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Subject: Hepatitis C Drug Therapy

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Position Statement	<u>Dosage/</u> <u>Administration</u>	Billing/Coding	Reimbursement	Program Exceptions	<u>Definitions</u>
Related Guidelines	<u>Other</u>	References	<u>Updates</u>		

DESCRIPTION:

<u>Hepatitis C virus</u> (HCV) is one of the leading causes of chronic liver disease in the United States. It is estimated that 3.9 million Americans have been infected with HCV with chronic liver disease developing in about 70% of the cases. About 1-5% of chronic liver disease cases due to HCV may die and HCV liver disease is the leading indication for liver transplants. The primary source for HCV infection is due to exposure to infected blood or blood products.

Currently the treatment of choice for chronic HCV infection in <u>compensated liver disease</u> is combination therapy. There are 6 major HCV <u>genotypes</u>, which predict treatment response. The majority of drug therapy data is dedicated to treating genotypes 1-4.

Production of hepatitis C protease inhibitors telaprevir (Incivek) and boceprevir (Victrelis) have been discontinued as of October 2014 and December 2015, respectively. In May 2018, AbbVie announced Technivie, Viekira Pak, and Viekira XR would be withdrawn from the US market in January 2019. Product will remain available through the end of 2018 to allow for current users to finish treatment. AbbVie has recommended that no new patients start treatment with Technivie, Viekira Pak, or Viekira XR beginning July 2018.

POSITION STATEMENT

Comparative Effectiveness

The Food and Drug Administration has deemed the drug(s) or biological product(s) in this coverage policy to be appropriate for self-administration or administration by a caregiver (i.e., not a healthcare professional). Therefore, coverage (i.e., administration) in a provider-administered setting such as an

outpatient hospital, ambulatory surgical suite, physician office, or emergency facility is not considered medically necessary.

NOTE:

- Glecaprevir-Pibrentasvir (Mavyret[™]), Ledipasvir-Sofosbuvir (Harvoni[®]), Sofosbuvir-Velpatasvir (Epclusa[®]), and Sofosbuvir-Velpatasvir-Voxilaprevir (Vosevi[™]) are the preferred direct acting antiviral products.
- Approval duration is based on genotype, treatment regimen, ribavirin and/or interferon eligibility, prior treatment (including treatment failures), baseline HCV RNA, transplant status, and number of doses received of requested regimen

Initiation of hepatitis C drug therapy **meets the definition of medical necessity** when all of the following criteria are met:

- 1. Indication for use is treatment of hepatitis C
- 2. Laboratory documentation of HCV genotype is provided
- 3. **ONE** of the following:
 - a. Member has been assessed for cirrhosis documentation from the medical record must be provided
 - b. Member is diagnosed with hepatocellular carcinoma awaiting liver transplantation documentation from the medical record must be provided
 - c. Member is a liver transplant recipient documentation from the medical record must be provided
- 4. Both the member and prescriber agree to provide documentation of virological response (i.e., HCV viral RNA level) upon completion of therapy
- 5. Member meets product specific criteria outlined in Table 1
- 6. Hepatitis C drug therapy does not exceed the maximum treatment duration for the requested regimen (see Table 2)

Continuation of hepatitis C drug therapy meets the definition of **medical necessity** when **ALL** of the following criteria are met:

- 1. Authorization/reauthorization has been previously approved by Florida Blue or another health plan, **OR** the member previously met all initiation criteria
- 2. Continuation of hepatitis C drug therapy does not exceed the maximum treatment duration for the requested regimen (see Table 2)

NOTE: Continuation of hepatitis C drug therapy will account for the number of doses the member has already received.

Approval duration: See Table 2

Table 1

Initiation criteria for	use of hepatitis C drug therapy
Product	Criteria
Brand	
Elbasvir-Grazoprevir	Use is a medical necessity when ALL criteria are met:
Zepatier	1. Member has a laboratory confirmed HCV genotype 1, 3, or 4
	 Use is alone (i.e., not in combination with any other hepatitis C therapy) OR in combination with ONE of the following:
	a. ribavirin
	b. sofosbuvir (Sovaldi)* – genotype 3 ONLY
	3. Member has a contraindication to treatment with each of the following direct acting antivirals (specific contraindication must be provided):
	a. glecaprevir-pibrentasvir (Mavyret)
	b. ledipasvir-sofosbuvir (Harvoni)
	c. sofosbuvir-velpatasvir (Epclusa)
	d. sofosbuvir-velpatasvir-voxilaprevir (Vosevi)
	 Total daily dose does not exceed one tablet (elbasvir 50 mg-grazoprevir 100 mg)
	Approval duration: See Table 2
Glecaprevir-	Use is a medical necessity when ALL criteria are met:
Pibrentasvir	1. Member has a laboratory confirmed HCV genotype 1, 2, 3, 4, 5, or 6
Mavyret	2. Use is alone (i.e., not in combination with any other hepatitis C therapy)
	 Total daily dose does not exceed three tablets (glecaprevir 100 mg- pibrentasvir 40 mg) or (if less than 12 years of age) five packets of oral pellets (glecaprevir 50 mg-pibrentasvir 20 mg)
	Approval duration: See Table 2
Ledipasvir-Sofosbuvir	Use is a medical necessity when ALL criteria are met:
Harvoni	1. Member has a laboratory confirmed HCV genotype 1, 4, 5, or 6
	 Use is alone (i.e., not in combination with any other hepatitis C therapy) OR in combination with ribavirin
	 Total daily dose does not exceed one tablet (ledipasvir 90 mg-sofosbuvir 400 mg-)
	Approval duration: See Table 2

Sofosbuvir	Use is a medical necessity when ALL criteria are met:
Sovaldi	 Member has a laboratory confirmed HCV genotype 3 Use is in combination with elbasvir-grazoprevir (Zepatier)* Total daily dose does not exceed 400 mg Approval duration: See Table 2
Sofosbuvir- Velpatasvir <i>Epclusa</i>	 Use is a medical necessity when ALL criteria are met: Member has a laboratory confirmed HCV genotype 1, 2, 3, 4, 5, or 6 Use is alone (i.e., not in combination with any other hepatitis C therapy)
Sofosbuvir- Velpatasvir- Voxilaprevir Vosevi	Use is a medical necessity when ALL criteria are met: 1. Member has a laboratory confirmed HCV genotype 1, 2, 3, 4, 5, or 6 2. Use is alone (i.e., not in combination with any other hepatitis C therapy) OR in combination with ribavirin 3. Total daily dose does not exceed one tablet (sofosbuvir 400 mg-velpatasvir 100 mg-voxilaprevir 100 mg) Approval duration: See Table 2
Prior authoriza	tion required. Member must meet prior authorization criteria for ALL prescribed

NOTE:

Table 2

- Glecaprevir-Pibrentasvir (Mavyret[™]), Ledipasvir-Sofosbuvir (Harvoni[®]), Sofosbuvir-Velpatasvir (Epclusa[®]), and Sofosbuvir-Velpatasvir-Voxilaprevir (Vosevi[™]) are the preferred direct acting antiviral products.
- Approval duration will account for any doses received since initiation of treatment.
- Treatment of a reinfection should follow criteria for initial treatment (i.e., treatment naïve)

Table 2: Maximum treatment/duration of approval for hepatitis C drug therapy

agents to be approved for combination therapy with the requested agent.

Genotype	Cirrhosis	Regimen	Required (ALL must be met for desired regimen)	Maximum Approval Duration
			Treatment naïve	12 weeks
		Epclusa [†] (or authorized	 Treatment naïve or treatment experienced with PEG-IFN +/- RBV +/- HCV protease inhibitor (telaprevir, boceprevir, simeprevir) Post-kidney transplantation 	12 weeks
		generic)	Treatment experienced with PEG-IFN +/- RBV +/- HCV protease inhibitor (telaprevir, boceprevir, simeprevir)	12 weeks
			Treatment experienced with sofosbuvir (Sovaldi) based regimen	12 weeks
		Harvoni [†] (or authorized generic) Harvoni [†] (or authorized generic) + RBV	 Treatment naïve or treatment experienced with PEG-IFN +/- RBV +/- HCV protease inhibitor (telaprevir, boceprevir, simeprevir) 	12 weeks
			Treatment experienced with sofosbuvir (Sovaldi) based regimen	12 weeks
1			Treatment naïve or treatment experienced with PEG-IFN +/- RBV	8 weeks
	Mavyret [†]	 Treatment naïve or treatment experienced with PEG-IFN +/- RBV +/- HCV protease inhibitor (telaprevir, boceprevir, simeprevir) Post-kidney transplantation 	12 weeks	
		Mavyret [†]	Treatment experienced with sofosbuvir (Sovaldi) based regimen	12 weeks
			Treatment experienced with simeprevir (Olysio) + sofosbuvir (Sovaldi)	12 weeks
			Treatment experienced with PEG-IFN +/- RBV + HCV protease inhibitor (telaprevir, boceprevir, simeprevir)	12 weeks
			Treatment experienced with NS5A inhibitor (daclatasvir, elbasvir, ledipasvir, ombitasvir, velpatasvir)	16 weeks
		Vosevi†	Treatment experienced with NS5A inhibitor (daclatasvir, elbasvir, ledipasvir, ombitasvir, or velpatasvir)	12 weeks

		 Treatment experienced with sofosbuvir (Sovaldi) based regimen 	12 weeks
		 Genotype 1a Treatment experienced with simeprevir (Olysio) + sofosbuvir (Sovaldi) based regimen 	12 weeks
	Zepatier	Treatment naïve or treatment experienced with PEG-IFN +/- RBV	12 weeks
		 Treatment experienced with PEG-IFN +/- RBV + HCV protease inhibitor (telaprevir, boceprevir, simeprevir) 	12 weeks
	Zepatier + RBV	 Genotype 1a with baseline NS5A RAS present Treatment naïve or treatment experienced with PEG-IFN +/- RBV +/- HCV protease inhibitor (telaprevir, boceprevir, simeprevir) 	16 weeks
	Epclusa [†] (or authorized generic)	 Treatment naïve or treatment experienced with PEG-IFN +/- RBV +/- HCV protease inhibitor (telaprevir, boceprevir, simeprevir) Allograft with or without compensated cirrhosis 	12 weeks
Post-liver transplant	Epclusa [†] (or authorized generic) + RBV	 Treatment naïve or treatment experienced with PEG-IFN +/- RBV +/- HCV protease inhibitor (telaprevir, boceprevir, simeprevir) Allograft with decompensated cirrhosis 	24 weeks
		 Treatment naïve or treatment experienced with PEG-IFN +/- RBV +/- HCV protease inhibitor (telaprevir, boceprevir, simeprevir) Allograft without cirrhosis 	12 weeks
	Harvoni [†] (or authorized	 Treatment naïve or treatment experienced with PEG-IFN +/- RBV +/- HCV protease inhibitor (telaprevir, boceprevir, simeprevir) Allograft with compensated cirrhosis 	12 weeks
	generic) + RBV	 Treatment naïve or treatment experienced with PEG-IFN +/- RBV +/- HCV protease inhibitor (telaprevir, boceprevir, simeprevir) Allograft with decompensated cirrhosis 	24 weeks

	Mavyret [†]	 Treatment naïve or treatment experienced with PEG-IFN +/- RBV +/- HCV protease inhibitor (telaprevir, boceprevir, simeprevir) Allograft with or without compensated cirrhosis 	12 weeks	
		Vosevi [†]	 Treatment experienced with NS5A inhibitor (daclatasvir, elbasvir, ledipasvir, ombitasvir, or velpatasvir) Allograft with or without compensated cirrhosis 	12 weeks
		Epclusa [†] (or authorized generic)	 Treatment naïve Treatment naïve or treatment experienced with PEG-IFN +/- RBV +/- HCV protease inhibitor (telaprevir, boceprevir, simeprevir) Post-kidney transplantation 	12 weeks
			 Treatment experienced with PEG-IFN +/- RBV +/- HCV protease inhibitor (telaprevir, boceprevir, simeprevir) 	12 weeks
			• Treatment experienced with sofosbuvir (Sovaldi) based regimen	12 weeks
			Treatment naïve	12 weeks
Col	Compensated	Harvoni [†] (or authorized generic)	 Treatment naïve or treatment experienced with PEG-IFN +/- RBV +/- HCV protease inhibitor (telaprevir, boceprevir, simeprevir) Post-kidney transplantation 	12 weeks
		Harvoni [†] (or authorized generic) + RBV	Treatment experienced with PEG-IFN +/- RBV +/- HCV protease inhibitor (telaprevir, boceprevir, simeprevir)	12 weeks
			Treatment naïve	8 or 12 weeks
		Mavyret [†]	 Treatment naïve or treatment experienced with PEG-IFN +/- RBV +/- HCV protease inhibitor (telaprevir, boceprevir, simeprevir) Post-kidney transplantation 	12 weeks
			Treatment experienced with PEG-IFN +/- RBV +/- HCV protease inhibitor (telaprevir, boceprevir, simeprevir)	12 weeks

		 Treatment experienced with sofosbuvir (Sovaldi) based regimen 	12 weeks
		• Treatment experienced with simeprevir (Olysio) + sofosbuvir (Sovaldi)	12 weeks
		 Treatment experienced with NS5A inhibitor (daclatasvir, elbasvir, ledipasvir, ombitasvir, velpatasvir) 	16 weeks
	Vosevi [†]	 Treatment experienced with NS5A inhibitor (daclatasvir, elbasvir, ledipasvir, ombitasvir, or velpatasvir) 	12 weeks
		 Treatment experienced with sofosbuvir (Sovaldi) based regimen 	12 weeks
	Zepatier	Treatment naïve or treatment experienced with PEG-IFN +/- RBV	12 weeks
	Zepatier + RBV	 Genotype 1a with baseline NS5A RAS present Treatment naïve or treatment experienced with PEG-IFN +/- RBV +/- HCV protease inhibitor (telaprevir, boceprevir, simeprevir) 	16 weeks
		 Treatment experienced with PEG-IFN +/- RBV + HCV protease inhibitor (telaprevir, boceprevir, simeprevir) 	12 weeks
	Epclusa [†] (or authorized generic)	 Treatment naïve or treatment experienced with PEG-IFN +/- RBV +/- HCV protease inhibitor (telaprevir, boceprevir, simeprevir) RBV contraindicated 	24 weeks
	Epclusa [†]	 Treatment naïve or treatment experienced with PEG-IFN +/- RBV +/- HCV protease inhibitor (telaprevir, boceprevir, simeprevir) 	12 weeks
Decompensated	(or authorized generic) + RBV	 Treatment experienced with a sofosbuvir (Sovaldi)- or NS5A (e.g.,daclatasvir, elbasvir, ledipasvir, ombitasvir)-based regimen 	24 weeks
	Harvoni [†] (or authorized generic)	 Treatment naïve or treatment experienced with PEG-IFN +/- RBV +/- HCV protease inhibitor (telaprevir, boceprevir, simeprevir) RBV contraindicated 	24 weeks
	Harvoni [†]	• Treatment naïve or treatment experienced with PEG-IFN +/- RBV +/-	12 weeks

		(or authorized generic)+ RBV	HCV protease inhibitor (telaprevir, boceprevir, simeprevir)	
			Treatment experienced with sofosbuvir- based regimen	24 weeks
		Epclusa [†] (or authorized generic)	 Treatment naïve or treatment experienced with PEG-IFN +/- RBV +/- HCV protease inhibitor (telaprevir, boceprevir, simeprevir) 	12 weeks
		genericy	Treatment experienced with sofosbuvir (Sovaldi) +/- RBV +/- PEG-IFN	12 weeks
			Treatment naïve or treatment experienced with PEG-IFN +/- RBV	8 weeks
		Mavyret [†]	 Treatment naïve or treatment experienced with PEG-IFN +/- RBV +/- HCV protease inhibitor (telaprevir, boceprevir, simeprevir) Post-kidney transplantation 	12 weeks
			Treatment experienced with sofosbuvir (Sovaldi) +/- RBV +/- PEG-IFN	12 weeks
		Vosevi [†]	 Treatment experienced with NS5A inhibitor (daclatasvir, elbasvir, ledipasvir, ombitasvir, or velpatasvir) 	12 weeks
2		Epclusa [†] (or authorized generic)	 Treatment naïve or treatment experienced with PEG-IFN +/- RBV +/- HCV protease inhibitor (telaprevir, boceprevir, simeprevir) Allograft with or without compensated cirrhosis 	12 weeks
	(or author generic) + Post-liver transplant Mavyre	Epclusa [†] (or authorized generic) + RBV	 Treatment naïve or treatment experienced with PEG-IFN +/- RBV +/- HCV protease inhibitor (telaprevir, boceprevir, simeprevir) Allograft with decompensated cirrhosis 	24 weeks
		Mavyret [†]	 Treatment naïve or treatment experienced with PEG-IFN +/- RBV +/- HCV protease inhibitor (telaprevir, boceprevir, simeprevir) Allograft with or without compensated cirrhosis 	12 weeks
		Vosevi [†]	 Treatment experienced with NS5A inhibitor (daclatasvir, elbasvir, ledipasvir, ombitasvir, or velpatasvir) Allograft with or without compensated cirrhosis 	12 weeks

	Epclusa [†] (or authorized generic)	 Treatment naïve or treatment experienced with PEG-IFN +/- RBV +/- HCV protease inhibitor (telaprevir, boceprevir, simeprevir) 	12 weeks
	genericy	 Treatment experienced with sofosbuvir (Sovaldi) +/- RBV +/- PEG-IFN 	12 weeks
		Treatment naïve	8 or 12 weeks
	Mavyret [†]	 Treatment naïve or treatment experienced with PEG-IFN +/- RBV +/- HCV protease inhibitor (telaprevir, boceprevir, simeprevir) Post-kidney transplantation 	12 weeks
		 Treatment experienced with PEG-IFN +/- RBV +/- sofosbuvir (Sovaldi) 	12 weeks
		Treatment experienced with sofosbuvir (Sovaldi) +/- RBV +/- PEG-IFN	12 weeks
	Vosevi [†]	 Treatment experienced with NS5A inhibitor (daclatasvir, elbasvir, ledipasvir, ombitasvir, or velpatasvir) 	12 weeks
	Epclusa [†] (or authorized generic)	 Treatment naïve or treatment experienced with PEG-IFN +/- RBV +/- HCV protease inhibitor (telaprevir, boceprevir, simeprevir) RBV contraindicated 	24 weeks
Decor	npensated Epclusa [†] (or authorized	 Treatment naïve or treatment experienced with PEG-IFN +/- RBV +/- HCV protease inhibitor (telaprevir, boceprevir, simeprevir) 	12 weeks
	generic) + RBV	Turnetus and according and accident	24 weeks
	Epclusa [†] (or authorized generic)	 Treatment naïve or treatment experienced with PEG-IFN +/- RBV +/- HCV protease inhibitor (telaprevir, boceprevir, simeprevir) 	12 weeks
3		Treatment naïve	8 weeks
	Mavyret [†]	 Treatment naïve or treatment experienced with PEG-IFN +/- RBV +/- HCV protease inhibitor (telaprevir, boceprevir, simeprevir) Post-kidney transplantation 	12 weeks

		Treatment experienced with PEG-IFN +/- RBV +/- sofosbuvir (Sovaldi)	16 weeks
		 Treatment experienced with PEG-IFN +/- RBV Y93H substitution is present 	12 weeks
	Vosevi [†] +/- RBV	 Treatment experienced with NS5A inhibitor (daclatasvir, elbasvir, ledipasvir, ombitasvir, or velpatasvir) 	12 weeks
		Treatment experienced with sofosbuvir (Sovaldi) based regimen	12 weeks
	Epclusa [†] (or authorized generic)	 Treatment naïve or treatment experienced with PEG-IFN + RBV +/- HCV protease inhibitor (telaprevir, boceprevir, simeprevir) Allograft with or without compensated cirrhosis 	12 weeks
	Epclusa [†] (or authorized generic) + RBV	 Treatment naïve or treatment experienced with PEG-IFN +/- RBV +/- HCV protease inhibitor (telaprevir, boceprevir, simeprevir) Allograft with decompensated cirrhosis 	24 weeks
	Mavyret [†]	Treatment naïve or treatment experienced	12 weeks
Post-liver transplant	Vosevi [†]	 Treatment experienced with NS5A inhibitor (daclatasvir, elbasvir, ledipasvir, ombitasvir, or velpatasvir) Allograft with or without compensated cirrhosis 	12 weeks
	Epclusa [†] (or authorized generic)	Treatment naïve	12 weeks
	Epclusa [†] (or authorized generic) + RBV	Treatment experienced with PEG-IFN +/- RBV +/- HCV protease inhibitor (telaprevir, boceprevir, simeprevir)	12 weeks
		Treatment naïve	8 or 12 weeks
	Mavyret [†]	 Treatment naïve or treatment experienced with PEG-IFN +/- RBV +/- HCV protease inhibitor (telaprevir, boceprevir, simeprevir) Post-kidney transplantation 	12 weeks
		Treatment experienced with PEG-IFN +/- RBV +/- sofosbuvir (Sovaldi)	16 weeks

		Variative PDV	Treatment naïveY93H substitution is present	12 weeks
			Treatment experienced with PEG-IFN +/- RBV	12 weeks
			 Treatment experienced with NS5A inhibitor (daclatasvir, elbasvir, ledipasvir, ombitasvir, or velpatasvir) 	12 weeks
			Treatment experienced with sofosbuvir (Sovaldi) based regimen	12 weeks
		Zepatier + Sovaldi	 Treatment experienced with PEG-IFN +/- RBV 	12 weeks
		Epclusa [†] (or authorized generic)	 Treatment naïve or treatment experienced with PEG-IFN +/- RBV +/- HCV protease inhibitor (telaprevir, boceprevir, simeprevir) RBV contraindicated 	24 weeks
	Decompensated	Epclusa [†] (or authorized generic) + RBV	 Treatment naïve or treatment experienced with PEG-IFN +/- RBV +/- HCV protease inhibitor (telaprevir, boceprevir, simeprevir) 	12 weeks
			 Treatment experienced with a sofosbuvir (Sovaldi)- or NS5A (e.g.,daclatasvir, elbasvir, ledipasvir, ombitasvir)-based regimen 	24 weeks
		Epclusa [†] (or authorized generic)	 Treatment naïve or treatment experienced with PEG-IFN +/- RBV +/- HCV protease inhibitor (telaprevir, boceprevir, simeprevir) 	12 weeks
	Absent	Harvoni [†] (or authorized generic)	 Treatment naïve or treatment experienced with PEG-IFN +/- RBV +/- HCV protease inhibitor (telaprevir, boceprevir, simeprevir) 	12 weeks
4			 Treatment naïve or treatment experienced with PEG-IFN +/- RBV +/- HCV protease inhibitor (telaprevir, boceprevir, simeprevir) Post-kidney transplantation 	12 weeks
		Mavyret [†]	Treatment naïve or treatment experienced with PEG-IFN + RBV +/- sofosbuvir (Sovaldi)	8 weeks
			 Treatment naïve or treatment experienced with PEG-IFN +/- RBV +/- HCV protease inhibitor (telaprevir, boceprevir, simeprevir) 	12 weeks

			Post-kidney transplantation	
		Vosevi [†]	 Treatment experienced with NS5A inhibitor (daclatasvir, elbasvir, ledipasvir, ombitasvir, or velpatasvir) 	12 weeks
		Zepatier	 Treatment naïve or treatment experienced with PEG-IFN +/- RBV 	12 weeks
		Zepatier + RBV	 On-treatment virologic failure with PEG- IFN +/- RBV 	16 weeks
		Epclusa [†] (or authorized generic)	 Treatment naïve or treatment experienced with PEG-IFN +/- RBV +/- HCV protease inhibitor (telaprevir, boceprevir, simeprevir) Allograft with or without compensated cirrhosis 	12 weeks
		Epclusa [†] (or authorized generic) + RBV	 Treatment naïve or treatment experienced with PEG-IFN +/- RBV +/- HCV protease inhibitor (telaprevir, boceprevir, simeprevir) Allograft with decompensated cirrhosis 	24 weeks
		Harvoni [†] (or authorized generic)	 Treatment naïve or treatment experienced with PEG-IFN +/- RBV +/- HCV protease inhibitor (telaprevir, boceprevir, simeprevir) Allograft without cirrhosis 	12 weeks
	Post-liver transplant	Harvoni [†] (or authorized generic) + RBV	 Treatment naïve or treatment experienced with PEG-IFN +/- RBV +/- HCV protease inhibitor (telaprevir, boceprevir, simeprevir) Allograft with compensated cirrhosis 	12 weeks
			 Treatment naïve or treatment experienced with PEG-IFN +/- RBV +/- HCV protease inhibitor (telaprevir, boceprevir, simeprevir) Allograft with decompensated cirrhosis 	24 weeks
		Mavyret [†]	 Treatment naïve or treatment experienced 	12 weeks
		Vosevi [†]	 Treatment experienced with NS5A inhibitor (daclatasvir, elbasvir, ledipasvir, ombitasvir, or velpatasvir) Allograft with or without compensated cirrhosis 	12 weeks
Co	ompensated	Epclusa [†]	• Treatment naïve	12 weeks

	(or authorized generic)	 Treatment experienced with PEG-IFN +/- RBV +/- HCV protease inhibitor (telaprevir, boceprevir, simeprevir) 	12 weeks
		Treatment naïve	12 weeks
	Harvoni [†] (or authorized generic)	 Treatment naïve or treatment experienced with PEG-IFN +/- RBV +/- HCV protease inhibitor (telaprevir, boceprevir, simeprevir) Post-kidney transplantation 	12 weeks
	Harvoni [†] (or authorized generic) + RBV	Treatment experienced with PEG-IFN +/- RBV +/- HCV protease inhibitor (telaprevir, boceprevir, simeprevir)	12 weeks
		Treatment naïve	8 or 12 weeks
	Mavyret [†]	 Treatment naïve or treatment experienced with PEG-IFN +/- RBV +/- HCV protease inhibitor (telaprevir, boceprevir, simeprevir) Post-kidney transplantation 	12 weeks
		Treatment experienced with PEG-IFN +/- RBV +/- sofosbuvir (Sovaldi)	12 weeks
	Vosevi [†]	Treatment experienced with NS5A inhibitor (daclatasvir, elbasvir, ledipasvir, ombitasvir, or velpatasvir)	12 weeks
	Zepatier	• Treatment naïve or treatment experienced with PEG-IFN +/- RBV	12 weeks
	Zepatier + RBV	On-treatment virologic failure with PEG- IFN +/- RBV	16 weeks
	Epclusa [†] (or authorized generic)	 Treatment naïve or treatment experienced with PEG-IFN +/- RBV +/- HCV protease inhibitor (telaprevir, boceprevir, simeprevir) RBV contraindicated 	24 weeks
Decompensated	Epclusa [†] (or authorized	 Treatment naïve or treatment experienced with PEG-IFN +/- RBV +/- HCV protease inhibitor (telaprevir, boceprevir, simeprevir) 	12 weeks
	generic) + RBV	 Treatment experienced with a sofosbuvir (Sovaldi)- or NS5A (e.g.,daclatasvir, elbasvir, ledipasvir, ombitasvir)-based regimen 	24 weeks

		Harvoni [†]	 Treatment naïve or treatment experienced with PEG-IFN +/- RBV +/- HCV protease inhibitor (telaprevir, boceprevir, simeprevir) RBV contraindicated 	24 weeks
		Harvoni [†] + RBV	 Treatment naïve or treatment experienced with PEG-IFN +/- RBV +/- HCV protease inhibitor (telaprevir, boceprevir, simeprevir) 	12 weeks
			Treatment experienced with sofosbuvir- based regimen	24 weeks
		Epclusa [†] (or authorized generic)	Treatment naïve or treatment experienced with PEG-IFN +/- RBV	12 weeks
			 Treatment naïve or treatment experienced with PEG-IFN +/- RBV 	12 weeks
	Absent or Compensated Mavyret Vosevi Harvoni (or authorized generic) Mavyret Vosevi	(or authorized	 Treatment naïve or treatment experienced with PEG-IFN +/- RBV +/- HCV protease inhibitor (telaprevir, boceprevir, simeprevir) Post-kidney transplantation 	12 weeks
			 No cirrhosis Treatment naïve or treatment experienced with PEG-IFN +/- RBV +/- sofosbuvir (Sovaldi) 	8 weeks
5		Mavyret [†]	 Treatment naïve or treatment experienced with PEG-IFN +/- RBV +/- HCV protease inhibitor (telaprevir, boceprevir, simeprevir) Post-kidney transplantation 	12 weeks
		Compensated cirrhosisTreatment naïve	8 or 12 weeks	
			 Compensated cirrhosis Treatment experienced with PEG-IFN +/- RBV +/- sofosbuvir (Sovaldi) 	12 weeks
		Vosevi [†]	Treatment experienced with NS5A inhibitor (daclatasvir, elbasvir, ledipasvir, ombitasvir, or velpatasvir)	12 weeks
	Post-liver transplant	Epclusa [†] (or authorized generic)	 Treatment naïve or treatment experienced with PEG-IFN +/- RBV +/- HCV protease inhibitor (telaprevir, boceprevir, simeprevir) Allograft with or without compensated cirrhosis 	12 weeks

	Epclusa [†] (or authorized generic) + RBV	 Treatment naïve or treatment experienced with PEG-IFN +/- RBV +/- HCV protease inhibitor (telaprevir, boceprevir, simeprevir) Allograft with decompensated cirrhosis 	24 weeks
	Harvoni [†] (or authorized generic)	 Treatment naïve or treatment experienced with PEG-IFN +/- RBV +/- HCV protease inhibitor (telaprevir, boceprevir, simeprevir) Allograft without cirrhosis 	12 weeks
	Harvoni [†] (or authorized	 Treatment naïve or treatment experienced with PEG-IFN +/- RBV +/- HCV protease inhibitor (telaprevir, boceprevir, simeprevir) Allograft with compensated cirrhosis 	12 weeks
	generic) + RBV	 Treatment naïve or treatment experienced with PEG-IFN +/- RBV +/- HCV protease inhibitor (telaprevir, boceprevir, simeprevir) Allograft with decompensated cirrhosis 	24 weeks
	Mavyret†	 Treatment naïve or treatment experienced with PEG-IFN +/- RBV +/- HCV protease inhibitor (telaprevir, boceprevir, simeprevir) Allograft with or without compensated cirrhosis 	12 weeks
	Vosevi [†]	 Treatment experienced with NS5A inhibitor (daclatasvir, elbasvir, ledipasvir, ombitasvir, or velpatasvir) Allograft with or without compensated cirrhosis 	12 weeks
	Epclusa [†] (or authorized generic)	 Treatment naïve or treatment experienced with PEG-IFN +/- RBV +/- HCV protease inhibitor (telaprevir, boceprevir, simeprevir) RBV contraindicated 	24 weeks
Decompensated	Epclusa [†] (or authorized	 Treatment naïve or treatment experienced with PEG-IFN +/- RBV +/- HCV protease inhibitor (telaprevir, boceprevir, simeprevir) 	12 weeks
	generic) + RBV	 Treatment experienced with a sofosbuvir (Sovaldi)- or NS5A (e.g.,daclatasvir, elbasvir, ledipasvir, ombitasvir)-based regimen 	24 weeks

		Harvoni [†] (or authorized generic)	 Treatment naïve or treatment experienced with PEG-IFN +/- RBV +/- HCV protease inhibitor (telaprevir, boceprevir, simeprevir) RBV contraindicated 	24 weeks
		Harvoni [†] (or authorized generic) + RBV	Treatment naïve or treatment experienced with PEG-IFN +/- RBV +/- HCV protease inhibitor (telaprevir, boceprevir, simeprevir)	12 weeks
		generic, r RDV	• Treatment experienced with sofosbuvir- based regimen	24 weeks
		Epclusa [†] (or authorized generic)	Treatment naïve or treatment experienced with PEG-IFN +/- RBV	12 weeks
			 Treatment naïve or treatment experienced with PEG-IFN +/- RBV 	12 weeks
	Absent or	Harvoni [†] (or authorized generic)	 Treatment naïve or treatment experienced with PEG-IFN +/- RBV +/- HCV protease inhibitor (telaprevir, boceprevir, simeprevir) Post-kidney transplantation 	12 weeks
			 No cirrhosis Treatment naïve or treatment experienced with PEG-IFN +/- RBV +/- sofosbuvir (Sovaldi) 	8 weeks
6 Compensated	Mavyret [†]	 Treatment naïve or treatment experienced with PEG-IFN +/- RBV +/- HCV protease inhibitor (telaprevir, boceprevir, simeprevir) Post-kidney transplantation 	12 weeks	
			Compensated cirrhosisTreatment naïve	8 or 12 weeks
			 Compensated cirrhosis Treatment experienced with PEG-IFN +/- RBV +/- sofosbuvir (Sovaldi) 	12 weeks
		Vosevi [†]	 Treatment experienced with NS5A inhibitor (daclatasvir, elbasvir, ledipasvir, ombitasvir, or velpatasvir) 	12 weeks
	Post-liver transplant	Epclusa [†] (or authorized generic)	 Treatment naïve or treatment experienced with PEG-IFN +/- RBV +/- HCV protease inhibitor (telaprevir, boceprevir, simeprevir) Allograft with or without compensated cirrhosis 	12 weeks

	Epclusa [†] (or authorized generic) + RBV	 Treatment naïve or treatment experienced with PEG-IFN +/- RBV +/- HCV protease inhibitor (telaprevir, boceprevir, simeprevir) Allograft with decompensated cirrhosis 	24 weeks
	Harvoni [†] (or authorized generic)	 Treatment naïve or treatment experienced with PEG-IFN +/- RBV +/- HCV protease inhibitor (telaprevir, boceprevir, simeprevir) Allograft without cirrhosis 	12 weeks
	Harvoni [†] (or authorized	 Treatment naïve or treatment experienced with PEG-IFN +/- RBV +/- HCV protease inhibitor (telaprevir, boceprevir, simeprevir) Allograft with compensated cirrhosis 	12 weeks
	generic) + RBV	 Treatment naïve or treatment experienced with PEG-IFN +/- RBV +/- HCV protease inhibitor (telaprevir, boceprevir, simeprevir) Allograft with decompensated cirrhosis 	24 weeks
	Mavyret†	 Treatment naïve or treatment experienced with PEG-IFN +/- RBV +/- HCV protease inhibitor (telaprevir, boceprevir, simeprevir) Allograft with or without compensated cirrhosis 	12 weeks
	Vosevi [†]	 Treatment experienced with NS5A inhibitor (daclatasvir, elbasvir, ledipasvir, ombitasvir, or velpatasvir) Allograft with or without compensated cirrhosis 	12 weeks
	Epclusa [†] (or authorized generic)	 Treatment naïve or treatment experienced with PEG-IFN +/- RBV +/- HCV protease inhibitor (telaprevir, boceprevir, simeprevir) RBV contraindicated 	24 weeks
Decompensated	Epclusa [†] (or authorized	 Treatment naïve or treatment experienced with PEG-IFN +/- RBV +/- HCV protease inhibitor (telaprevir, boceprevir, simeprevir) 	12 weeks
	generic) + RBV	 Treatment experienced with a sofosbuvir (Sovaldi)- or NS5A (e.g.,daclatasvir, elbasvir, ledipasvir, ombitasvir)-based regimen 	24 weeks

Harvoni [†] (or authorized generic)	 Treatment naïve or treatment experienced with PEG-IFN +/- RBV +/- HCV protease inhibitor (telaprevir, boceprevir, simeprevir) RBV contraindicated 	24 weeks
Harvoni [†] (or authorized	 Treatment naïve or treatment experienced with PEG-IFN +/- RBV +/- HCV protease inhibitor (telaprevir, boceprevir, simeprevir) 	12 weeks
generic) + RBV	• Treatment experienced with sofosbuvir- based regimen	24 weeks

[†]Preferred direct-acting antiviral product

PEG-IFN: peginterferon; RAS: resistance-associated substitution; RBV: ribavirin

DOSAGE/ADMINISTRATION:

THIS INFORMATION IS PROVIDED FOR INFORMATIONAL PURPOSES ONLY AND SHOULD NOT BE USED AS A SOURCE FOR MAKING PRESCRIBING OR OTHER MEDICAL DETERMINATIONS. PROVIDERS SHOULD REFER TO THE MANUFACTURER'S FULL PRESCRIBING INFORMATION FOR DOSAGE GUIDELINES AND OTHER INFORMATION RELATED TO THIS MEDICATION BEFORE MAKING ANY CLINICAL DECISIONS REGARDING ITS USAGE.

Dosage and administration varies considerably between specific indications for use and treatment history. A brief overview of selected products is provided in Table 3, however it is strongly recommended that the prescriber refer to current guidelines and product-specific labeling for complete dosing and administration instructions.

Table 3

Dosage/Administration of he	Dosage/Administration of hepatitis C drug therapy		
Product	Dosage/Administration		
Brand			
Elbasvir-Grazoprevir	One tablet daily		
Zepatier	Available as 50 mg elbasvir-100 mg grazoprevir tablet		
Glecaprevir-Pibrentasvir	Three tablets once daily with food or weight-based dosing for		
Mavyret	children less than 12 years of age and less than 45 kg		
	Available as 100 mg glecaprevir-40 mg pibrentasvir tablet or 50 mg		
	glecaprevir-20 mg pibrentasvir oral pellet packet		
Ledipasvir-Sofosbuvir	One tablet daily		
Harvoni	Available as a 90 mg ledipasivir-400 mg sofosbuvir tablet		
Sofosbuvir	400 mg once daily with food		
Sovaldi	Available as a 400 mg capsule		
Sofosbuvir-velpatasvir	One tablet daily or weight-based dosing for children less than 12		
Epclusa	years of age		
	Available as a 400 mg sofosbuvir-100 mg velpatasvir tablet, 200 mg		
	sofosbuvir-500 mg velpatasvir tablet, 200 mg sofosbuvir-500 mg		

	velpatasvir oral pellet packet, or 150 mg sofosbuvir-37.5 mg
	velpatasvir oral pellet packet
Sofosbuvir-velpatasvir-	One tablet daily
voxilaprevir	Available as a 400 mg sofosbuvir-100 mg velpatasvir-100 mg
Vosevi	voxilaprevir tablet

PRECAUTIONS:

Specific precautions and warnings are highlighted in Table 4.

Table 4

Precautions and warnings of	f hepatitis C drug therapy
Product	Precautions/Warnings
Brand	
Elbasvir-Grazoprevir	Contraindicated in patients with moderate or severe hepatic
Zepatier	impairment (Child-Pugh B or C). Contraindicated for co-
	administration with OATP1B1/3 inhibitors, strong CYP3A inducers,
	and efavirenz.
Glecaprevir-Pibrentasvir	Contraindicated in patients with severe hepatic impairment (Child-
Mavyret	Pugh C) and for co-administration with atazanavir and rifampin
Ledipasvir-Sofosbuvir	Bradycardia with amiodarone coadministration: Serious
Harvoni	symptomatic bradycardia may occur in patients taking amiodarone,
	particularly in patients also receiving beta blockers, or those with
	underlying cardiac comorbidities and/or advanced liver disease.
	Coadministration of amiodarone is not recommended. In patients
	without alternative, viable treatment options, cardiac monitoring is
	recommended.
	Use with P-gp inducers (e.g., rifampin, St. John's wort) may alter
	concentrations of ledipasvir and sofosbuvir
Sofosbuvir	Bradycardia with amiodarone coadministration: Serious
Sovaldi	symptomatic bradycardia may occur in patients taking amiodarone,
Sovarar	particularly in patients also receiving beta blockers, or those with
	underlying cardiac comorbidities and/or advanced liver disease.
	Coadministration of amiodarone is not recommended. In patients
	without alternative, viable treatment options, cardiac monitoring is
	recommended.
	Use with caution in combination with drugs that are potent P-gp
	inducers in the intestine (e.g., rifampin, St. John's wort)
Sofosbuvir-velpatasvir	Bradycardia may occur in patients taking amiodarone, particularly in
Epclusa .	patients also receiving beta blockers, or those with underlying
•	cardiac comorbidities and/or advanced liver disease.
	Coadministration of amiodarone is not recommended. In patients

	without alternative viable treatment options, cardiac monitoring is
	recommended.
Sofosbuvir-velpatasvir-	Contraindicated for co-administration with rifampin.
voxilaprevir	Bradycardia with amiodarone coadministration: Serious
Vosevi	symptomatic bradycardia may occur in patients taking amiodarone,
	particularly in patients also receiving beta blockers, or those with
	underlying cardiac comorbidities and/or advanced liver disease.
	Coadministration of amiodarone is not recommended. In patients
	without alternative, viable treatment options, cardiac monitoring is
	recommended.

BILLING/CODING INFORMATION:

HCPCS Coding:

J8499	Prescription drug, oral, non-chemotherapeutic, not otherwise specified
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ICD-10 Diagnosis Codes That Support Medical Necessity:

B18.2	Chronic viral hepatitis C
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REIMBURSEMENT INFORMATION:

Refer to section entitled **POSITION STATEMENT**.

PROGRAM EXCEPTIONS:

Federal Employee Program (FEP): Follow FEP guidelines.

State Account Organization (SAO): Follow SAO guidelines.

Medicare Advantage Products: No National Coverage Determination (NCD) or Local Coverage Determination (LCD) were found at the time of the last guideline revised date.

Medicare Part D: Florida Blue has delegated to Prime Therapeutics authority to make coverage determinations for the Medicare Part D services referenced in this guideline.

If this Medical Coverage Guideline contains a step therapy requirement, in compliance with Florida law 627.42393, members or providers may request a step therapy protocol exemption to this requirement if based on medical necessity. The process for requesting a protocol exemption can be found at Coverage Protocol Exemption Request.

DEFINITIONS:

Compensated liver disease: condition is stable and no clinical signs of physical or metabolic changes such as, but not limited to hepatic encephalopathy, variceal bleeding, ascites, acute changes in bilirubin, albumin, platelet count or bleeding time that cannot be not attributed to other organic cause(s).

Decompensated cirrhosis: moderate or severe hepatic impairment (Child Pugh class B or C)

Early Virological Response (EVR): $a \ge 2 \log 10$ reduction in HCV RNA level compared to baseline HCV RNA level (partial EVR) or HCV RNA negative at treatment week 12 (complete EVR).

Genotype: the genetic constitution of an organism or cell, as distinct from its expressed features or phenotype.

Hepatitis C virus: a viral infection caused by a single-stranded RNA virus of the family Flaviviridae (species Hepatitis C virus of the genus Hepacivirus) that tends to persist in the blood serum and is usually transmitted by infected blood (as by injection of an illicit drug, blood transfusion, or exposure to blood or blood products) and that accounts for most cases of chronic liver disease.

Null responder: those that fail to achieve a 2log10 HCV RNA decrease at week 24 of treatment.

Phenotype: the observable properties of an organism that are produced by the interaction of the genotype and the environment.

Rapid Virological Response (RVR): those that are HCV RNA negative at treatment week 4.

Responder: those that achieve an HCV RNA decrease greater than or equal to 2log10 by treatment week 12 followed by a sustained virological response (SVR) at the end of treatment.

Partial responder: those that achieve an HCV RNA decrease greater than or equal to 2log10 by treatment week 12, but fail to have a sustained virological response (SVR) at the end of treatment.

Poor interferon responder: those with less than 1 log 10 decline in viral load at treatment week 4.

Relapse: HCV RNA levels undetectable at end of treatment, but are detectable again at post treatment follow-up.

Sustained Virological Response (SVR): a negative HCV RNA level 24 weeks after cessation of treatment.

RELATED GUIDELINES:

None applicable.

OTHER:

Tab	Table 5: Metavir Fibrosis Scores		
FO	No fibrosis		
F1	Portal fibrosis without septa		
F2	Portal fibrosis with few septa		
F3	Numerous septa without cirrhosis		
F4	Cirrhosis		

Table 6: Child Turcotte Pugh (CTP) classification of the severity of cirrhosis					
	Class A	Class B	Class C		
Total points	5–6	7–9	10–15		
Factor	1 Point	2 Points	3 Points		

Total bilirubin (μmol/L)	<34	34–50	>50
Serum albumin (g/L)	>35	28–35	<28
Prothrombin time/international	<1.7	1.71-2.30	>2.30
normalized ratio			
Ascites	None	Mild	Moderate to Severe
Hepatic encephalopathy	None	Grade I–II	Grade III–IV
		(or suppressed with	(or refractory)
		medication)	

Table 7: Interferon ineligible

IFN ineligible is defined as one or more of the below:

- Intolerance to IFN
- Autoimmune hepatitis and other autoimmune disorders
- Hypersensitivity to PEG or any of its components
- Decompensated hepatic disease
- Major uncontrolled depressive illness
- A baseline neutrophil count below 1500/μL, a baseline platelet count below 90,000/μL or baseline hemoglobin below 10 g/dL
- A history of preexisting cardiac disease

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COMMITTEE APPROVAL:

This Medical Coverage Guideline (MCG) was approved by the Florida Blue Pharmacy Policy Committee on 07/09/25.

GUIDELINE UPDATE INFORMATION:

01/01/06	New Medical Coverage Guideline.
12/15/06	Reviewed: reformatted, added note regarding MCG for hepatitis C only and combo
	therapy is therapy of choice, and added HCPCS codes and updated CPT coding. MCG
	revised to include Medicare Part D as a program exception.
08/15/07	Review and revision to guideline; consisting of reformatting guideline, adding criteria for
	treating genotypes 1 and 4 hepatitis C beyond 12 weeks, updated internet links and
	updated references.

11/15/07	Devision of avidalina, consisting of shanning all references to longth of the growth was
11/15/07	Revision of guideline; consisting of changing all references to length of therapy to weeks
07/45/00	to be consistent with Position Statement.
07/15/08	Review and revision to guideline; consisting of adding Alferon® N to MCG, adding note
	stating that Roferon® brand of interferon alfa-2a has been discontinued by the
	manufacturer, adding genotypes 5 & 6 to the guideline, adding boxed warning under
	precautions section and updating references and links.
01/01/09	Annual HCPCS coding update: revised descriptor codes J9212, J9213, J9214 and J9215;
00/15/00	deleted 90772; added 96372.
08/15/09	Review and revision to guideline; consisting of adding consideration for up to 72 weeks
	for genotype 1 infection, highlight "notes" in position statement, updating precautions
	section and references.
09/15/10	Review and revision to guideline; consisting of updating references.
01/15/11	Revision to guideline; consisting of adding ICD-10 codes.
07/13/11	Review and revision to guideline; consisting of addition of treatment post liver
	transplant and triple combination therapy to the position statement, added definition of
	<u>responder</u> , <u>partial responder</u> , <u>null responder</u> , HCPCS code update, deleted Roferon A,
	added drug information for Incivek TM (telaprevir) and Victrelis® (boceprevir).
10/15/11	Updated guideline to add note preferred pegylated interferon product Pegasys,
	removed Alferon N from the description and coding section, updated coverage to
	cirrhosis to include compensated only, updated to include justification for need of
	monotherapy, added pediatric use dual therapy, updated the definitions.
08/15/12	Review and revision to guideline; consisting of updated position statement and drug-
	drug interaction lists to include new data and recommendations regarding the drug-drug
	interactions with boceprevir and ritonavir boosted HIV therapies, updated the
	formatting to include all criteria in a table format, added consideration for extended
	dosing in genotype 2, 3, added definition of compensated liver disease.
02/15/13	Revision to guideline; consisting of updating format in the note for dual therapy.
	Removed viral titer quantification < 100 IU/ml from boceprevir TW 8.
08/15/13	Review and revision to guideline; consisting of updating the coverage to include HCV
	recurrence in liver transplant. Updated response guided therapy chart to provide 24
	week of therapy results for telaprevir (Incivek™). Minor formatting changes. Updated
	ICD-9, ICD-10, and Program Exceptions section. Updated sources.
02/01/14	Revision to guideline; consisting of updating the position statement to include HCV
	therapy with simeprevir and sofosbuvir and updated the Response Guided Therapy for
	Table 1 and Therapy by Genotype Table 2. Updated introduction, product information
	and references.
02/15/14	Revision to guideline; consisting of updating position statement due to updated ASSLD
	guidelines in the treatment of Hepatitis C infection.
08/15/14	Review and revision to guideline; consisting of description, position statement,
	dosage/administration, precautions/warning, references.
10/15/14	Revision to guideline; consisting of position statement, dosage/administration,
	precautions/warnings, references, description.
11/15/14	Revision to guideline; consisting of position statement.
•	•

12/15/14	Revision to guideline; consisting of position statement, dosage/administration, coding, other.
02/15/15	Revision to guideline; consisting of position statement, dosage/administration,
	precautions/warnings, references.
04/01/15	Revisin to guideline; consisting of position statement, dosage/administration,
	precautions/warnings
06/15/15	Revision to guideline; consisting of position statement
08/15/15	Review and revision to guidelines; consisting of precautions, program exceptions,
	references
09/15/15	Revision to guideline; consisting of updating position statement, dosage/administration,
	precautions, coding, and references.
11/01/15	Revision: ICD-9 Codes deleted.
12/18/15	Revision to guideline; consisting of updating position statement and
	dosage/administration.
03/15/16	Revision to guideline; consisting of updating position statement and
	dosage/administration, precautions/warnings, references.
08/15/16	Review and revision to guideline; consisting of updating position statement and
	dosage/administration, precautions/warnings, references.
09/16/16	Revision of guideline to update position statement, dosage/administration,
	precautions/warnings, references.
11/15/16	Revision to guideline; consisting of updating Tabe 2 with "PEG-IFN + RBV".
02/15/17	Revision to guideline; consisting of updating position statement, precautions, coding.
8/15/17	Review and revision to guideline consisting of updating position statement and
	references.
09/15/17	Revision to guideline to include Vosevi in position statement, dosing and administration, references.
11/15/17	Revision to guideline to include Mayvret in position statement, dosing and
	administration, references.
08/15/18	Revision to guideline to update position statement, references.
11/15/18	Revision to guideline to update position statement
8/15/19	Revision to guideline to update references.
12/15/19	Updated position statement
07/15/20	Updated position statement
08/15/20	Updated position statement.
08/15/21	Updated position statement.
09/15/21	Updated position statement with new oral pellet formulations of Mavyret and Epclusa
12/15/21	Updated position statement to remove 8 week Harvoni requirement for GT 1
08/15/22	Revision to guideline; consisting of updating position statement, references
08/15/25	Revision to guideline; consisting of updating position statement, dosage/administration,
	precautions, and references.