



**FOR PATIENTS WHO DO NOT MEET CRITERIA FOR SIMPLIFIED TREATMENT, SEE BELOW****Pre-treatment Assessment/On-Treatment Monitoring and Follow-up Recommendations Available at:**<https://www.hcvguidelines.org/treatment-naive/simplified-treatment>Providers are urged to check an online drug interaction site such as: <https://www.hep-druginteractions.org/checker>**Please attach documentation of the following:**

<input type="checkbox"/> Quantitative HCV RNA viral load (done within 6 months or consistently positive in past results for $\geq 1$ year span of time with the last being in the last 5 years) <input type="checkbox"/> Child-Turcotte-Pugh (CTP) Score: _____ Date: _____ <input type="checkbox"/> Patient does not have limited life expectancy (less than 12 months) due to non-liver-related comorbid conditions. <input type="checkbox"/> Within 60 days of request in women of childbearing age: pregnancy test*	<input type="checkbox"/> HCV Genotype verified by lab * Genotype: (circle) 1a 1b 2 3 4 5 6 <b>Labs below done within the last 6 months</b> <input type="checkbox"/> Fibrosis score*: _____ Date: _____ method: _____ <input type="checkbox"/> CBC* <input type="checkbox"/> Hepatic function panel*: albumin, total and direct bilirubin, ALT, AST <input type="checkbox"/> Calculated glomerular filtration rate: eGFR* _____ <input type="checkbox"/> Quantitative HCV RNA viral load* _____ <input type="checkbox"/> HIV antigen/antibody test* _____ <input type="checkbox"/> Hepatitis B surface antigen* _____
<input type="checkbox"/> Prescriber is, or has consulted with, a gastroenterologist, hepatologist, ID specialist or other Hepatitis specialist. Consult must be within the past year with documentation of recommended regimen.*	<input type="checkbox"/> Provider certifies they have checked an up-to-date drug interaction list or on-line list such as: <a href="https://www.hep-druginteractions.org/checker">https://www.hep-druginteractions.org/checker</a> .

**PEDIATRIC NOTE: FDA approved pediatric formulations of direct acting antivirals (DAA) and DAA approved for pediatric use will be approved for those under the age of eighteen when used in accordance with current AASLD guidelines including for indication and age. Prior authorization is still required prior to the first dose and for treatment naive children when used in accordance with the table below.**

GT	Age (years)	Weight (kg)	Drug/Dose	Weeks
Any	$\geq 3$ to 11	<20	Mavyret 50/20mg Pellet Pack (3 packets once daily)	8
		$\geq 20$ to <30	Mavyret 50/20mg Pellet Pack (4 packets once daily)	8
		$\geq 30$ to <45	Mavyret 50/20mg Pellet Pack (5 packets once daily) -OR- sofosbuvir/velpatasvir 400/100 mg tablet	8 12
Any	$\geq 12$	$\geq 45$	Mavyret 100/40 mg tablets (3 tablets once daily)-OR-	8
			sofosbuvir/velpatasvir 400/100 mg tablet	12

**NOTE: Adult Guidelines have changed substantially; most recommendations are largely genotype non-specific; exceptions are noted in red**

**ADULT: Treatment naïve (includes those treated in the past with IFN/RBV or IFN + 1<sup>st</sup> generation protease inhibitors)****No cirrhosis**

- Mavyret 100/40 mg, three (3) tablets daily for 8 weeks
- sofosbuvir/velpatasvir 400/100 mg, one tablet daily for 12 weeks

**Compensated cirrhosis**

- Mavyret 100/40 mg, three (3) tablets daily for 8 weeks (GT4 AND HCV coinfecting patients with compensated cirrhosis, IDSA/AASLD guidelines recommend 12 weeks of treatment)
- sofosbuvir/velpatasvir 400/100, one tablet daily for 12 weeks (for GT3, add weight based RBV if Y93H positive)

**ADULT: Treatment experienced (with or without compensated cirrhosis)****Sofosbuvir-based regimen**

- Mavyret 100/40 mg, three (3) tablets daily for 16 weeks

**NS3/4 protease inhibitor inclusive regimen (e.g. Zepatier)**

- Vosevi 400/100/100 mg, one tablet daily for 12 weeks

**Mavyret**

- Vosevi 400/100/100 mg, one tablet daily for 12 weeks (if compensated cirrhosis, add weight based RBV)

**Vosevi or sofosbuvir + Mavyret**

- 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****Genotype, treatment history, and extent of liver disease:**


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**Drug names, doses and durations:**


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**Clinical rationale for selecting regimens other than those outlined above:**


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**Abbreviations RBV=ribavirin; PI=protease inhibitor; DAA=direct acting antiviral**

**# low dose ribavirin = 600 mg/day and increase as tolerated**

**For ANY regimen that includes ribavirin**

- For women of childbearing potential** (and male patients with female partners of childbearing potential):
  - 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

- Agreement that partners will use two forms of effective contraception during treatment and for at least 6 months after stopping
- Verification that monthly pregnancy tests will be performed throughout treatment

**For ribavirin-ineligible\*\*:** (Patients with CrCl <50 ml/min (moderate or severe renal dysfunction, ESRD, HD) should have dosage reduced

- 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/mm<sup>3</sup>
- ANC <1500 cells/mm<sup>3</sup>
- Hb <12 gm/dl in women or <13 g/dl in men
- Other: \_\_\_\_\_

By completing this form, I hereby certify that the above request is true, accurate and complete. That the request is medically necessary, does not exceed the medical needs of the member, and is clinically supported in your medical records. I also understand that any misrepresentations or concealment of any information requested in the prior authorization request may subject me to audit and recoupment.

**Provider Signature:** \_\_\_\_\_ **Date of Submission:** \_\_\_\_\_

**\*MUST MATCH PROVIDER LISTED ABOVE**