beyond 12 weeks will be denied if the patient's HCV RNA level has increased by greater than 10-fold (>1 log₁₀ IU/mL) on repeat testing at week 6 (or thereafter) or if the prescriber has not submitted or has not obtained a viral load prior to treatment week 12. **Denial of continuation due to lack of efficacy does not prevent the approval of an alternative regimen if indicated by AASLD guidelines.**
- 6) Coverage shall be for one <u>successful</u> course of therapy in a lifetime. Success of therapy shall be judged by undetectable quantitative HCV RNA levels measured at 12 weeks following completion of therapy (SVR12). If RNA levels have not been submitted, then it will be assumed that therapy was successful. **Re-infection will not be covered.** Exceptions may be allowed on a case-by-case basis.
- 7) Lost or stolen medication replacement request will not be authorized.





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ATTACHMENT A: HepC Treatment Algorithm and Preferred Regimens

(not all regimens available are listed; most cost-effective regimens listed below)

Ge	notype 1a			
	Treatment naïve, no cirrhosis, HCV viral load < 6 million copies/ml → Regimen 1 or 12 (HIV negative only) or 7 (only if			
	negative for NS5A resistance associated polymorphisms) or 5			
	Treatment naïve, no cirrhosis, HCV viral load \geq 6 million \Rightarrow Regimen 1 or 7 (only if negative for NS5A resistance associated			
	polymorphisms¥) or 5			
Ц	Treatment naïve, compensated cirrhosis, Child-Pugh A ONLY → Regimen 2 or 7 (only if negative for NS5A resistance associated polymorphisms¥) or 5			
	Treatment experienced (PEG-IFN + ribavirin ONLY), not cirrhotic → Regimen 1 or 7 (only if negative for NS5A resistance associated polymorphisms) or 5			
	Treatment experienced (PEG-IFN + ribavirin ONLY), compensated cirrhosis (Child-Pugh A ONLY) \rightarrow Regimen 7 (only if negative for NS5A resistance associated polymorphisms¥) or 5 or 2			
	Treatment experienced (PEG-IFN + ribavirin +NS3 protease inhibitor, no prior NS5A, no sofosbuvir), no cirrhosis → Regimen 9 (only if negative for NS5A resistance associated polymorphisms¥) or 5 or 2			
	Treatment experienced (PEG-IFN + ribavirin + protease inhibitor, no prior NS5A, no sofosbuvir), compensated cirrhosis, Child-Pugh A ONLY → Regimen 9 (only if negative for NS5A resistance associated polymorphisms¥) or 5 or 2			
	Treatment experienced (sofosbuvir + ribavirin +/- PEG-IFN OR simeprevir, no NS5A), no cirrhosis → Regimen 2			
	Treatment experienced (sofosbuvir + ribavirin +/- PEG-IFN OR simeprevir, no NS5A), compensated cirrhosis, Child-Pugh A ONLY → Regimen 2			
	Treatment experienced, any NS5A inhibitor but NO NS3/4A protease inhibitor (prior therapy ONLY with			
	daclatasvir+sofosbuvir, ledipasvir+sofosbuvir or sofosbuvir +velpatasvir), no cirrhosis or compensated cirrhosis, Child-			
	Pugh A ONLY → Regimen 3 or 10			
Ц	Treatment experienced, any NS5A inhibitor (ledipasvir (Harvoni), velpatasvir (Epclusa/Vosevi), elbasvir (Zepatier),			
	dasabuvir (Viekira), pibrentasvir (Mavyret) and daclatasvir (Daklinza), including those given with a NS3/4A protease			
	inhibitor, non-cirrhotic or compensated cirrhosis (Child-Pugh A ONLY) > Regimen 10			
	Re-infection of allograft liver after transplant, no cirrhosis → Regimen 2			
<u> </u>	Re-infection of allograft liver after transplant, compensated cirrhosis (Child Pugh A ONLY) Regimen 13			
	Re-infection of allograft liver after transplant, decompensated cirrhosis (Child-Pugh B and C only) Regimen 14 Resigned as a prior soft of the pugh soft of			
	Decompensated cirrhosis, no prior sofosbuvir or NSSA → Regimen 6 (low dose ribavirin if Child-Pugh Class C)			
	Decompensated cirrhosis, no prior sofosbuvir or NS5A, ribavirin ineligible**→ Regimen 4			
	Decompensated cirrhosis, prior treatment with sofosbuvir or NS5A → Regimen 6 (low dose ribavirin if Child-Pugh Class C)			
	notype 1b			
	Treatment naïve, no cirrhosis, HCV viral load <6 million copies/ml → Regimen 1 or12 (HIV negative only) or 7 or 5			
	Treatment naïve, no cirrhosis, HCV viral load ≥6 million → Regimen 1 or 7 or 5			
<u> </u>	Treatment naïve, compensated cirrhosis, Child-Pugh A ONLY → Regimen 2 or 7 or 5			
	Treatment experienced (PEG-IFN + ribavirin ONLY), not cirrhotic → Regimen 1 or 7 or 5			
	Treatment experienced (PEG-IFN + ribavirin ONLY), compensated cirrhosis (Child-Pugh A ONLY) → Regimen 7 or 5 or 2			
	Treatment experienced (PEG-IFN + ribavirin + protease inhibitor), no prior NS5A, no prior sofosbuvir, no cirrhosis ->			
	Regimen 9 or 5 or 2			
Ц	Treatment experienced (PEG-IFN + ribavirin + protease inhibitor), no prior NS5A, no prior sofosbuvir, compensated cirrhosis, Child-Pugh A ONLY → Regimen 9 or 5 or 2			
	Treatment experienced (sofosbuvir + ribavirin +/- PEG-IFN OR simeprevir, no NS5A), no cirrhosis → Regimen 5 or 2			
	Treatment experienced (sofosbuvir + ribavirin +/- PEG-IFN OR simeprevir, no NS5A), compensated cirrhosis, Child-Pugh A			





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	ONLY → Regimen 5 or 2
	Treatment experienced, any NS5A inhibitor but NO NS3/4A protease inhibitor (prior therapy ONLY with
	daclatasvir+sofosbuvir, ledipasvir+sofosbuvir or sofosbuvir +velpatasvir), no cirrhosis or compensated cirrhosis, Child-
_	Pugh A ONLY → Regimen 3 or 10
Ц	Treatment experienced, any NS5A inhibitor (ledipasvir (Harvoni), velpatasvir (Epclusa/Vosevi), elbasvir (Zepatier), dasabuvir (Viekira), pibrentasvir (Mavyret) and daclatasvir (Daklinza), including those given with a NS3/4A protease
	inhibitor, non-cirrhotic or compensated cirrhosis (Child-Pugh A ONLY) \rightarrow Regimen 10
	Re-infection of allograft liver after transplant, no cirrhosis → Regimen 2
<u> </u>	Re-infection of allograft liver after transplant, roo envisors > Negamen 2 Re-infection of allograft liver after transplant, compensated cirrhosis (Child-Pugh A ONLY) → Regimen 13
<u> </u>	Re-infection of allograft liver after transplant, decompensated cirrhosis (Child-Pugh B and C only) \rightarrow Regimen 14
<u> </u>	Decompensated cirrhosis, no prior sofosbuvir or NSSA → Regimen 6 (low dose ribavirin if Child-Pugh Class C)
<u> </u>	Decompensated cirrhosis, no prior sofosbuvir or NSSA, ribavirin ineligible**→ Regimen 4
<u> </u>	Decompensated cirrhosis, prior treatment with sofosbuvir or NS5A → Regimen 6 (low dose ribavirin if Child-Pugh Class C)
	notype 2
	Treatment naïve, no cirrhosis → Regimen 1 or 5
<u> </u>	Treatment naïve, compensated cirrhosis, Child-Pugh A ONLY → Regimen 5 or 2
<u> </u>	Treatment experienced (PEG-IFN + ribavirin), no cirrhosis → Regimen 1 or 5
<u> </u>	Treatment experienced (PEG-IFN + ribavirin), no cirriosis → Regimen 1 or 3 Treatment experienced (PEG-IFN + ribavirin), compensated cirrhosis (Child-Pugh A ONLY) → Regimen 5 or 2
<u> </u>	Treatment experienced (sofosbuvir + ribavirin) → Regimen 5 or 2
	· · · · · · · · · · · · · · · · · · ·
	Decompensated cirrhosis, no prior sofosbuvir or NS5A failure → Regimen 6 or if RBV ineligible**ONLY→ Regimen 4
	Decompensated cirrhosis, prior sofosbuvir or NS5A failure → Regimen 16 (low dose ribavirin if Child-Pugh C)
	Re-infection of allograft liver after transplant, no cirrhosis → Regimen 2
	Re-infection of allograft liver after transplant, compensated cirrhosis, → Regimen 15 or 6 or 2
	Re-infection of allograft liver after transplant, decompensated cirrhosis → Regimen 15 or 6
	notype 3
<u></u>	Treatment naïve, no cirrhosis → Regimen 1 or 5
	Treatment naïve, with cirrhosis, Child-Pugh A ONLY → Regimen 5 or 2
<u> </u>	Treatment experienced (PEG-IFN + ribavirin), no cirrhosis, Y93H neg → Regimen 5 or 3
<u> </u>	Treatment experienced (PEG-IFN + ribavirin), no cirrhosis, Y93H positive → Regimen 6 or 3
	Treatment experienced (PEG-IFN + ribavirin), compensated cirrhosis, Child-Pugh A ONLY \rightarrow Regimen 6 or 3, if RBV ineligible only** \rightarrow Regimen 8
	Treatment experienced (any direct acting antiviral including NS5A), no or compensated cirrhosis, Child-Pugh A ONLY → Regimen 10; if prior NS5A AND cirrhosis → Regimen 11
	Decompensated cirrhosis, no prior sofosbuvir or NS5A failure \rightarrow Regimen 6 or, if RBV ineligible ONLY** ** \rightarrow Regimen 4
	Decompensated cirrhosis, prior sofosbuvir or NS5A failure → Regimen 16 (low dose ribavirin if Child-Pugh C)
	Re-infection of allograft liver after transplant, no cirrhosis → Regimen 2
	Re-infection of allograft liver after transplant, compensated cirrhosis → Regimen 15 or 6 or 2
	Re-infection of allograft liver after transplant, decompensated cirrhosis → Regimen 15 or 6
Ge	notype 4
	Treatment naïve, no cirrhosis → Regimen 1 or 7 or 5
	Treatment naïve, compensated cirrhosis (Child-Pugh A ONLY) → Regimen 7 or 5 or 2
	Treatment experienced (PEG-IFN + ribavirin), no cirrhosis → Regimen 1 or 7 (only if prior virologic relapse after PEG-IFN therapy) or 5
	Treatment experienced (PEG-IFN + ribavirin), compensated cirrhosis, Child-Pugh A ONLY→ Regimen 5 or 7 (only if prior virologic relapse after PEG-IFN therapy) or Regimen 2
	Treatment experienced (any direct acting antiviral including NS5A), with or without compensated cirrhosis (Child-Pugh A





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	$ONLY) \rightarrow \frac{Regimen}{10}$		
Ţ	Decompensated cirrhosis, no prior sofosbuvir or NS5A \rightarrow Regimen 6 (low dose ribavirin if Child-Pugh Class C)		
Ū	☐ Decompensated cirrhosis, no prior sofosbuvir or NS5A, ribavirin ineligible** → Regimen 4		
Ţ	Decompensated cirrhosis, prior treatment with sofosbuvir or NS5A \rightarrow Regimen 6 (low dose ribavirin if Child-Pugh Class C)		
Ţ	Re-infection of allograft liver after transplant, no cirrhosis → Regimen 2		
Ţ	Re-infection of allograft liver after transplant, compensated cirrhosis (Child-Pugh A ONLY) → Regimen 13		
Ţ	Re-infection of allograft liver after transplant, decompensated cirrhosis (Child-Pugh B and C only) → Regimen 14		
	notype 5		
Ţ	☐ Treatment naive, no cirrhosis → Regimen 1 or 5		
Ţ	☐ Treatment naïve, compensated cirrhosis, Child-Pugh A ONLY → Regimen 5 or 2		
Ţ	Treatment experienced (PEG-IFN + ribavirin), without cirrhosis \rightarrow Regimen 1 or 5		
Ţ	Treatment experienced (PEG-IFN + ribavirin), compensated cirrhosis (Child-Pugh A ONLY) \rightarrow Regimen 5 or 2		
Ţ	☐ Treatment experienced (any DAA including NS5A), with no or compensated cirrhosis (Child-Pugh A ONLY) → Regimen 10		
Ţ	Decompensated cirrhosis, no prior sofosbuvir or NS5A \rightarrow Regimen 6 (low dose ribavirin if Child-Pugh Class C)		
Ţ	☐ Decompensated cirrhosis, no prior sofosbuvir or NS5A, ribavirin ineligible** → Regimen 4		
Ţ	Decompensated cirrhosis, prior treatment with sofosbuvir or NS5A \rightarrow Regimen 6 (low dose ribavirin if Child-Pugh Class C)		
Ţ	Re-infection of allograft liver after transplant, no cirrhosis $\rightarrow \frac{\text{Regimen}}{2}$		
Ţ	Re-infection of allograft liver after transplant, compensated cirrhosis (Child-Pugh A ONLY) \rightarrow Regimen 13		
Ţ	Re-infection of allograft liver after transplant, decompensated cirrhosis (Child-Pugh B and C only) \rightarrow Regimen 14		
	Genotype 6		
Ţ	☐ Treatment naïve, no cirrhosis → Regimen 1 or 5		
Ţ	☐ Treatment naïve, compensated cirrhosis (Child-Pugh A ONLY) → Regimen 5 or 2		
Ţ	Treatment experienced (PEG-IFN + ribavirin), without cirrhosis \rightarrow Regimen 1 or 5		
Ţ	Treatment experienced (PEG-IFN + ribavirin), compensated cirrhosis (Child-Pugh A ONLY) \rightarrow Regimen 5 or 2		
Ţ	Treatment experienced (any direct acting antiviral, including NS5A) with or without compensated cirrhosis (Child-Pugh A		
_	ONLY) → Regimen 10		
	Decompensated cirrhosis, no prior sofosbuvir or NS5A→Regimen 6 (low dose ribavirin if Child-Pugh Class C)		
	Decompensated cirrhosis, no prior sofosbuvir or NS5A, ribavirin ineligible** → Regimen 4		
_	Decompensated cirrhosis, prior treatment with sofosbuvir or NS5A → Regimen 6 (low dose ribavirin if Child-Pugh Class C)		
_	Re-infection of allograft liver after transplant, no cirrhosis Regimen 2		
_	Re-infection of allograft liver after transplant, compensated cirrhosis (Child-Pugh A ONLY) → Regimen 13		
	Re-infection of allograft liver after transplant, decompensated cirrhosis (Child-Pugh B and C only) \rightarrow Regimen 14		





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<u>Preferred REGIMENS Key for HepC Treatment Algorithm (Attachment A)</u>:

т.	iviavyret (giecapre	evir/pibrentasvir) 100/40 mg; three (3) tablets daily for 56 days (8 weeks) 🗀							
2.	Mavyret (glecaprevir/pibrentasvir) 100/40 mg; three (3) tablets daily for 84 days (12 weeks) □								
3.	Mavyret (glecaprevir/pibrentasvir) 100/40 mg; three (3) tablets daily for 112 days (16 weeks) □								
4.	Epclusa (sofosbuvir/velpatasvir) 400/100 mg daily for 168 days (24 weeks) □								
5.	Epclusa (sofosbuv	Epclusa (sofosbuvir/velpatasvir) 400/100 mg daily for 84 days (12 weeks)							
6.	Epclusa (sofosbuvir/velpatasvir) 400/100 mg daily + weight-based ribavirin for 84 days (12 weeks)								
7.	Zepatier (elbasvir/grazoprevir) 50/100 mg daily for 84 days (12 weeks) □								
8.	Zepatier (elbasvir/grazoprevir) 50/100 mg daily + sofosbuvir 400 mg daily for 84 days (12 weeks) □								
9.	Zepatier (elbasvir/grazoprevir) 50/100 mg daily + weight based ribavirin for 84 days (12 weeks) □								
10.	Vosevi (sofosbuvir/velpatasvir/voxilaprevir) 400/100/100 mg, one tablet daily for 84 days (12 weeks) □								
	Vosevi (sofosbuvir/velpatasvir/voxilaprevir) 400/100/100 mg, one tablet daily + weight-based ribavirin for 84 days (12 weeks) □								
12.	Harvoni (ledipasvi	ir/sofosbuvir) 90/400 mg daily for 56 days (8 weeks) 🗖							
13.	Harvoni (ledipasvi	Harvoni (ledipasvir/sofosbuvir) 90/400 mg daily + weight-based ribavirin for 84 days (12 weeks) □							
14.	Harvoni (ledipasvir/sofosbuvir) 90/400 mg daily + low dose ribavirin for 84 days (12 weeks) □								
15.	Daklinza^(daclata	svir) 60 mg plus Sovaldi (sofosbuvir) 400 mg daily + low initial dose of ribavirin for 84 days (12 weeks)							
16.	Epclusa (sofosbuv	vir/velpatasvir) 400/100 mg daily + weight based ribavirin for 168 days (24 weeks) □							
	concurrent CYP3A # low dose ribavir	a (daclatasvir) MUST BE ADJUSTED with certain co-administered drugs (reduced to 30 mg daily with A4 inhibitors and increased to 90 mg daily with concurrent moderate CYP3A4 inducers) rin = 600 mg/day and increase as tolerated slymorphisms at amino acid positions 28, 30, 31, or 93							
	•	inical rationale with the completed PA form if choosing a regimen that is beyond those found elines, or if selecting regimens other than those outlined above.							
WICHIII	ine current guide	miles, of it selecting regimens other than those outlined above.							
	**Pationts wh	o are ribavirin-ineligible must have at least one of the following reasons documented:							
_		History of severe or unstable cardiac disease							
		Pregnant women and men with pregnant partners							
		Diagnosis of hemoglobinopathy (e.g., thalassemia major, sickle cell anemia)							
		Hypersensitivity to ribavirin							
		Baseline platelet count <70,000 cells/mm3							
1		ANC <1500 cells/mm3							
	П	Hh <12 gm/dl in women or <13 g/dl in men							

Patients with CrCl <50 ml/min (moderate or severe renal dysfunction, ESRD, HD) should have dosage reduced





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General Mechanism of Action for Available Agents

- **Daklinza** (daclatasvir) is an HCV NS5A inhibitor.
- **Epclusa** (sofosbuvir/velpatasvir) is a fixed-dose combination of sofosbuvir, a hepatitis C virus (HCV) nucleotide analog NS5B polymerase inhibitor and velpatasvir, an HCV NS5A inhibitor.
- Harvoni (ledipasvir/sofosbuvir) is a fixed-dose combination of ledipasvir, an HCV NS5A inhibitor, and sofosbuvir, an HCV nucleotide analog NS5B polymerase inhibitor.
- Mavyret (glecaprevir/pibrentasvir) is a fixed-dose combination of glecaprevir, an HCV NS3/4A protease inhibitor, and pibrentasvir, an HCV NS5A inhibitor, and is indicated for the treatment of patients with chronic HCV genotype (GT) 1, 2, 3, 4, 5 or 6 infection without cirrhosis and with compensated cirrhosis (Child-Pugh A). Mavyret is also indicated for the treatment of adult patients with 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. (NOTE: GT1 is the only genotype that can be retreated with Mavyret after previous NS5A or NS3/4A protease inhibitor therapy)
- Olysio (simeprevir) is an HCV NS3/4A protease inhibitor.
- Sovaldi (sofosbuvir) is an HCV nucleotide analog NS5B polymerase inhibitor.
- **Technivie** (ombitasvir/paritaprevir) is a fixed-dose combination of ombitasvir, an HCV NS5A inhibitor, paritaprevir, an HCV NS3/4A protease inhibitor, and ritonavir, a CYP3A inhibitor.
- **Viekira XR** (dasabuvir/ombitasvir/paritaprevir) includes dasabuvir, an HCV non-nucleoside NS5B palm polymerase inhibitor, ombitasvir, an HCV NS5A inhibitor, paritaprevir, an HCV NS3/4A protease inhibitor, and ritonavir, a CYP3A inhibitor.
- Vosevi (sofosbuvir/velpatasvir/voxilaprevir) is a fixed-dose combination of sofosbuvir, a hepatitis C virus (HCV) nucleotide analog NS5B polymerase inhibitor, velpatasvir, an HCV NS5A inhibitor, and voxilaprevir, an HCV NS3/4A protease inhibitor, and is indicated for the treatment of adult patients with chronic HCV infection without cirrhosis or with compensated cirrhosis (Child-Pugh A) who have:
 - genotype 1, 2, 3, 4, 5, or 6 infection and have previously been treated with an HCV regimen containing an NS5A inhibitor.
 - genotype 1a or 3 infection and have previously been treated with an HCV regimen containing sofosbuvir without an NS5A inhibitor.
 - *** Additional benefit of VOSEVI over sofosbuvir/velpatasvir was not shown in adults with genotype 1b, 2, 4, 5, or 6 infection previously treated with sofosbuvir without an NS5A inhibitor.
- **Zepatier** (elbasvir/grazoprevir) is a fixed-dose combination product containing elbasvir, an HCV NS5A inhibitor, and grazoprevir, an HCV NS3/4A protease inhibitor.





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References

- American Association for the Study of Liver Diseases Infectious Diseases Society of America: Recommendations for testing, managing and treating hepatitis C. Available at: http://hcvguidelines.org/ Accessed September 05, 2017.
- 2) LexiComp Clinical Drug Information Accessed November 22, 2016.
- 3) Epclusa [package insert]. Foster City, CA; Gilead, June 2016.
- Viekira XR[™] [package insert]. Abbvie, Revised 7/2016
- 5) Daklinza [package insert]. Bristol-Myers Squibb Company, Feb 2016.
- 6) Sovaldi [package insert]. Foster City, CA; Gilead, August 2015.
- 7) Olysio [package insert]. Janssen Therapeutics; Titusville, NJ. April 2015.
- 8) Technivie® [package insert]. Abbvie, Revised 7/2015
- 9) Viekira Pak™ [package insert]. Abbvie, Revised 4/2016
- 10) Zepatier [package insert]. Merck, January, 2016.
- 11) Harvoni [package insert]. Foster City, CA; Gilead, February 2016.
- 12) Poynard T, Ratziu V, Benmanov Y, DiMartino V, Bedossa P, Opolon P. Fibrosis in patients with hepatitis c: detection and significance. *Semin Liver Dis.* 2000;20(1). Retrieved from www.medscape.com. Accessed February 26, 2014.
- 13) Heidelbaugh JJ and Bruderly M. Cirrhosis and Chronic Liver Failure: Part I. Diagnosis and Evaluation. *Am Fam Physician*. 2006 Sep 1;74(5):756-762.
- 14) Mavyret [package insert]. Abbvie. August, 2017.

Attachment A Change Log:

Ver 2016.3C Created by Laureen Biczak (GHS) and edited by BMT 6/7/2016

Ver 2016.4D Created by Laureen Biczak (CHC)

Ver 2016.4E Created by Laureen Biczak (CHC)

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