



MEDICATION POLICY

Generic Name: Ombitasvir, Paritaprevir, and Ritonavir

Therapeutic Class or Brand Name: Technivie™

Applicable Drugs (if Therapeutic Class): N/A

Date of Origin: 9/22/15

Date Last Reviewed/Revised: 10/27/17

GPI Code: 1235990360

Prior Authorization Criteria (may be considered medically necessary when criteria I through VII are met):

- I. Documented diagnosis of chronic hepatitis C (CHC) genotype 4 infection.
- II. Documentation that patient meets ONE of the following criteria A, B, or C:
 - A. Has a Metavir score of F3 (advanced fibrosis).
 - B. Is post-liver transplant.
 - C. Has clinically severe extrahepatic manifestations of hepatitis C infection as evidenced by one of the following 1 or 2:
 1. Type 2 or 3 essential mixed cryoglobulinemia with end-organ manifestations (i.e. vasculitis).
 2. Proteinuria, nephrotic syndrome, or membranoproliferative glomerulonephritis.
- III. Patient must have a documented contraindication to Mavyret™ and Zepatier™.
- IV. Documentation of patient's Hepatitis C treatment history and baseline viral load.
- V. Documentation that patient's hepatitis C drug therapy is prescribed as outlined in the table under Authorization in the Approval Length section.
- VI. Minimum age requirement: 18 years old.
- VII. Prescriber is a Gastroenterologist, Infectious Disease Specialist, or Hepatologist.

Exclusion Criteria:

- As retreatment when there has been relapse after, or no response to, a prior treatment course with Daklinza™ (daclatasvir), Epclusa® (sofosbuvir/velpatasvir), Harvoni® (ledipasvir/sofosbuvir), Incivek® (telaprevir), Mavyret™ (glecaprevir/pibrentasvir), Olysio® (simeprevir), Sovaldi® (sofosbuvir), Technivie™ (ombitasvir, paritaprevir, and ritonavir), Victrelis® (boceprevir), Viekira Pak™/XR™ (dasabuvir, ombitasvir, paritaprevir, ritonavir), Vosevi® (sofosbuvir/velpatasvir/voxilaprevir), or Zepatier™ (elbasvir/grazoprevir).

Disclaimer: Medication Policies are developed to help ensure safe, effective and appropriate use of selected medications. They offer a guide to coverage and are not intended to dictate to providers how to practice medicine. Refer to Plan for individual adoption of specific Medication Policies. Providers are expected to exercise their medical judgment in providing the most appropriate care for their patients.



| |
|--------------------------|
| MEDICATION POLICY |
|--------------------------|

- Moderate (Child-Pugh B) to severe (Child-Pugh C) hepatic impairment.
- Known hypersensitivity (i.e. toxic epidermal necrolysis (TEN) or Stevens-Johnson syndrome) to ritonavir.
- Coadministration of Technivie™ with drugs that are highly dependent on CYP3A for clearance, moderate or strong inducers of CYP3A, or any of the drugs listed in the table below:

| Drug Class | Drugs within class |
|--|--|
| Alpha1-adrenoreceptor antagonist | Alfuzosin HCL |
| Anticonvulsants | Carbamazepine, phenytoin, phenobarbital |
| Anti-anginal | Ranolazine |
| Antiarrhythmic | Dronedarone |
| Anti-gout | Colchicine |
| Antimycobacterial | Rifampin |
| Antipsychotic | Lurasidone, pimozide |
| Ergot derivatives | Ergotamine, dihydroergotamine, ergonovine, methylergonovine |
| Ethinyl estradiol-containing products | Ethinyl estradiol-containing medications such as combined oral contraceptives |
| GI Motility Agent | Cisapride |
| Herbal Product | St. John's Wort (<i>Hypericum perforatum</i>) |
| HIV-Antiviral Agents | Atazanavir, atazanavir/ritonavir, darunavir/ritonavir, efavirenz, lopinavir/ritonavir, rilpivirine |
| HMG-CoA Reductase Inhibitors | Atorvastatin, lovastatin, pravastatin (if > 40mg/day), simvastatin |
| Immunosuppressants | Everolimus, sirolimus, tacrolimus |
| Long-acting beta-adrenoceptor agonist | Salmeterol |
| Non-nucleoside reverse transcriptase inhibitor | Efavirenz |
| Other protease inhibitors, NS5A inhibitors, or polymerase inhibitors used to treat chronic hepatitis C virus infection | Daklinza™ (daclatasvir), Epclusa® (sofosbuvir/velpatasvir), Harvoni® (ledipasvir/sofosbuvir), Mavyret™ (glecaprevir/pibrentasvir), Olysio® (simeprevir), Victrelis® (boceprevir), Viekira Pak™/XR™ (dasabuvir, ombitasvir, paritaprevir, ritonavir), Vosevi® (sofosbuvir/velpatasvir/voxilaprevir), Zepatier™ (elbasvir/grazoprevir) |
| Phosphodiesterase-5 (PDE5) inhibitor | Sildenafil when dosed as Revatio® for the treatment of pulmonary arterial hypertension (PAH) |
| Sedatives/hypnotics | Triazolam, orally administered midazolam |

Other Criteria:

Disclaimer: Medication Policies are developed to help ensure safe, effective and appropriate use of selected medications. They offer a guide to coverage and are not intended to dictate to providers how to practice medicine. Refer to Plan for individual adoption of specific Medication Policies. Providers are expected to exercise their medical judgment in providing the most appropriate care for their patients.



MEDICATION POLICY

- Technivie™ should be discontinued in patients who develop evidence of hepatic decompensation or failure.

Quantity/Days Supply Restrictions:

- 1 monthly carton (56 tablets) per 28 days.

Approval Length:

- **Authorization:** See table directly below.

| Drug Therapy | Cirrhosis | Authorization Duration | |
|-----------------|-----------|------------------------|------------------|
| | | G4 | |
| | | TN | TE |
| Technivie™+ RBV | No | 12w | 12w ¹ |
| | Comp | 12w | 12w ¹ |

TN = treatment naïve; TE = treatment experienced; Comp = compensated; RBV = ribavirin; pegIFN = peginterferon; w = weeks

¹For patients who have failed pegIFN/RBV.

- **Re-Authorization:** N/A

Appendix:

N/A

References:

1. <http://hcvguidelines.org/full-report-view>.
2. http://www.rxabbvie.com/pdf/technivie_pi.pdf.
3. Medi-Span.

Disclaimer: Medication Policies are developed to help ensure safe, effective and appropriate use of selected medications. They offer a guide to coverage and are not intended to dictate to providers how to practice medicine. Refer to Plan for individual adoption of specific Medication Policies. Providers are expected to exercise their medical judgment in providing the most appropriate care for their patients.



MEDICATION POLICY

| Historical Tracking Of Changes Made To Policy | |
|--|--|
| 10/27/2017 | <ol style="list-style-type: none"> 1. Changed “I. Documented diagnosis of chronic hepatitis C (CHC) genotype 4 infection without cirrhosis” to “I. Documented diagnosis of chronic hepatitis C (CHC) genotype 4 infection” under Prior Authorization Criteria. 2. Changed “III. Patient must have a documented contraindication to Zepatier™ and Epclusa®” to “III. Patient must have a documented contraindication to Mavyret™ and Zepatier™” under Prior Authorization Criteria. 3. Added “Mavyret™ (glecaprevir/pibrentasvir)” and “Vosevi® (sofosbuvir/velpatasvir/voxilaprevir)” to list of drugs following the statement “As retreatment when there has been relapse after, or no response to, a prior treatment course with...” under Exclusion Criteria. 4. Added “Atorvastatin” under Exclusion Criteria to table under “Coadministration of Technivie™ with...”, line entitled “HMG-CoA Reductase Inhibitors”. 5. Added “Immunosuppressants: Everolimus, sirolimus, tacrolimus” under Exclusion Criteria to table under “Coadministration of Technivie™ with...”. 6. Added “Mavyret™ (glecaprevir/pibrentasvir)” and “Vosevi® (sofosbuvir/velpatasvir/voxilaprevir)” under Exclusion Criteria to table under “Coadministration of Technivie™ with...”, line entitled “Other protease inhibitors, NS5A inhibitors, or polymerase inhibitors used to treat chronic hepatitis C virus infection”. |
| 7/30/2016 | <ol style="list-style-type: none"> 1. Changed “Viekira Pak™ (ombitasvir, paritaprevir, and ritonavir /dasabuvir)” to “Viekira Pak™/XR™ (dasabuvir, ombitasvir, paritaprevir, ritonavir)” in list of drugs following the statement “As retreatment when there has been relapse after, or no response to, a prior treatment course with...” under Exclusion Criteria. 2. Changed “Viekira Pak™ (ombitasvir, paritaprevir, and ritonavir/dasabuvir)” to “Viekira Pak™/XR™ (dasabuvir, ombitasvir, paritaprevir, ritonavir)” under Exclusion Criteria to table under “Coadministration of Technivie™ with...”, line entitled “Other protease inhibitors, NS5A inhibitors, or polymerase inhibitors used to treat chronic hepatitis C virus infection”. |
| 7/21/2016 | <ol style="list-style-type: none"> 1. Changed “III. Patient must have a documented contraindication to Zepatier™” to “III. Patient must have a documented contraindication to Zepatier™ and Epclusa®” under Prior Authorization Criteria. 2. Added “Epclusa® (sofosbuvir/velpatasvir)” under Exclusion Criteria to: 1) List of drugs following the statement “As retreatment when there has been relapse after, or no response to, a prior treatment course with...; 2) table under “Coadministration of Technivie™ with...”, line entitled “Other protease inhibitors, NS5A inhibitors, or polymerase inhibitors used to treat chronic hepatitis C virus infection”. 3. Added “Anti-anginal: Ranolazine”; “Antiarrhythmic: Dronedarone”; “Anti-gout: Colchicine”; “Antipsychotic: Lurasidone, pimozide”; and “GI Motility Agent: Cisapride” on table under “Coadministration of Technivie™ with...” under Exclusion Criteria. 4. Added “(elbasvir/grazoprevir)” following Zepatier™ to table under “Coadministration of Technivie™ with...”, line entitled “Other protease inhibitors, NS5A inhibitors, or polymerase inhibitors used to treat chronic hepatitis C virus infection” under Exclusion Criteria. 5. Unhighlighted authorization duration lengths on table below Authorization under Approval Length. 6. Added “pegIFN = peginterferon” beneath table below Authorization under Approval Length. |
| 3/21/2016 | <ol style="list-style-type: none"> 1. Changed “member” to “patient” throughout policy. 2. Changed “III. Documentation of member’s Hepatitis C treatment history” to “III. Patient must have a documented contraindication to Zepatier™; IV. Documentation of member’s Hepatitis C treatment history and baseline viral load; V. Documentation that patient’s hepatitis C drug therapy is prescribed as outlined in the table under Authorization in the Approval Length section” under Prior Authorization Criteria. 3. Changed “As retreatment when there has been relapse after, or no response to, a prior treatment course with Daklinza™ (daclatasvir), Harvoni® (ledipasvir/sofosbuvir), Incivek® (telaprevir), Olysio® |

Disclaimer: Medication Policies are developed to help ensure safe, effective and appropriate use of selected medications. They offer a guide to coverage and are not intended to dictate to providers how to practice medicine. Refer to Plan for individual adoption of specific Medication Policies. Providers are expected to exercise their medical judgment in providing the most appropriate care for their patients.



MEDICATION POLICY

Historical Tracking Of Changes Made To Policy

| | <p>(simeprevir), Sovaldi® (sofosbuvir), Technivie™ (ombitasvir, paritaprevir, and ritonavir), Victrelis® (boceprevir), or Viekira Pak™ (ombitasvir, paritaprevir, and ritonavir /dasabuvir)” to “As retreatment when there has been relapse after, or no response to, a prior treatment course with Daklinza™ (daclatasvir), Harvoni® (ledipasvir/sofosbuvir), Incivek® (telaprevir), Olysio® (simeprevir), Sovaldi® (sofosbuvir), Technivie™ (ombitasvir, paritaprevir, and ritonavir), Victrelis® (boceprevir), Viekira Pak™ (ombitasvir, paritaprevir, and ritonavir /dasabuvir), or Zepatier™ (elbasvir/grazoprevir)” under Exclusion Criteria.</p> <p>4. Added “Zepatier™” to “Other protease inhibitors, NS5A inhibitors, or polymerase inhibitors used to treat chronic hepatitis C virus infection” line on table underneath Exclusion Criteria.</p> <p>5. Changed table following Authorization under Approval Length from:</p> <table border="1" style="margin-left: 20px; border-collapse: collapse; width: 60%;"> <thead> <tr> <th style="background-color: #cccccc;">Patient Characteristics</th> <th style="background-color: #cccccc;">Treatment</th> <th style="background-color: #cccccc;">Authorization Duration</th> </tr> </thead> <tbody> <tr> <td>Genotype 4 without cirrhosis</td> <td>Technivie™ + ribavirin*</td> <td>12 weeks</td> </tr> </tbody> </table> <p style="margin-left: 20px;">*Technivie™ administered without ribavirin for 12 weeks may be considered for treatment-naïve patients who cannot take or tolerate ribavirin.</p> <p>to:</p> <table border="1" style="margin-left: 20px; border-collapse: collapse; width: 60%;"> <thead> <tr> <th rowspan="2" style="background-color: #ffcccc;">Drug Therapy</th> <th rowspan="2" style="background-color: #ccffcc;">Cirrhosis</th> <th colspan="2" style="background-color: #cccccc;">Authorization Duration</th> </tr> <tr> <th style="background-color: #cccccc;">G4</th> <th style="background-color: #cccccc;">G4</th> </tr> <tr> <th></th> <th></th> <th>TN</th> <th>TE</th> </tr> </thead> <tbody> <tr> <td rowspan="2" style="background-color: #ffcccc;">Technivie™+ RBV</td> <td style="background-color: #ccffcc;">No</td> <td>12w</td> <td>12w¹</td> </tr> <tr> <td style="background-color: #ccffcc;">Comp</td> <td>12w</td> <td>12w¹</td> </tr> </tbody> </table> <p style="margin-left: 20px;">TN = treatment naïve; TE = treatment experienced; Comp = compensated; RBV = ribavirin; w = weeks ¹For patients who have failed pegIFN/RBV.</p> | Patient Characteristics | Treatment | Authorization Duration | Genotype 4 without cirrhosis | Technivie™ + ribavirin* | 12 weeks | Drug Therapy | Cirrhosis | Authorization Duration | | G4 | G4 | | | TN | TE | Technivie™+ RBV | No | 12w | 12w ¹ | Comp | 12w | 12w ¹ |
|------------------------------|--|-------------------------|------------------|------------------------|------------------------------|-------------------------|-----------------|--------------|-----------|------------------------|--|----|----|--|--|----|----|-----------------|----|-----|------------------|------|-----|------------------|
| Patient Characteristics | Treatment | Authorization Duration | | | | | | | | | | | | | | | | | | | | | | |
| Genotype 4 without cirrhosis | Technivie™ + ribavirin* | 12 weeks | | | | | | | | | | | | | | | | | | | | | | |
| Drug Therapy | Cirrhosis | Authorization Duration | | | | | | | | | | | | | | | | | | | | | | |
| | | G4 | G4 | | | | | | | | | | | | | | | | | | | | | |
| | | TN | TE | | | | | | | | | | | | | | | | | | | | | |
| Technivie™+ RBV | No | 12w | 12w ¹ | | | | | | | | | | | | | | | | | | | | | |
| | Comp | 12w | 12w ¹ | | | | | | | | | | | | | | | | | | | | | |
| 11/18/2015 | <p>1. Changed “I. Documented diagnosis of chronic hepatitis C (CHC) genotype 4 infection” to “I. Documented diagnosis of chronic hepatitis C (CHC) genotype 4 infection without cirrhosis” under Prior Authorization Criteria.</p> <p>2. Changed “A. Has a Metavir score of F3 (advanced fibrosis) or F4 (compensated cirrhosis)” to “A. Has a Metavir score of F3 (advanced fibrosis)” under “II. Documentation that member meets ONE of the following criteria A, B, or C:” under Prior Authorization Criteria.</p> <p>3. Added “Technivie™ should be discontinued in patients who develop evidence of hepatic decompensation or failure” under Other Criteria.</p> <p>4. Changed “Genotype 4” to “Genotype 4 without cirrhosis” under Patient Characteristics on table under Approval Length.</p> | | | | | | | | | | | | | | | | | | | | | | | |

Disclaimer: Medication Policies are developed to help ensure safe, effective and appropriate use of selected medications. They offer a guide to coverage and are not intended to dictate to providers how to practice medicine. Refer to Plan for individual adoption of specific Medication Policies. Providers are expected to exercise their medical judgment in providing the most appropriate care for their patients.