**Direct Acting Antivirals for Hepatitis C (HCV)**

**Genotype 4**

Harvoni® (ledipasvir 90mg & sofosbuvir 400mg)
Olysio® (simeprevir 150mg)
Sovaldi® (sofosbuvir 400mg)
Technivie® (ombitasvir 12.5mg, paritaprevir 75mg, & ritonavir 50mg)
Zepatier® (elbasvir 50mg & grazoprevir 100mg)

**Indications:**

“Harvoni is a fixed-dose combination of ledipasvir, a hepatitis C virus (HCV) NS5A inhibitor, and sofosbuvir, an HCV nucleotide analog NS5B polymerase inhibitor, and is indicated with or without ribavirin for the treatment of chronic hepatitis C virus (HCV) genotype 1, 4, 5, or 6 infection.”

“Olysio is a hepatitis C virus (HCV) NS3/4A protease inhibitor indicated for the treatment of chronic hepatitis C (CHC) genotype 1 or 4 infection as a component of a combination antiviral treatment regimen.”

“Sovaldi is a hepatitis C virus (HCV) nucleotide analog NS5B polymerase inhibitor indicated for the treatment of genotype 1, 2, 3 or 4 chronic hepatitis C virus (HCV) infection as a component of a combination antiviral treatment regimen.”

“Technivie is a fixed-dose combination of ombitasvir, a hepatitis C virus NS5A inhibitor, paritaprevir, a hepatitis C virus NS3/4A protease inhibitor, and ritonavir, a CYP3A inhibitor and is indicated in combination with ribavirin for the treatment of patients with genotype 4 chronic hepatitis C virus (HCV) infection without cirrhosis.”

“Zepatier is a fixed-dose combination product containing elbasvir, a hepatitis C virus (HCV) NS5A inhibitor, and is indicated with or without ribavirin for treatment of chronic HCV genotypes 1 or 4 infection in adults.”

**Table 1: FDA Labeled Indications**

<table>
<thead>
<tr>
<th></th>
<th>Daklinza + Sovaldi</th>
<th>Harvoni</th>
<th>Olysio + Sovaldi</th>
<th>Sovaldi</th>
<th>Technivie</th>
<th>VieKira</th>
<th>Zepatier</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Genotype 1a</strong></td>
<td>Yes</td>
<td>Yes</td>
<td>Yes</td>
<td>Yes</td>
<td></td>
<td>Yes</td>
<td>Yes</td>
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<tr>
<td><strong>Genotype 1b</strong></td>
<td>Yes</td>
<td>Yes</td>
<td>Yes</td>
<td>Yes</td>
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<tr>
<td><strong>Genotype 2</strong></td>
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<td>Yes</td>
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<tr>
<td><strong>Genotype 3</strong></td>
<td></td>
<td></td>
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<td>Yes</td>
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<tr>
<td><strong>Genotype 4</strong></td>
<td></td>
<td></td>
<td></td>
<td>Yes</td>
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<td>Yes</td>
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<tr>
<td><strong>Genotype 5</strong></td>
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<td>Yes</td>
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<tr>
<td><strong>Genotype 6</strong></td>
<td></td>
<td></td>
<td></td>
<td>Yes</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
**Quantity Limit**

- Harvoni – One tablet once per day (28 tablets /28 days)
- Sovaldi – One tablet per day (28 tablets/28 days)
- Technivie – Two tablets once per day with food (56 tablets/28 days)
- Zepatier – One tablet once per day (28 tablets /28 days)

**Additional Considerations**

Ongoing patient engagement is encouraged throughout the treatment course for optimal outcomes.

Combination treatment with ribavirin is contraindicated in women who are pregnant or may become pregnant and men whose female partners are pregnant because of the risks for birth defects and fetal death associated with ribavirin.

**Criteria for Approval: Treatment Naïve**

1. Adult patient age ≥ 18 years old; **AND**
2. Documentation of HCV genotype, subtype, and HCV viral load is included in the authorization request; **AND**
3. Meets diagnosis and disease severity of Hepatitis C, Genotype (GT) 4 and Metavir Fibrosis score F2-F4 or equivalent (includes extrahepatic manifestations of advancing disease); **AND**
4. To confirm the Metavir fibrosis stage, at least one of the following tests or procedures must be submitted: biopsy, elastography, FibroSure, FibroTest, HepaScore, or FibroScan *(Note: APRI and FIB-4 will not be accepted as confirmation of the Metavir fibrosis stage)*; **AND**
5. The Hepatitis C disease activity score must be submitted with the authorization request; **AND**
6. Documentation of previously trialed HCV therapies, dates of therapy, whether full therapy was completed or discontinued early, and if discontinued early, the reason for the discontinuation is included in the authorization request; **AND**
7. The patient agrees to complete regimen; **AND**
8. The prescriber agrees to maintain HCV RNA levels obtained at 12-weeks and 24-weeks post-therapy completion to demonstrate Sustained Virologic Response (SVR); **AND**
9. Patient has been tested for the use of illicit drugs, controlled substances, and alcohol within the previous 90 days (results submitted with the request); **AND**
   - If the test is positive for alcohol or illicit substances, the prescriber must submit documentation that the patient is actively attending a treatment program for substance abuse.
   - If the test is positive for a prescription controlled substance or a metabolite, the prescriber must document whether the patient has an active prescription for the attributable controlled substance (prescribers may consider using the Alaska PDMP, available at [http://alaskapdmp.com](http://alaskapdmp.com) as a tool to aid in the review).
   - If the test is positive for an unprescribed controlled substance or metabolite, the prescriber must submit documentation that the patient is actively attending a treatment program for substance abuse.
   - If the prescriber believes that the test documents an inaccurate or false positive result, an explanation must be submitted, and will be considered on a case-by-case basis.
10. If HCV/HIV co-infected, prescriber must provide documentation of CD4 count, HIV viral load, and HIV treatment regimen; **AND**

11. Regimens containing Olysio will not be approved.

**Criteria for Renewal Authorization Approval, with Approval Duration:**

1. For regimens with durations longer than 12 weeks, HCV RNA must be submitted for treatment weeks 4 and 8; **AND**

2. HCV RNA < 25 IU/mL at treatment week 4; **AND**

3. The prescriber must maintain documentation in the patient’s medical chart of the following information: HCV RNA level at treatment weeks 4 and 8, as well as HCV RNA levels after completion of therapy at week 12 post therapy (SVR12), and week 24 post therapy (SVR24).
   - This information shall be made available upon request.

4. Based on HCV genotype and prior treatment experience
   - Refer to Table 2 for regimen durations; authorization duration will be approved as follows:

<table>
<thead>
<tr>
<th>Regimen Duration</th>
<th>Authorization Duration</th>
</tr>
</thead>
<tbody>
<tr>
<td>12 weeks</td>
<td>12 weeks</td>
</tr>
<tr>
<td>16 weeks</td>
<td>12 weeks + 4 weeks*</td>
</tr>
<tr>
<td>24 weeks</td>
<td>12 weeks + 12 weeks*</td>
</tr>
</tbody>
</table>

*or under the discretion of Alaska Medicaid

5. Lost or stolen medication replacement requests will **not** be authorized.
**Authorized Regimens: Treatment Naïve**

<table>
<thead>
<tr>
<th>Genotype</th>
<th>Regimen</th>
<th>Duration</th>
</tr>
</thead>
<tbody>
<tr>
<td>GT 4 Metavir F2-4</td>
<td>Harvoni</td>
<td>12 weeks</td>
</tr>
<tr>
<td></td>
<td>Zepatier</td>
<td>12 – 16 weeks*</td>
</tr>
<tr>
<td>GT 4 Metavir F2-3</td>
<td>Sovaldi + PegIFN + ribavirin</td>
<td>12 weeks</td>
</tr>
<tr>
<td></td>
<td>Technivie + ribavirin</td>
<td>12 weeks</td>
</tr>
</tbody>
</table>

- **GT 4 Hepatocellular carcinoma awaiting liver transplantation AND meets Milan Criteria ›**
  - Sovaldi + ribavirin†,‡,§* 48 weeks or until liver transplant

- **GT 4 with decompensated cirrhosis [Child-Pugh B or C]**
  - Restricted to specialist

- **Mixed genotype**
  - Restricted to specialist

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1Prescribers are advised to review FDA approved labeling and other available clinical resources when determining appropriate regimens based on contraindications and warnings including clinically relevant drug-drug and drug-disease interactions as well as considerations for HIV/HCV co-infected individuals to ensure appropriate monitoring schema are taken into consideration. Refer to Table 4: Additional Criteria for Denial for limitations of use. Weight based ribavirin; §Refer to FDA approved labeling for use in individuals with impaired renal function. *If ribavirin is not used as part of the regimen indicated, Alaska Medicaid reserves the right to not extend treatment duration beyond what was initially authorized. ›Milan criteria: In single hepatocellular (HC) carcinomas, tumor ≤ 5 cm in diameter, OR In multiple HC carcinomas, no more than 3 tumor nodules, each ≤ 3 cm in diameter, AND No extrahepatic manifestations of the cancer or evidence of vascular invasion of tumor. •Duration varies depending on clinical situation and whether treatment naïve or PegIFN + Ribavirin experienced.

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**Table 3:**

<table>
<thead>
<tr>
<th>Authorized Clinical Reasons Ribavirin Cannot be Used</th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>Pregnancy</td>
<td>Hemoglobin &lt; 8.5 g/dL</td>
</tr>
<tr>
<td>Creatinine Clearance &lt; 50 mL/min</td>
<td>Hemoglobinopathies (e.g. sickle cell disease, thalassemia major)</td>
</tr>
<tr>
<td>Documented previous severe ribavirin hypersensitivity reaction</td>
<td>History of significant or unstable cardiac disease</td>
</tr>
</tbody>
</table>
Harvoni, Olysio, Sovaldi, Technivie, and Zepatier Criteria for Denial

1. Patient has not been tested for the use of illicit drugs, controlled substances, and alcohol within the previous 90 days (or results have not been submitted with the request); OR
   - If the test is positive for alcohol or illicit substances, the patient is not actively attending a treatment program for substance abuse.
   - If the test is positive for a prescription controlled substance or a metabolite, the prescriber has not documented whether the patient has an active prescription for the attributable controlled substance.
   - If the test is positive for an unprescribed controlled substance or metabolite, the patient is not actively attending a treatment program for substance abuse.

2. Diagnostic/disease severity/disease activity evidence based on biopsy, elastography, FibroSure, FibroTest, HepaScore, or FibroScan is not submitted with the request, or an APRI or FIB4 score alone is submitted to demonstrate disease severity/activity; OR

3. HCV RNA results not submitted with the request; OR

4. For regimens containing ribavirin, patient is pregnant or lactating; OR

5. Patient has a Child-Pugh score greater than 6 (class B or C) and treatment is not being managed by a liver disease specialist; OR

6. Regimens containing Olysio will not be approved.

<table>
<thead>
<tr>
<th>Table 4: Additional Criteria for Denial†</th>
</tr>
</thead>
<tbody>
<tr>
<td>Harvoni Genotype 2 or 3 infection</td>
</tr>
<tr>
<td>Taking a concomitant drug that has a significant clinical interaction or is contraindicated</td>
</tr>
<tr>
<td>Olysio Any genotype</td>
</tr>
<tr>
<td>Taking a concomitant drug that has a significant clinical interaction or is contraindicated</td>
</tr>
<tr>
<td>Sovaldi Genotype 5 or 6 infection</td>
</tr>
<tr>
<td>Taking a concomitant drug that has a significant clinical interaction or is contraindicated</td>
</tr>
<tr>
<td>Technivie Genotype 1, 2, 3, 5 or 6 infection</td>
</tr>
<tr>
<td>Taking a concomitant drug that is contraindicated (e.g., highly dependent on CYP3A for clearance &amp; for which elevated plasma concentrations are associated with serious events, or drugs that are moderate or strong inducers of CYP3A) or has a significant clinical interaction</td>
</tr>
<tr>
<td>Zepatier Genotype 2, 3, 5 or 6 infection</td>
</tr>
<tr>
<td>Concomitant use with OATP1B1/3 inhibitors, strong CYP3A inducers, or efavirenz.</td>
</tr>
</tbody>
</table>

† Prescribers are advised to review FDA approved labeling and other available clinical resources when determining appropriate regimens based on contraindications and warnings including clinically relevant drug-drug and drug-disease interactions as well as considerations for HIV/HCV co-infected individuals to ensure appropriate monitoring schema are taken into consideration.
ALASKA MEDICAID
Prior Authorization Criteria

Denial Due to Lack of Information:

If incomplete information is submitted on any prior authorization request, prescribers will have 7 calendar days to respond to the request for additional information, or the request will be non-clinically denied due to lack of information.

A re-review is possible with the submittal of a new complete PA request.

Criteria for Approval: Treatment Experienced/Retreatment Patients

1. Adult patient age \( \geq \) 18 years old; AND
2. Documentation of HCV genotype, HCV subtype, and HCV viral load is included in the authorization request; AND
3. Meets diagnosis and disease severity of Hepatitis C, Genotype 4 (GT 4), and Metavir Fibrosis score F2-F4 equivalent; AND
4. To confirm the Metavir fibrosis stage, the following tests or procedures will be accepted: biopsy, elastography, FibroSure, FibroTest, HepaScore, or FibroScan (Note: APRI and FIB-4 will not be accepted as confirmation of the Metavir fibrosis stage); AND
5. The Hepatitis C disease activity score must be submitted with the authorization request; AND
6. The patient agrees to complete regimen; AND
7. The prescriber agrees to maintain HCV RNA levels obtained at 12-weeks and 24-weeks post-therapy completion to demonstrate Sustained Virologic Response (SVR); AND
8. For patients previously treated with an NS5A inhibitor, NS5B inhibitor or a NS3/4a protease inhibitor, polymorphism testing results MUST be submitted; AND
9. If HCV/HIV co-infected, must provide documentation of CD4 count, HIV viral load, and HIV treatment regimen; AND
10. Documentation of previously trialed HCV therapies, dates of therapy, whether full therapy was completed or discontinued early, and, if discontinued early, the reason for the discontinuation is included in the authorization request; AND
11. Patient has been tested for the use of illicit drugs, controlled substances, and alcohol within the previous 90 days (results submitted with request); AND
   - If the test is positive for alcohol or illicit substances, the prescriber must submit documentation that the patient is actively attending a treatment program for substance abuse.
   - If the test is positive for a prescription controlled substance or a metabolite, the prescriber must document whether the patient has an active prescription for the attributable controlled substance (prescribers may consider using the Alaska PDMP, available at http://alaskapdmp.com as a tool to aid in the review).
   - If the test is positive for an unprescribed controlled substance or metabolite, the prescriber must submit documentation that the patient is actively attending a treatment program for substance abuse.
   - If the prescriber believes that the test documents an inaccurate or false positive result, an explanation must be submitted.
12. Regimens containing Olysio will not be approved.
<table>
<thead>
<tr>
<th>Genotype</th>
<th>Failed Treatment</th>
<th>Retreatment Regimen</th>
<th>Duration</th>
</tr>
</thead>
<tbody>
<tr>
<td>GT 4 Metavir F2-4</td>
<td>PegIFN + Ribavirin</td>
<td>Refer to Table 2 regimens</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Prior treatment with Direct Acting Antiviral based regimens (e.g. NS5A-inhibitor or NS5B polymerase inhibitor based regimens)</td>
<td>Harvoni + Ribavirin</td>
<td>12 - 24 weeks*</td>
</tr>
<tr>
<td></td>
<td>Sovaldi ± Ribavirin ± PegIFN</td>
<td>Technivie + Ribavirin (F2-3 only)</td>
<td>12 - 24 weeks*</td>
</tr>
</tbody>
</table>

\*Prescribers are advised to review FDA approved labeling and other available clinical resources when determining appropriate regimens based on contraindications and warnings including clinically relevant drug-drug and drug-disease interactions as well as considerations for HIV/HCV co-infected individuals to ensure appropriate monitoring schema are taken into consideration. °Refer to Table 4: Additional Criteria for Denial for limitations of use. ‡Weight based ribavirin; §Refer to FDA approved labeling for use in individuals with impaired renal function. *If ribavirin is not used as part of the regimen indicated, Alaska Medicaid reserves the right to not extend treatment duration beyond what was initially authorized. ▪ Duration varies depending on clinical situation and Metavir Fibrosis score.

Harvoni, Sovaldi, Technivie, and Zepatier Criteria for Denial

Same as listed above in the section for Treatment Naïve patients.

Criteria for Retreatment Renewal Authorization Approval, with Approval Duration:

Same as listed above in the section for Treatment Naïve patients.

References

6. FDA Antiviral Drugs Advisory Committee Meeting, October 25, 2013; Background Package for NDA 204671 sofosbuvir (GS-7977).
8. Poynard T, Ratziu V, Benmanov Y, DiMartino V, Bedossa P, Opolon P. Fibrosis in patients with hepatitis...


