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Screening for Hepatitis C Virus Infection in Adolescents and Adults: A Systematic Review Update for the U.S. Preventive Services Task Force

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Abstract

Background: Prior reviews on hepatitis C (HCV) infection screening and treatment used by the U.S. Preventive Services Task Force (USPSTF) to inform its 2013 recommendation found interferon-containing antiviral therapies associated with sustained virologic response (SVR) rates of 68 percent to 78 percent and an association between SVR after antiviral therapy and improved clinical outcomes. Interferon-containing regimens were associated with a high rate of harms. Since the prior reviews, interferon-containing antiviral therapies have been replaced by all-oral direct acting antiviral (DAA) regimens.

Purpose: To systematically review the evidence on screening for HCV infection in asymptomatic adults and adolescents, including effects of DAA regimens and interventions to prevent mother-to-child transmission.

Data Sources: We searched the Cochrane Central Register of Controlled Trials and Cochrane Database of Systematic Reviews, Ovid MEDLINE and ClinicalTrials.gov through February 2019 and manually reviewed reference lists.

Study Selection: Randomized controlled trials (RCTs), non-randomized trials, and cohort studies of HCV screening, antiviral therapy, and interventions to prevent mother-to-child transmission of HCV infection on SVR and clinical outcomes; and cohort studies on the association between an SVR after antiviral therapy versus no SVR and clinical outcomes. Treatment studies focused on populations without cirrhosis who are more likely to be asymptomatic and identified by screening.

Data Extraction: One investigator abstracted data, and a second investigator checked data abstraction for accuracy. Two investigators independently assessed study quality using methods developed by the USPSTF.

Data Synthesis (Results): No study evaluated the benefits of HCV screening versus no screening, or the yield of repeat versus one-time screening. Previously reviewed studies found that HCV screening might be associated with negative psychological and social consequences, but had important methodological limitations; no new studies were identified. One new study found similar diagnostic yield of risk-based and birth cohort screening, but it was retrospective and assumed perfect implementation of risk-based screening. Ten trials reported improvements in some quality of life and functional outcomes following DAA treatment compared with prior to treatment, but differences were small, studies were open-label, and there was no non-DAA comparison group. Forty-nine trials found DAA regimens associated with pooled SVR rates that ranged from 95.5 percent to 98.9 percent across genotypes; rates of serious adverse events (1.9%) and withdrawal due to adverse events (0.4%) were low. Seven trials reported SVR rates in adolescents with DAA therapy similar to those observed in adults. An SVR after antiviral therapy was associated with decreased risk of all-cause mortality (13 studies, pooled hazard ratio [HR] 0.40, 95% confidence interval [CI] 0.28 to 0.56), liver mortality (4 studies, pooled HR 0.11, 95% CI, 0.04 to 0.27), cirrhosis (4 cohorts in 3 studies, pooled HR 0.36, 95% CI, 0.33 to 0.40), and hepatocellular carcinoma (20 studies, pooled HR 0.29, 95% CI, 0.23 to 0.38) versus no SVR, after adjustment for potential confounders. New evidence on interventions to reduce the risk of mother-to-infant transmission was limited and did not change the conclusion from the prior review that no intervention has been clearly demonstrated to reduce risk.

Limitations: Most DAA trials were not randomized and did not have a non-DAA comparison group, almost all DAA trials relied on SVR as the main efficacy outcome, observational studies varied in how well they adjusted for confounders, and few studies evaluated the effectiveness of DAA regimens in adolescents.

Conclusions: The USPSTF previously determined that HCV screening is highly accurate. Currently recommended all-oral DAA regimens are associated with very high SVR rates (95.5% to 98.9% across genotypes) and few harms relative to older antiviral therapies. An SVR after antiviral therapy is associated with improved clinical outcomes compared with no SVR, after adjusting for potential confounders. Direct evidence on the benefits of HCV screening remains unavailable, and direct evidence on the effects of antiviral therapy on clinical outcomes remains limited but indicates improved long-term outcomes.

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Chapter 1. Introduction and Background

Purpose

The purpose of this report is to systematically review the evidence on screening for hepatitis C virus (HCV) infection in asymptomatic adults and adolescents without known liver enzyme abnormalities. This report updates prior (2013) U.S. Preventive Services Task Force (USPSTF) reviews on screening for HCV infection^{1,2} and prenatal screening,^{2,3} and a comparative effectiveness review on antiviral treatments.^{4,5} Although prior reports focused on benefits and harms of screening and treatment in adults, this report expands the population to include adolescents. For treatments, this report focuses on currently recommended direct acting antiviral (DAA) therapies and interventions to potentially reduce risk of mother-to-child transmission. It will be used by the USPSTF to update its 2013 recommendation on screening for HCV infection in adults and potentially inform a new recommendation on HCV screening in adolescents.

In 2013, the USPSTF recommended screening for HCV infection in adults at high risk for infection and recommended offering one-time screening for HCV infection in adults born between 1945 and 1965 ("birth cohort" screening) (**B Recommendation**).⁶ This recommendation represented a change from the prior (2004) USPSTF recommendation, which found insufficient evidence to recommend for or against HCV screening in adults at high risk for infection (**I recommendation**); the 2004 USPSTF recommendation did not address birth cohort screening and recommended against HCV screening in persons not at increased risk (**D recommendation**).⁷ The USPSTF did not issue a recommendation specifically on prenatal HCV screening, but noted that antiviral therapies were contraindicated during pregnancy and found inadequate evidence that labor management and breastfeeding strategies in HCV-infected persons are effective at reducing risk for mother-to-child transmission.

The basis for the change in the 2013 USPSTF recommendation was evidence that newer antiviral therapies are more effective than prior therapies in achieving the intermediate outcome of sustained virologic response (SVR) and evidence showing that SVR after antiviral therapy is associated with improved clinical outcomes (all-cause and liver-related mortality and hepatocellular carcinoma [HCC]), with few serious treatment-related harms that generally resolve after treatment discontinuation.⁶ The USPSTF also considered the prevalence of HCV infection in high-risk persons (e.g., \geq 50% in persons who inject drugs [PWID]) and in persons born between 1945 and 1965 (3% to 4%), and modeling studies that indicated cost-effectiveness of the birth cohort screening strategy.^{8,9} The USPSTF found few serious adverse events with liver biopsy performed for the diagnostic evaluation of persons with HCV infection and noted that fewer biopsies were being performed due to the availability of accurate noninvasive tests for evaluating liver fibrosis. The USPSTF had previously found that screening tests are highly accurate for diagnosing HCV infection (overall sensitivity 94% and specificity 97%).⁷

Condition Background

Condition Definition

HCV is a single-stranded, positive-sense ribonucleic acid (RNA) virus of the family Flaviviridae. HCV infection can range from mild and self-limited to a serious, lifelong illness that can result in cirrhosis, liver failure, and HCC.¹⁰ In most cases (78% to 85%), acute HCV leads to chronic HCV.¹⁰ HCV is primarily acquired by exposures to infected blood, with injection drug use the strongest risk factor. In the United States, approximately 70 to 77 percent of HCV infections are caused by genotype 1 (subtypes 1a or 1b), 13 to 16 percent by genotype 2, 12 percent by genotype 3, and less than 5 percent by genotypes 4, 5, or 6 combined.^{11,12}

Prevalence and Burden of Disease/Illness

HCV is the most common chronic bloodborne pathogen in the United States. The number of U.S. residents with past or current HCV infection (positive for anti-HCV antibody) is estimated at 4.1 million (range 3.4 million to 4.9 million); of these, an estimated 2.4 million (range 2.0 million to 2.8 million) are currently infected, defined as having HCV detectable in the blood (viremia).^{10,13,14} Approximately three-quarters (78% to 85%) of those who test positive for anti-HCV antibody have chronic infection;^{10,15} those with anti-HCV antibody but no viremia are considered to have cleared the infection. The estimated prevalence of chronic HCV infection during the years 2013 to 2016 was approximately 1.0 percent (95% confidence interval [CI], 0.8 to 1.1%).¹⁶ Persons born between 1945 and 1965 comprise approximately 27 percent of the U.S. population but account for approximately three-quarters of all HCV infection,¹⁶ and are at 6.0- to 9.5-fold increased risk of having HCV infection compared with younger adults.^{17,18} Males are at increased risk for HCV infection compared with females (odds ratio [OR] 1.6, 95% CI, 1.1 to 2.4), and non-Hispanic black persons are at increased risk compared with 62 other races/ethnicities (OR 1.6, 95% CI, 1.1 to 2.3), excluding American Indian/Alaska Natives.¹⁸ American Indian/Alaska Natives, who are often not included in national seroprevalence surveys, have higher HCV-related mortality than non-Hispanic black persons.¹⁹ Reported cases of acute HCV infection increased approximately 3.5-fold from 2010 through 2016.²⁰ After adjusting for under-ascertainment and under-reporting, an estimated 41,200 (95% CI, 32,600 to 140,600) new HCV infections occurred in the United States in 2016.²⁰ The increase in acute HCV incidence has most impacted young, white PWID living in non-urban areas.²¹⁻²³

Data also indicate an increase in the number of reproductive aged women (15 to 44 years of age) with HCV infection.^{24,25} An estimated 29,000 females with HCV infection give birth annually in the United States, resulting in 1,700 cases of infected infants.²⁵ Trends in HCV epidemiology, prevalence, and incidence are discussed in more detail in Contextual Question 1.

Etiology and Natural History

HCV infection is a leading cause of complications from chronic liver disease. The number of deaths due to HCV infection ranged from 18,650 to 19,629 from 2012 to 2015 (4.9 to 5.0 deaths/100,000) and decreased to 18,153 in 2016 (4.5 deaths/100,000).²⁰ Despite likely

underestimation, HCV-related mortality exceeds mortality associated with 60 other nationally notifiable infectious conditions combined.²⁶ According to the Centers for Disease Control and Prevention, of every 100 persons infected with HCV, approximately 60-70 will develop chronic liver disease, 5 to 20 will develop cirrhosis over a period of 20 to 30 years, and 1 to 5 will die from the consequences of liver cancer or cirrhosis.²⁷ HCV without cirrhosis is associated with worse quality of life and symptoms (e.g., fatigue) compared with not having HCV infection.²⁸⁻³² Other extrahepatic manifestations of HCV infection include mixed cryogloblinemias, non-Hodgkin lymphoma, type II diabetes mellitus and insulin resistance, cardiovascular disease, and renal disease.³³

The natural course of chronic HCV infection varies. Some patients with chronic HCV infection have only mild liver disease after decades of infection or never develop histologic evidence of liver disease.³⁴ In other patients, inflammation and fibrosis of the liver may progress to cirrhosis, which can lead to end-stage liver disease or HCC. In persons with cirrhosis due to HCV infection, the annual incidence of HCC is 1 to 4 percent.³⁵ Once cirrhosis develops, patients have a much higher risk of death, and some may benefit from liver transplantation. Until recently, chronic HCV was the leading indication for liver transplantation in the United States.^{36,37} The number of HCV-related liver transplants in the United States declined from a peak of 1,905 in 2014 to 1,535 in 2016.³⁶ Well-established predictors of advanced fibrosis in those with chronic HCV infection include older age at infection, longer duration of infection, male sex, concomitant HIV or hepatitis B virus (HBV) infection, and greater alcohol use.^{34,38,39} Other factors that may be associated with increased risk of fibrosis include insulin resistance, hepatic steatosis, higher viral load, and the presence of certain human leukocyte antigen (HLA) class II polymorphisms. Once a person develops advanced (METAVIR stage 3) fibrosis, the risk of progression to cirrhosis is around 10 percent per year.⁴⁰

Estimating the proportion of patients in the general population with HCV infection who progress to cirrhosis is difficult because the time of acquisition is often unclear and important endpoints often do not occur until after decades of infection; in addition, reasons for the variability in progression are not completely understood.⁴¹ Six retrospective cohort studies of HCV-infected adults with known time of infection (based on an identified exposure, often to contaminated blood products during young adulthood) reported cirrhosis in 0 to 10 percent of patients after at least 10 years of followup.^{29,42-48} Studies of community cohorts estimate cirrhosis in an average of 7 percent of persons after 20 years of HCV infection, with rates about twice as high in clinical and referral cohorts.^{38,49} One study of females infected by contaminated batches of anti-D immunoglobulin in 1980 found that approximately 14 percent of those who remained viremic had cirrhosis after 35 years.⁵⁰ Other studies suggest that progression to cirrhosis may accelerate after 20 years of chronic infection.^{47,51}

Mother-to-child (vertical) transmission is believed to be the main route of HCV infection acquisition in children. In a meta-analysis of the risk of vertical HCV infection, the pooled transmission rate was 5.8 percent among females with HCV monoinfection and 10.8 percent among those with HCV/HIV coinfection.⁵²

Risk Factors

HCV is primarily acquired via percutaneous exposures to infected blood. The strongest risk factor for HCV infection is injection drug use. The prevalence of HCV infection in PWID varies widely depending on age, duration of injection drug use, and other factors (such as availability and use of needle exchange programs).⁵³ Recent surveys of active PWID indicate that approximately one third of those aged 18 to 30 years are HCV-infected. Older PWID typically have a higher prevalence (approximately 70% to 90%) of HCV infection.²⁷ Although large population-based studies⁵⁴⁻⁵⁶ report independent associations between HCV infection and some high-risk sexual behaviors (multiple sexual partners, unprotected sex, and/or sex with a person infected with HCV infection or using injection drugs), the efficiency of transmission via sexual contact appears to be low; high-risk sexual behaviors may be a marker for unacknowledged drug use or other risk factors. Transfusions prior to 1992 are a risk factor for HCV infection but are no longer an important source of infection due to the implementation of effective screening programs for donated blood.^{57,58}

Rationale for Screening/Screening Strategies

Screening for HCV infection in asymptomatic adults who have no history of liver disease or known liver enzyme abnormalities may identify infected patients at earlier stages of disease, before they develop serious or irreversible liver damage. Studies estimate that around 50 percent (range 43 to 72%) of persons in the United States with chronic HCV infection are unaware of their status.^{18,57-60} Antiviral treatment, has become increasingly effective at achieving sustained aviremia (clearance of HCV infection). Screening for HCV infection might also help prevent transmission by decreasing high-risk injection drug use and other risky behaviors in those who test positive or through successful treatment of HCV,⁶¹ and could identify those who might benefit from hepatitis A or HBV vaccinations, alcohol cessation counseling, identification and management of extrahepatic manifestations, or other interventions. Screening is an important component of the National Academies of Sciences, Engineering, and Medicine report on eliminating HCV as a public health problem by the year 2030.⁶² Shorter-term goals of the National Viral Hepatitis Action Plan are to increase the proportion of persons aware of their positive HCV infection status to 66 percent and to decrease the number of HCV-related deaths by 25 percent by the year 2020.⁶³

Although prenatal HCV infection could identify infected females, a challenge is the lack of antiviral therapies proven to be effective for reducing risk of perinatal transmission and approved for use in pregnancy.¹ Older antiviral therapies were contraindicated in pregnancy due to teratogenic risks. Due to the lack of data on safety of newer DAA regimens during pregnancy and breastfeeding, clinical practice guidelines do not recommend antiviral therapy during pregnancy.^{64,65} However, even in the absence of antiviral therapy proven to be safe and effective during pregnancy, identification of HCV infection during pregnancy could facilitate decision making around the management and use of interventions during labor and delivery or in the perinatal period that might reduce risk of perinatal transmission, and identify females who could benefit from antiviral treatment later and infants who should be tested for HCV infection. A potential alternative strategy for preventing mother-to-child transmission is identification and treatment of HCV infection prior to pregnancy.²⁴

Interventions/Treatment

The goal of antiviral treatment for chronic HCV infection is to prevent the long-term health complications associated with HCV infection, such as cirrhosis, hepatic decompensation, and liver cancer. However, it is extremely difficult to design and carry out clinical trials long and large enough to provide direct evidence related to these outcomes. The SVR rate, typically defined as the proportion of patients who experience a decline in HCV RNA to undetectable levels 12 or 24 weeks following completion of antiviral treatment, is the standard marker of successful treatment in clinical trials. Most studies now focus on SVR at 12 weeks. Long-term recurrence of hepatitis C viremia occurs in less than one percent of patients with an SVR at 12 or 24 weeks of therapy; therefore, an SVR is considered equivalent to a cured infection.⁶⁶⁻⁶⁸ Studies have consistently found an association between achieving an SVR after antiviral therapy and reductions in mortality, liver failure, and cancer, though such analyses are susceptible to residual confounding.⁶⁹⁻⁷²

A major advance in antiviral treatment for HCV infection has been the development and adoption of all-oral DAA regimens without interferon. Such regimens are associated with substantially higher SVR rates than previous antiviral regimens, shorter duration of treatment (8 to 12 weeks instead of 24 to 48 weeks), and improved tolerability.⁷³ SVR rates with older antiviral regimens are shown in **Table 1**. DAA regimens are highly effective for HCV genotype 1 infection, the most common genotype in the United States and historically associated with lower SVR rates when treated with interferon-only regimens.

Given the rapid pace of development for HCV antiviral therapies, guidance for antiviral therapy for HCV is rapidly evolving (**Tables 2** and **3**).⁷⁴ Several newer DAA regimens are pangenotypic,⁷⁵ meaning that they are effective across all common genotypes, and most currently recommended regimens do not require use of ribavirin. Whereas antiviral therapy was previously reserved for patients with more advanced fibrosis, the American Association for the Study of Liver Disease (AASLD) and the Infectious Diseases Society of America (IDSA) now recommend treatment for all patients with chronic HCV, except those with short life expectancies that cannot be remediated by treating HCV, by transplantation, or by other directed therapy.⁶⁵ The U.S. Food and Drug Administration (FDA) recently approved three HCV regimens for children 12 to 17 years of age (**Table 4**).^{76,77} Although HCV antiviral therapy has traditionally most frequently been administered in specialty settings, studies have demonstrated similar SVR rates without any negative impacts on safety in community-based and primary care settings.^{78,79}

Recommendations regarding the diagnostic workup and pretreatment assessment for HCV are also evolving. Whereas liver biopsy was previously recommended in all patients with HCV infection in order to determine the severity of fibrosis, the AASLD-IDSA guideline currently also recommends blood tests or transient elastography as noninvasive options for fibrosis assessment.^{65,74,80,81} Given the availability of noninvasive tests to stage HCV infection, rates of biopsy have declined substantially, though precise data on current biopsy rates are lacking.

Current Clinical Practice/Recommendations of Other Groups

U.S.-based screening guidelines are summarized in **Table 5**. All are consistent in recommending HCV screening in persons born between 1945 and 1965 and in persons with risk factors for HCV infection. Data on rates of birth cohort screening are limited, though a study of U.S. veterans found an increased rate of testing in this age group compared with other age groups.⁸²

Guidelines from the European Association for the Study of the Liver (EASL)⁸³ and the World Health Organization (WHO)⁸⁴ are generally consistent with the above screening guidance. In 2017, the Canadian Task Force on Preventive Health Care recommended against screening for HCV in adults not at elevated risk (including persons born between 1945 and 1965 or other birth cohorts).⁸⁵ The Canadian recommendation was based on the reasoning that most persons with HCV infection have risk factors that can be identified using risk-based guidelines. However, the Canadian Association for the Study of the Liver recommends screening of high-risk persons and persons born between 1945 and 1975.⁸⁶

The CDC⁸⁷ and the American College of Obstetricians and Gynecologists⁸⁸ recommend offering HCV screening to pregnant people with risk factors.

Chapter 2. Methods

Key Questions and Analytic Framework

This systematic review followed a standard protocol in accordance with USPSTF procedures.⁸⁹ The scope and Key Questions (KQs) for this report were determined by the USPSTF and informed by evidence gaps identified from the prior reviews.^{1-3,5,90} Three additional contextual questions on recent epidemiologic trends in HCV infection, modeling analyses, and behavioral effects of current antiviral therapies were requested by the USPSTF. The KQs and Contextual Questions are shown below. Investigators created an analytic framework incorporating the KQs and outlining the patient populations, interventions, outcomes, and potential adverse effects, as well as the direct and indirect pathways from screening to health outcomes (**Figure 1**).

Key differences between this report and the prior reviews are inclusion of adolescents in addition to adults; evaluation of new all-oral, DAA regimens. We also removed previously reviewed questions on harms of liver biopsy, given its reduced role in evaluation of patients with HCV infection, and on effects of counseling or immunizations in persons with HCV infection, given limited evidence and likely small magnitude of effects relative to antiviral treatments. This report focuses on effects of treatments in populations more likely to be identified by screening (i.e., asymptomatic and without advanced liver disease), and excludes poor quality studies (e.g., cohort studies that did not perform statistical adjustment) that were included in prior USPSTF reviews. We did not re-review the diagnostic accuracy of HCV screening, which the prior review found to be highly accurate.⁹¹

Key Questions

- 1a. Does screening for HCV infection in pregnant and nonpregnant adolescents and adults without known abnormal liver enzyme levels reduce HCV-related mortality and morbidity or affect quality of life?
- 1b. Does prenatal screening for HCV infection reduce risk of vertical transmission of HCV infection?
- 2. What is the effectiveness of different risk- or prevalence-based methods for screening for HCV infection on clinical outcomes?
- 3. What is the yield (number of new diagnoses per tests performed) of one-time versus repeat screening or alternative screening strategies for HCV infection, and how does the screening yield vary in different risk groups?
- 4. What are the harms of screening for HCV infection (e.g., anxiety and labeling)?
- 5. What are the effects of interventions during labor and delivery or the perinatal period on risk of vertical transmission of HCV infection?
- 6. What is the effectiveness of currently recommended antiviral treatments in improving health outcomes in patients with HCV infection?*
- 7. What is the effectiveness of currently recommended antiviral treatments in achieving a SVR in patients with HCV infection?*
- 8. What are the harms of currently recommended antiviral treatments?*

9. What is the association between experiencing SVR following antiviral treatment and reduction in risk of HCV-related adverse health outcomes?

* Subpopulations of interest for KQs 6, 7, and 8 include those defined by age, race/ethnicity, sex, drug use, receipt of medications for treatment of opioid use disorder, stage of disease, HCV genotype, and pregnancy status (including nonpregnant women of childbearing age).

Contextual Questions

Three Contextual Questions were also requested by the USPSTF to help inform the report.

Contextual Questions are addressed by narratively summarizing key evidence; they are not reviewed using systematic review methodology.

- Based on population level estimates, what are recent trends in the epidemiology, prevalence, and incidence of HCV infection in the United States, including in primary care settings, over the past 5 to 10 years?
- What are the effects of different risk- or prevalence-based methods for screening for HCV infection in modeling studies?
- What is the effect of antiviral treatments on behavioral outcomes?

Search Strategies

We searched the Cochrane Central Register of Controlled Trials and Cochrane Database of Systematic Reviews (through February 8, 2019), and Ovid MEDLINE (1946 through February 8, 2019) for relevant studies. Search strategies are available in **Appendix A1**. We also searched ClinicalTrials.gov for ongoing studies, and reviewed the reference lists of relevant review articles and studies meeting inclusion criteria. We also carried forward studies in the prior USPSTF report that met inclusion criteria for this update.^{2,90}

Study Selection

Two reviewers independently evaluated each study to determine its inclusion eligibility based on predetermined inclusion and exclusion criteria developed for each KQ (**Appendix A2**).

The target population for screening was asymptomatic, pregnant and nonpregnant adolescents (ages 12 to 17 years) and adults without prior HCV infection. For treatment, the target population was persons with HCV infection likely to be identified by screening. However, no trial enrolled screen-detected patients, and trials did not report presence of symptoms. To evaluate patients more likely to be asymptomatic and identified by screening, we restricted inclusion of antiviral treatment studies to those in which up to 20 percent of participants had cirrhosis at baseline. For antiviral regimens with few studies meeting this threshold and for studies on the association between SVR after antiviral therapy and clinical outcomes, we

permitted a threshold up to 25 percent. We included studies of patients previously treated with interferon-based therapy (interferon or pegylated interferon with or without ribavirin) or boceprevir or telaprevir with pegylated interferon and ribavirin, because data indicate similar SVR rates in these treatment-experienced compared with treatment-naive patients.⁷³ Included interventions were HCV screening and alternative screening strategies; mode of delivery, labor management strategies, and breastfeeding practices; currently recommended (including alternative) DAA regimens for evaluation of clinical outcomes, SVR rates and harms; and DAA regimens or interferon-based treatment for evaluation of mortality and long-term clinical outcomes.⁷⁴ For analysis of SVR rates, we included studies in which ribavirin or dasabuvir was not used as recommended (e.g., ombitasvir / paritaprevir / ritonavir / dasabuvir that omitted ribavirin for genotype 1a infection or used ribavirin for genotype 1b infection, or did not include dasabuvir for genotype 1 infection) (**Tables 2** and **3**), because SVR rates were similar to recommended regimens with these variations, but performed sensitivity analyses without them.

For analysis of adverse events, we restricted inclusion to trials in which ribavirin was administered as recommended. DAA regimens were restricted to recommended doses and durations. We excluded trials that focused on persons coinfected with HIV or HBV infection, transplant patients, or with advanced renal disease.

For KQs on screening and treatment, we included randomized trials. For questions on screening, perinatal (labor and delivery or breastfeeding) interventions, effects of DAA regimens on clinical outcomes, and the association between SVR after antiviral therapy and clinical outcomes, we also included cohort studies that reported risk estimates adjusted for potential confounders. We included trials of current DAA regimens versus placebo, an older antiviral regimen, or another DAA regimen (including regimens not currently recommended). We also included trials of DAA regimens without one of these comparisons, because there were few comparative trials. Clinical trials were defined as studies in which patients were prospectively allocated to treatment by the study investigator using pre-defined inclusion criteria and followup methods. Included outcomes were mortality, morbidity (e.g., cirrhosis, hepatic decompensation, liver transplant, extrahepatic manifestations of HCV infection), quality of life, HCV transmission, harms (e.g., labeling, anxiety, drug-related and treatment-related harms), screening yield (number of new diagnoses per tests performed), and perinatal transmission. We restricted inclusion to English-language articles, and we excluded studies published only as abstracts. Studies of non-human subjects were excluded, and studies had to report original data. The selection of literature is summarized in the literature flow diagram (Appendix A3), and Appendix A4 provides a list of included studies. Appendix A5 lists excluded studies with reasons for exclusion.

Data Abstraction and Quality Rating

We constructed evidence tables summarizing the data from each study. One investigator abstracted details about the study design, patient population, setting, interventions, analysis, followup, and results. A second investigator reviewed abstracted data for accuracy. Two investigators independently applied criteria developed by the USPSTF⁸⁹ to rate the quality of each study as good, fair, or poor (**Appendix A6**). Discrepancies were resolved through a consensus process. In accordance with the USPSTF Procedure Manual, we excluded studies rated poor quality due to important methodological shortcomings that severely undermine their

reliability;⁸⁹ this applied to studies utilized in the prior USPSTF review that were rated poor quality and were excluded in the current report.

Data Synthesis

We performed a random effects meta-analysis to summarize the proportion of patients experiencing SVR and adverse events with current DAA regimens. We used a generalized linear mixed effects model with a logit link, allowing the inclusion of studies in which the proportion of patients with the event were 0 percent or 100 percent. We combined arms of comparable interventions within the same study so each study was represented once in a meta-analysis, in order to avoid overweighting. For SVR, we performed separate analyses for each genotype (1 through 6); for adverse events, results were pooled across genotypes. For SVR and adverse events, analyses were stratified according to DAA regimen. Subgroup and sensitivity analyses were performed on geographic settings (United States or Europe, multinational, or other), fibrosis stage (cirrhosis excluded or some [up to 20% of patients] with cirrhosis), prior treatment status (naïve or experienced to interferon-based therapies, boceprevir or telaprevir), and quality. For SVR, we performed sensitivity analysis by excluding studies in which ribavirin or dasabuvir was not used as recommended. For analyses of adverse events, we excluded trials of ribavirincontaining regimens except for ombitasvir / paritaprevir / ritonavir / dasabuvir with ribavirin, which is recommended for genotype 1b infection.

We also performed a random effects meta-analysis of adjusted hazard ratios (HRs) of SVR after antiviral therapy versus no SVR on clinical outcomes (mortality, liver-related mortality, cirrhosis, and HCC) using a linear mixed effects model. In some cases the adjusted HR for SVR versus no SVR had to be calculated from other estimates (e.g., from adjusted HRs for SVR and no SVR vs. no treatment). In these situations we calculated the adjusted HR for SVR versus no SVR based on the HRs for SVR versus no treatment and no SVR versus no treatment and their reported CIs, assuming a correlation of 0 between the two HRs. Because HRs are typically positively correlated, this assumption results in more conservative (i.e., wider) CIs for the calculated HR. Subgroup analysis were performed on duration of study (5 years or less vs. more than 5 years), geographic setting (United States/Europe vs. Asia) and whether the study had full adjustment of confounding variables (age, sex, fibrosis stage and genotype) or did not adjust for one or more of these populations. We also performed sensitivity analysis by excluding studies with potential overlapping populations in order to ensure that results were not sensitive to double counting of patients.

For all meta-analyses, statistical heterogeneity was assessed using the variance parameter of the random effects, the Cochran Q-test and *I*² statistic.⁹² For pooled proportions of SVR and adverse events, the Cochran Q-test and *I*² statistic were based on the Freeman-Tukey double arcsine transformed proportions.⁹³ All meta-analyses were conducted using SAS 9.4 (SAS Institute Inc., Cary, NC, USA.) and forest plots were created using Stata/SE 14.0 (StataCorp, College Station, TX).

We also conducted random effects meta-analysis on adverse events with DAA regimens versus placebo and DAA regimens versus telaprevir / pegylated interferon / ribavirin using RevMan 5.3.5 (the Nordic Cochrane Centre, Copenhagen). Analyses were stratified by DAA regimen.

There were too few trials evaluating these comparisons to conduct additional sensitivity or subgroup analyses.

We assessed the aggregate internal validity (quality) of the body of evidence for each KQ ("good", "fair", "poor") using methods developed by the USPSTF, based on the number, quality and size of studies, consistency of results between studies, and directness of evidence in the Summary of Evidence.⁸⁹ We determined aggregate internal validity using the totality of evidence (new studies identified for the update plus studies carried forward from the prior USPSTF report).

External Review

The draft research plan was posted for public comment and modified prior to finalization. The draft report will be reviewed by content experts, USPSTF members, Agency for Healthcare Research and Quality Medical Officers, and collaborative partners, and will be posted for public comment.

Chapter 3. Results

A total of 7,170 new references from electronic database searches and manual searches of recently published studies were reviewed, and 700 full-text papers were evaluated for inclusion. We included a total of 97 studies (reported in 94 publications). Eighty-four studies were newly identified as part of this update, and 13 were carried forward from the previous review. Included studies and quality ratings are described in **Appendix B**.

Key Question 1a. Does Screening for HCV Infection in Pregnant and Nonpregnant Adolescents and Adults Without Known Abnormal Liver Enzyme Levels Reduce HCV-Related Mortality and Morbidity or Affect Quality of Life?

As in the prior USPSTF review, no study directly assessed effects of HCV screening versus no screening on clinical outcomes such as HCV-related mortality and morbidity or quality of life.

Key Question 1b. Does Prenatal Screening for HCV Infection Reduce Risk of Vertical Transmission of HCV Infection?

As in the prior USPSTF review, no study assessed effects of prenatal HCV screening versus no screening on risk of vertical transmission of HCV infection.

Key Question 2. What Is the Effectiveness of Different Riskor Prevalence-Based Methods for Screening for HCV Infection on Clinical Outcomes?

As in the prior USPSTF review, no study directly assessed the effectiveness of different risk- or prevalence-based methods for HCV screening on clinical outcomes.

Key Question 3. What Is the Yield (Number of New Diagnoses per Tests Performed) of One-Time Versus Repeat Screening or Alternative Screening Strategies for HCV Infection, and How Does the Screening Yield Vary in Different Risk Groups?

Summary

• The prior USPSTF review included five studies that found screening strategies that targeted multiple risk factors associated with sensitivities of more than 90 percent and numbers needed to screen to identify one case of HCV infection of less than 20. More

narrowly targeted screening strategies were associated with numbers needed to screen of less than two, but missed up to two-thirds of infected patients.

• One new study found that applying risk-based guidelines perfectly would result in 24.7 percent of the population tested and 82 percent of HCV cases identified (number needed to screen 14.6), compared with 45 percent of the population tested and 76 percent of HCV cases identified with birth cohort screening (number needed to screen 28.7), but assumed perfect implementation of risk-based testing.

Evidence

The prior USPSTF review included five poor quality studies⁹⁴⁻⁹⁸ that found screening strategies that targeted multiple risk factors associated with sensitivities of more than 90 percent and numbers needed to screen to identify one case of HCV infection of less than 20.² More narrowly targeted screening strategies were associated with numbers needed to screen of less than two, but missed up to two-thirds of infected patients.

One new study that retrospectively applied screening criteria to patients in the 2003 to 2006 National Health and Nutrition Examination Survey (NHANES) database compared the yield of risk-based HCV screening (based on then-current AASLD guidelines) versus birth cohort screening.⁹⁹ It found that applying risk-based guidelines perfectly would result in 24.7 percent of the general population tested and identify 82 percent of the HCV exposed population, with a number needed to screen to identify one case of HCV infection of 14.6. Applying the birth cohort strategy would result in 45 percent of the general population tested and identify 76 percent of the HCV exposed population, with a number needed to screen to identify one case of 28.7. Although this analysis suggests that the two strategies would identify a similar proportion of HCV infected persons, it would require perfect implementation of risk-based testing, which has not occurred in actual practice.

No study evaluated the yield of one-time versus repeat screening, the yield of alternative screening strategies in different risk groups, or the yield of currently recommended screening (i.e., 1945 to 1965 birth cohort plus risk-based screening) versus expanded screening strategies. Studies that modeled effects of alternative screening strategies are addressed in Contextual Question 2.

Key Question 4. What Are the Harms of Screening for HCV Infection (e.g., Anxiety and Labeling)?

The prior USPSTF review included five studies^{31,100-103} of persons with HCV infection that suggested potential negative psychological and social effects of screening, but the quality of evidence was assessed as poor due to small sample sizes and methodological shortcomings, included no unscreened comparison group, reliance on retrospective recall, and poorly defined outcomes.² All of the studies were conducted in the context of treatment with older interferon-containing regimens. No new study meeting inclusion criteria evaluated harms associated with HCV screening.

Key Question 5. What Are the Effects of Interventions During Labor and Delivery or the Perinatal Period on Risk of Vertical Transmission of HCV Infection?

Summary

- Five observational studies (four included in the prior USPSTF review) found no clear association between the mode of delivery and risk of mother-to-infant transmission of HCV infection, after adjustment for potential confounders.
- One observational study included in the prior USPSTF review found prolonged (longer than 6 hours) rupture of membranes associated with increased risk for HCV transmission versus less prolonged (6 hours or less) rupture after adjusting for maternal demographic characteristics, HCV RNA level, intravenous drug use, and smoking status during pregnancy (adjusted OR 9.3, 95% CI, 1.5 to 180).¹⁰⁴ No new study evaluated this association.
- One observational study included in the prior USPSTF review found internal fetal monitoring associated with increased risk of mother-to-infant transmission of HCV infection versus external monitoring, after adjustment for maternal demographic characteristics, HCV viral load, intravenous drug use history, and smoking status in pregnancy (adjusted OR 6.7, 95% CI, 1.1 to 35.9).¹⁰⁴ No new study evaluated this association.
- Three observational studies (two included in the prior USPSTF review) found no clear association between breastfeeding and risk of mother-to-infant transmission of HCV infection after adjustment for potential confounders; in the two good quality studies adjusted OR estimates were close to 1.¹⁰⁵⁻¹⁰⁷

Evidence

Mode of Delivery

The prior USPSTF review² included 14 observational studies in 16 publications (sample sizes of 56 to 1,034 mother-infant pairs) that found no clear association between the mode of delivery (vaginal vs. cesarean delivery) and risk of mother-to-infant transmission of HCV.^{104-106,108-120} Twelve studies found no statistically significant association between the mode of delivery and risk of HCV transmission;^{104-106,109-112,114-120} most estimates were imprecise, and findings were inconsistent, with point estimates that favored vaginal delivery in some studies and cesarean delivery in others. Most of the studies included in the prior review did not meet inclusion criteria for the current review: eight were rated poor quality^{109,111-113,116-120} and ten did not conduct multivariate analyses.¹⁰⁹⁻¹²⁰ No study reported baseline characteristics according to mode of delivery or matched women on key potential confounders.

Restricting inclusion to the four studies (total 1,717 mother-infant pairs) in the prior review that met current inclusion criteria (fair or good quality and multivariate analysis performed) resulted in a similar conclusion of no clear association between the mode of delivery and risk of HCV transmission (**Table 6**; **Appendix B Table 1**).^{104-106,108} One of the studies was conducted in the

United States¹⁰⁴ and the other three in Europe. Although one fair quality study (424 motherinfant pairs) found elective cesarean associated with decreased risk of HCV transmission versus vaginal delivery or emergent (after onset of labor) cesarean after adjusting for HIV status and breastfeeding (adjusted OR 0.0, 95% CI, 0.0 to 0.87),¹⁰⁵ the other three studies, including two good quality studies,^{104,106} found no association between the mode of delivery and HCV transmission risk. One good quality study (1,034 mother-infant pairs) found no statistically significant association between the mode of delivery and risk of HCV transmission, though there was a trend towards higher risk with elective cesarean versus vaginal or emergent (after onset of labor) cesarean, after adjusting for infant sex, prematurity, and breastfeeding status (adjusted OR 1.59, 95% CI, 0.88 to 2.86),¹⁰⁶ and another good quality study (181 mother-infant pairs) found no association between the mode of delivery (elective cesarean, emergent cesarean or vaginal) and risk of mother-to-infant transmission in univariate analysis; mode of delivery was excluded from the multivariate model.¹⁰⁴ The fourth, fair quality study (78 mother-infant pairs) found no association between cesarean (not specified as elective or emergent) versus vaginal delivery and risk of transmission (data not reported).¹⁰⁸

One additional Italian study (1,301 mother-infant pairs) not included in the prior USPSTF review also found no statistically significant association between the mode of delivery (cesarean vs. vaginal delivery) and risk of mother-to-infant transmission of HCV infection (adjusted OR 0.83, 95% CI, 0.65 to 1.08). Cesarean deliveries were not specified as elective or emergent¹⁰⁷ (**Table 6; Appendix B Tables 1-3**). The study was rated good quality (**Table 6; Appendix B Table 4**).

Rupture of Membranes

Evidence on the association between duration of rupture of membranes during labor and risk of HCV transmission is limited. The prior USPSTF review included one good quality United States cohort study (189 mother-infant pairs) that found prolonged rupture (longer than 6 hours) of membranes associated with increased risk for HCV transmission versus less prolonged rupture (6 hours or less) after adjusting for maternal demographic characteristics, HCV RNA level, intravenous drug use, and smoking status during pregnancy (adjusted OR 9.3, 95% CI, 1.5 to 180)¹⁰⁴ (**Table 7**; **Appendix B Tables 1-3**). However, there were only 7 cases of perinatal HCV infection, and the estimate was very imprecise. A smaller (63 mother-infant pairs) Australian study¹¹⁶ included in the prior USPSTF review found that mean duration of membrane rupture was longer in mothers in whom HCV transmission occurred compared with those in whom transmission did not occur, but did not meet current inclusion criteria because it did not attempt to adjust for potential confounders and was rated poor quality. We identified no new studies on the association between the duration of rupture of membranes and risk of HCV transmission that met inclusion criteria.

Fetal Monitoring

Evidence on the association between use of fetal monitoring methods during labor and risk of HCV transmission is limited. The prior USPSTF review included one good quality U.S.-based study (188 mother-infant pairs) that found internal fetal monitoring associated with increased risk of mother-to-infant transmission of HCV infection versus external monitoring, after adjustment for maternal demographic characteristics, HCV viral load, intravenous drug use history, and smoking status in pregnancy (adjusted OR 6.7, 95% CI, 1.1 to 35.9)¹⁰⁴ (**Table 8**;

Appendix B Tables 1-3). However, there were only 7 cases of perinatal HCV infection and the estimate was imprecise. Although the prior USPSTF review included two other studies on the association between fetal monitoring and risk of HCV transmission, neither met current inclusion criteria because they did not report adjusted risk estimates.^{112,114} One of the studies¹¹² did not compare internal fetal monitoring to no internal monitoring and the other study¹¹⁴ found no association between internal fetal monitoring and transmission risk (relative risk [RR] 1.24, 95% CI, 0.70 to 2.2). We identified no new studies on the association between the use of fetal monitoring methods and risk of HCV transmission that met inclusion criteria.

Breastfeeding

The prior USPSTF review² included 14 observational studies^{104-106,109,111,115-124} (total of 2,971 mother-infant pairs) that found no association between breastfeeding by women infected with HCV and risk of transmission to infants. No study reported a statistically significant association, though some estimates were very imprecise due to few cases of HCV transmission. Most of the studies included in the prior review did not meet inclusion criteria for the current review: ten were rated poor quality,^{108-114,116-120} and twelve did not conduct multivariate analyses.^{104,108-120}

Restricting the analysis to the two studies^{105,106} in the prior review that meet current inclusion criteria (fair or good quality and multivariate analysis performed) resulted in a similar conclusion of no association between breastfeeding and risk of HCV transmission (**Table 9**; **Appendix B Tables 1-3**).^{104-106,108} One large (1,034 mother-infant pairs) European study found no association between breastfeeding by HCV-infected women without HIV infection and risk of HCV transmission to infants (followed until at least 18 months of age), after adjusting for infant sex, prematurity, and mode of delivery (adjusted OR 0.92, 95% CI, 0.50 to 1.70). A fair quality European study (414 mother-infant pairs) also found no association between breastfeeding and risk of HCV transmission to infants (duration of followup 24 months), after adjusting for HIV status (5% of mothers were HIV-infected) and mode of delivery (adjusted OR 1.52, 95% CI, 0.35 to 5.12). Although the point estimate was consistent with increased risk associated with breastfeeding, the estimate was imprecise.

One additional good quality Italian cohort study¹⁰⁷ (1,281 mother-infant pairs) not included in the prior systematic review also found no association between breastfeeding and risk of HCV transmission to infants, after adjusting for maternal HCV viral load, HIV status (14% of mothers were HIV-infected), injection drug use, and mode of delivery (adjusted OR 0.95, 95% CI, 0.58 to 1.40) (**Table 9**; **Appendix B Tables 1-4**). Duration of followup was 24 months.

Key Question 6. What Is the Effectiveness of Currently Recommended Antiviral Treatments in Improving Health Outcomes in Patients With HCV Infection?

Summary

Adults

- The prior review included no randomized trials or observational studies on the effects of then-current antiviral regimens on long-term (e.g., more than 2 years) clinical outcomes; no new randomized trial evaluated effects of current DAA regimens on long-term clinical outcomes.
- Ten new trials reported quality of life and functional outcomes before and after treatment with a current DAA regimen.
 - A pooled analysis of four trials found sofosbuvir / velpatasvir associated with an average improvement of 5.5 to 6.1 points (0 to 100 scale) on 26 measures related to quality of life or function at 24 weeks (12 weeks post-treatment) in persons without cirrhosis.
 - A pooled analysis of three trials found sofosbuvir / ledipasvir associated with small but statistically significant improvements from baseline to 24 weeks (12 weeks post-treatment) on multiple quality of life and functional domains in persons with no to mild fibrosis at baseline.
 - Three trials of DAA regimens not included in the pooled analyses (two trials of ombitasvir / paritaprevir / ritonavir / dasabuvir and one trial of elbasvir / grazoprevir) found DAA use associated with small changes from baseline to 12 weeks post-treatment on the 36-Item Short Form Health Survey (SF-36) physical (improvement 0.5 to 1.4 points) or mental component (improvement 2.5 to 3.0 points) summary scales (0 to 100 scale).
- Thirty-one trials reported mortality 12 to 36 weeks following completion of therapy with a DAA regimen. Twenty-one trials reported no deaths; in the other ten trials, there were 17 deaths (0.4% [17/3,848] overall).
- Three large (n=34,206; 17,836; and 6,850) cohort studies evaluated the association between use of DAA regimens, interferon-based treatment, and no antiviral therapy and risk of cardiovascular events and HCC.
 - One retrospective study (n=34,206) found DAA therapy and interferon-based therapy each associated with similarly decreased risk of cardiovascular events relative to no therapy (incidence per 1,000 person-years 16.3 for DAA therapy, 23.5 for interferon-based therapy, and 30.4 for no therapy; p<0.001 for DAA therapy or interferon-based therapy vs. no therapy).
 - One study (n=17,836) found no difference between interferon-based treatment versus DAA therapy in risk of HCC (incidence rate per 1,000 person-years of followup 7.48 vs. 7.92; p=0.72); both regimens were associated with lower incidence of HCC than no therapy.
 - One study (n=6,850) found no difference between DAA therapy versus no antiviral therapy and risk of HCC (adjusted HR 1.02, 95% CI, 0.40 to 2.61)

among persons without known cirrhosis at baseline after 33 months followup; effects on all-cause mortality favored DAA therapy, but the difference was not statistically significant (adjusted HR 0.74, 95% CI, 0.43 to 1.28).

Adolescents

- Three trials of DAA therapy in adolescents found quality of life improved from baseline based on Pediatric Quality of Life Inventory scores.
- Three short-term trials of DAA regimens in adolescents reported no deaths.

Evidence

Adults

The prior review identified no randomized trials or observational studies on the effects of thencurrent antiviral regimens (triple therapy with telaprevir or boceprevir, pegylated interferon, and ribavirin or dual therapy with pegylated interferon and ribavirin) for chronic HCV infection on long-term (more than 2 year) clinical outcomes.^{5,90} Two trials in the prior review reported shortterm mortality with triple therapy versus dual therapy, but events were few and estimates were imprecise, with no clear differences.^{125,126} There were a total of 9 deaths in over 1,700 persons across the two trials.

No new randomized trial evaluated effects of current DAA regimens on long-term clinical outcomes. Randomized trials of older (non-DAA) antiviral therapy versus no antiviral therapy that evaluated long-term clinical outcomes did not meet inclusion criteria because they enrolled persons with cirrhosis at baseline,¹²⁷⁻¹³² utilized non-standard therapy (indefinite treatment with interferon),¹³³ or were rated poor quality (not clearly randomized).¹³⁴

Ten trials reported quality of life and functional outcomes before and after receipt of current DAA regimens; seven trials were included in two pooled analyses^{135,136} and three additional trials (reported in 2 publications) not in the pooled analyses also reported these outcomes (**Appendix B Tables 5, 10, and 11**).^{137,138} One trial of sofosbuvir / velpatasvir that reported quality of life and functional outcomes was included in a pooled analysis and is not reported separately here.^{139,140} The trials were all open-label and none reported comparisons of DAA therapy versus placebo or non-DAA therapy.

Thirty-one trials (in 28 publications)^{139,141-167} reported short-term mortality with current DAA regimens (**Appendix B Tables 10 and 11**). A multicenter prospective cohort study conducted in France¹⁶⁸ and two retrospective cohort studies^{169,170} based on a national Veterans Affairs (VA) database, Electronically Retrieved Cohort of HCV Infected Veterans (ERCHIVES), evaluated the association between treatment with a DAA regimen versus no treatment and other clinical outcomes (cardiovascular outcomes and HCC) after adjusting for potential confounders (**Appendix B Table 5**).

Quality of Life and Function

Ten trials reported quality of life and functional outcomes before and after treatment with a current DAA regimen (**Appendix B Tables 5, 10, and 11**). Seven trials were included in two post-hoc pooled analyses: one analysis¹³⁵ included three trials (n=1,005) of sofosbuvir / ledipasvir and one analysis¹³⁶ included four trials (n=1,701) of sofosbuvir / velpatasvir. The trials varied with regard to whether antiviral therapy was administered with or without ribavirin. Two additional trials (reported in 1 publication, n=309 and 148) of ombitasvir / paritaprevir / ritonavir / dasabuvir (with or without ribavirin)¹³⁷ and one additional trial of elbasvir / grazoprevir (n=129) also reported quality of life or function.¹³⁸ All studies used an open-label design, and the quality of life and functional measures assessed in the trials differed. In addition, the trials included in the pooled analyses lacked a non-DAA regimen comparison group.

A pooled analysis of four trials found sofosbuvir / velpatasvir associated with an average improvement of 5.5 to 6.1 points on 26 measures related to quality of life or function at 24 weeks (12 weeks post-treatment) in persons without cirrhosis.¹³⁶ Changes from baseline were not statistically significant. Findings were similar when the regimen was administered with or without ribavirin. The average improvement was based on 26 outcomes derived from the SF-36, the Functional Assessment of Chronic Illness Therapy-Fatigue (FACIT-F), the Chronic Liver Disease Questionnaire-HCV version (CLDQ-HCV), and the Work Productivity Activity Index: Specific Health Problem (WPAI-SHP) measures, standardized to a 0 to 100 scale.

A pooled analysis of three trials found sofosbuvir / ledipasvir associated with statistically significant improvements from baseline to 24 weeks (12 weeks post-treatment) on multiple quality of life and functional domains in persons with no to mild fibrosis at baseline.¹³⁵ Estimates were similar when sofosbuvir / ledipasvir was administered with or without ribavirin. Mean differences were less than 3 points on the 0 to 100 SF-36 physical and mental component summary scales, 10 to 11 points on the 0 to 160 FACIT-F scale, 0.5 to 0.6 points on the CLDQ-HCV, less than 0.1 point on the 0 to 1 WPAI-SHP scales, and 0.04 to 0.05 points on the six-dimensional health state short-form (SF-6D) health utility scale; the latter measure was derived from the SF-36 instrument.

Three trials not included in pooled analyses also reported small improvements in some measures of quality of life or function.^{137,138} Two trials found ombitasvir / paritaprevir / ritonavir / dasabuvir associated with small changes from baseline to 12 weeks post-treatment on the SF-36 physical (improvement 0.5 to 1.4 points) or mental component (improvement 2.5 to 3.0 points) summary scales.¹³⁷ Estimates were similar when the regimen was administered with or without ribavirin and among treatment-naïve and -experienced patients. In both trials, there were no statistically significant differences between the DAA regimen versus telaprevir / pegylated interferon / ribavirin on the SF-36 (differences -1.1 to -1.5 points on the mental component and -1.3 to +0.9 points on the physical component summary scales). Changes from baseline following treatment with ombitasvir / paritaprevir / ritonavir / dasabuvir on the WPAI-SHP scale were also very small. Another trial found elbasvir / grazoprevir use associated with small but statistically significant improvements from baseline in SF-36 mental and physical component scores (mean change of 2 points each).¹³⁸ There was no effect of elbasvir / grazoprevir on patient fatigue, based on FACIT-F scale score.

Mortality

Thirty-one trials (in 28 publications; n=21 to 558; total N=3,848) reported mortality 12 to 36 weeks following completion of therapy with a DAA regimen (**Appendix B Tables 10 and 11**).^{139,141-167} The trials were not designed or powered to assess mortality, and 21 studies reported no deaths. There were 17 deaths in the remaining ten studies (0.4% overall). The regimens evaluated in these trials were sofosbuvir / velpatasvir (8 deaths in 884 patients; 0.9%),^{139,146,147,150} ombitasvir / paritaprevir / ritonavir / dasabuvir with ribavirin (4 deaths in 187 patients; 2%),^{149,162} grazoprevir / elbasvir (2 deaths in 732 patients; 0.3%),^{164,166} glecaprevir / pibrentasvir (2 deaths in 1,172 patients; 0.2%),¹⁶⁷ and sofosbuvir / daclatasvir (one death in 115 patients; 0.9%).¹⁶⁷ Ten of the 17 deaths were reported in three trials that enrolled persons reporting recent injection drug use (26% to 66% at baseline) or use of opioid substitution therapy (3% to 85% at baseline).^{149,150,167}

Other Clinical Outcomes

Three large, fair-quality cohort studies evaluated the association between antiviral treatment versus no treatment and clinical outcomes (cardiovascular events, HCC, or all-cause mortality).¹⁶⁸⁻¹⁷⁰ Two studies^{169,170} were conducted using the VA ERCHIVES database, and one study¹⁶⁸ was conducted in France.

Two large (n=17.836 and 34.206), retrospective analyses of VA patients evaluated the association between use of DAA regimens, interferon-based treatment, and no antiviral therapy and risk of cardiovascular events and HCC (Appendix B Tables 5 and 6).^{169,170} The studies included primarily male (3 to 4% female), HCV-infected veterans. Mean age ranged from 54 to 62 years; approximately 20 percent of the population had cirrhosis at baseline. One study found DAA therapy and interferon-based therapy each associated with decreased risk of cardiovascular events, including acute myocardial infarction, congestive heart failure, and stroke (incidence rate per 1,000 person-years of followup: 16.3 for DAA therapy, 23.5 for interferon-based therapy, and 30.4 for no therapy; p<0.001 for DAA therapy vs. no therapy and for interferon-based therapy vs. no therapy).¹⁶⁹ The proportion of patients with at least 5 years followup was 82% for interferon-based therapy, 3.7% for DAA therapy, and 43% for no therapy (mean followup not reported). The other study found no difference between interferon-based treatment versus DAA therapy in risk of HCC (incidence rate per 1,000 person-years of followup 7.48 vs. 7.92; p=0.72).¹⁷⁰ Both types of antiviral therapy regimens were associated with lower incidence of HCC than no therapy (incidence rate per 1,000 person years 10.90). The mean duration of followup was 7.4 years for persons treated with interferon-based therapy and 1.1 years for persons treated with DAA therapy (mean not reported for untreated patients).

A third, smaller (n=6,850) study conducted in France found no difference between DAA therapy versus no antiviral therapy in risk of HCC (adjusted HR 1.02, 95% CI, 0.40 to 2.61) in persons not known to have cirrhosis at baseline after a median of 33 months followup.¹⁶⁸ Effects on all-cause mortality favored DAA therapy, but the difference was not statistically significant (adjusted HR 0.74, 95% CI, 0.43 to 1.28). There were too few events to estimate effects on liver-related mortality or decompensated cirrhosis. Some differences between this analysis and the VA studies described above include availability of results for the subgroup of persons without cirrhosis at baseline, a much higher proportion of female patients (approximately 50%),

restriction to DAA therapy, prospective design, and similar duration of followup in treated and untreated patients.

No study evaluated effects of treatment with DAA regimens on risk of HCV transmission.

Adolescents

Data on health outcomes associated with DAA regimens in adolescents is available from one fair quality, open-label trial¹⁷¹ and post-hoc, before-after analyses of two other fair quality trials (Appendix B Tables 7 and 8).^{172,173} The studies included a total of 200 patients, mean age was 14 to 15 years, the proportion of females ranged from 40 to 63 percent, and patients did not have known cirrhosis. The studies utilized ledipasvir and sofosbuvir in adolescents with genotype 1 infection,¹⁷² sofosbuvir and ribavirin in adolescents with genotype 2 or 3 infection,¹⁷³ and glecaprevir / pibrentasvir in patients with genotype 1, 2, 3 or 4 infection.¹⁷¹ Quality of life was assessed based on change from baseline on the Pediatric Ouality of Life Inventory.¹⁷⁴ The Pediatric Quality of Life Inventory comprises four domains: Physical, Emotional, Social and School Functioning, and the total score is determined by averaging the scores from each of the four domains. In adolescents with genotype 1 infection treated with ledipasvir and sofosbuvir, caregiver-reported total quality of life scores were significantly improved from baseline at 24 weeks post-treatment (0-100 scale; mean change 5.2 points; p=0.009). However, there was no significant change in patients' self-reported total scores (mean change 1.9 points; p=0.12). Only the Emotional Functioning domain was rated as significantly improved from baseline by both caregivers (mean change 9.32 points, p<0.001) and patients (mean change 3.66, p=0.04).¹⁷² In adolescents with genotype 2 or 3 infection treated with sofosbuvir and ribavirin, scores improved on the self-reported Social Functioning score by 4.8 points (p=0.02) and on the parent-proxyreported School Functioning score by 13.0 points (p=0.0065). Adolescents treated with glecaprevir / pibrentasvir also experienced a small improvement in total quality of life score (mean change 2.3 points) though the statistical significance (p-value not reported) and timing of the assessment in this study is unclear.

Three studies of DAA regimens (sample sizes 30 to 100; total N=182) reported no deaths, but were not designed to assess long-term clinical outcomes (duration of followup \leq 48 weeks; **Appendix B Tables 7 and 8**). Two of the studies evaluated DAA regimens FDA-approved for use in adolescents (ledipasvir and sofosbuvir¹⁷⁵ and sofosbuvir and ribavirin¹⁷³) and one study evaluated a DAA regimen currently recommended for use in adults but not FDA-approved for use in adolescents (sofosbuvir and daclatasvir¹⁷⁶).

Key Question 7. What Is the Effectiveness of Currently Recommended Antiviral Treatments in Achieving an SVR in Patients With HCV Infection?

Summary

Adults

- The prior review found triple therapy with telaprevir or boceprevir associated with higher likelihood of SVR than dual therapy with pegylated interferon and ribavirin in persons with genotype 1 infection. SVR rates were 68 percent to 72 percent with triple therapy and 38 percent to 46 percent with dual therapy.
- One new randomized trial found sofosbuvir / velpatasvir associated with very high likelihood of SVR versus placebo in persons with mixed genotype (1, 2, 4, 5, or 6) infection (99% vs. 0%, RR 231.6, 95% CI, 14.6 to 3,680).¹³⁹ Across genotypes, the SVR rate with sofosbuvir / velpatasvir ranged from 97 percent to 100 percent.
- Two new randomized trials found ombitasvir / paritaprevir / ritonavir / dasabuvir (with or without ribavirin) associated with increased likelihood of SVR versus telaprevir / pegylated interferon / ribavirin in persons with genotype 1 infection who were treatment-naïve (98% vs. 80%, RR 1.22, 95% CI, 1.09 to 1.37) or who had previously received interferon therapy (99% vs. 66%, RR 1.50, 95% CI, 1.22 to 1.85).¹³⁷
- Forty-nine new trials found current DAA regimens associated with pooled SVR rates that ranged from 95.5 percent to 98.9 percent:
 - Genotype 1 infection (32 trials): Pooled SVR 97.7 percent (95% CI, 96.6% to 98.4%, I²=82%)
 - Genotype 2 infection (5 trials): Pooled SVR 98.9 percent (95% CI, 97.5% to 99.5%, I²=4%)
 - Genotype 3 infection (6 trials): Pooled SVR 95.5 percent (95% CI, 91.6% to 97.7%; I²=66%)
 - Genotype 4 infection (10 trials): Pooled SVR 98.2 percent (95% CI, 94.7% to 99.4%; I²=50%)
 - Genotype 5 infection (4 trials): Pooled SVR 96.0 percent (95% CI, 88.3% to 98.7%; I²=0%)
 - Genotype 6 infection (5 trials): Pooled SVR 98.2 percent (95% CI, 95.4% to 99.3%, I²=0%).
 - Mixed genotype 1 to 6 (2 trials): Pooled SVR 95.4% (95% CI, 89.4% to 98.1%; $I^2=0\%$).
- SVR estimates were consistent in analyses stratified by DAA regimen, study quality, inclusion of persons with cirrhosis at baseline, and geographic setting; and when analyses were restricted to trials that utilized ribavirin as recommended or to treatment-naïve patients.
- SVR estimates were similar in trials that stratified patients according to age (17 trials, primarily using a 55- or 65-year threshold), sex (17 trials), race or ethnicity (11 trials), or treatment-experience (five trials).

Adolescents

- Seven new trials (total N=348) reported SVR rates of 97 percent to 100 percent with DAA regimens in adolescents with HCV infection.
 - Four trials evaluated DAA regimens currently recommended and FDA-approved for use in adolescents (ledipasvir / sofosbuvir, sofosbuvir / ribavirin or glecaprevir / pibrentasvir) and three trials evaluated DAA regimens currently recommended for adults but not FDA-approved for use in adolescents.
 - Results were consistent across genotypes and in treatment-naïve and -experienced patients.

Evidence

Adults

The prior review found higher SVR rates in persons with HCV genotype 1 infection treated with triple therapy with telaprevir or boceprevir plus pegylated interferon and ribavirin than with dual therapy with pegylated interferon and ribavirin.^{5,90} Findings were consistent for a 48-week boceprevir regimen (2 trials, SVR rates 70% vs. 38%, RR 1.8, 95% CI, 1.6 to 2.1),^{126,177} a 24-week, fixed-duration telaprevir regimen (3 trials, SVR rates 68% vs. 46%, RR 1.5, 95% CI, 1.3 to 1.8),¹⁷⁸⁻¹⁸⁰ and a 24- or 48-week, response-guided telaprevir regimen (1 trial, SVR rate 72% vs. 44%, RR 1.6, 95% CI, 1.4 to 1.9).¹²⁵ The prior review also included 5 trials of dual therapy with pegylated interferon and ribavirin for genotype 2 or 3 infection that reported pooled SVR rates of 78 percent (95% CI, 67% to 88%) for 24 weeks of treatment and 68 percent (56% to 78%) for 12 to 16 weeks of therapy.¹⁸¹⁻¹⁸⁴ None of the studies in the prior review evaluated current DAA regimens.

Forty-nine new trials (in 44 publications) reported effects of current DAA treatment regimens on SVR in patients with HCV infection (Table 10; Appendix B Tables 10 and 11).^{137,139,141-167,185-} ¹⁹⁹ Sample sizes ranged from 10 to 706 (total N=9,917), mean age ranged from 45 to 68 years, and the proportion of female participants ranged from 18 to 64 percent. Twenty-four trials (in 20 publications) were multinational (primarily United States, Australia and/or Europe),^{137,139,143,144,149,150,155,158,160,164,166,167,185-189,191,196,198} 11 (in 10 publications) were conducted in the United States and/or Canada,^{146,147,153,154,157,161,190,192-194} eight in Asia,^{145,151,152,156,163,165,197,199} two in France,^{141,142} two in Egypt,^{162,195} and one each in Brazil,¹⁵⁹ and New Zealand.¹⁴⁸ The eight trials conducted in Asia did not report race. In the other studies, among those that reported race, the majority of participants were white (range 60 to 100%^{139,141,142,146,147,153-155,157,158,160-162,166,185-188,190-194} with the exception of one study conducted in New Zealand in which 16 percent of participants were white¹⁴⁸ and one study conducted primarily in Asian countries in which 28% of participants were white.¹⁶⁴ Twenty-one trials (in 19 publications) enrolled patients with genotype 1 infection, ^{137,145,149,151-156,159-161,163,167,185-188,190-} ^{194,197} one trial genotype 2,^{147,199} three trials genotype 3,^{147,157,158,167} three trials genotype 4, 141,162,189,195,200 one trial each for genotypes $5^{142,143}$ and 6, 143,148 and nine trials mixed genotypes (three trials genotypes 1 through 6;^{146,150,165} one trial genotypes 1, 2, 4 and 6;¹³⁹ two trials genotypes 2 through 6;^{144,196} two trials genotypes 1, 4 and 6;^{166,198} and one trial genotypes 1 and 4¹⁶⁴). Thirty-one trials (in 28 publications) excluded patients with cirrhosis^{137,144,146,147,154,155,159-} 162,167,186,188-194,196,197,199 or reported results in the subgroup of patients without

cirrhosis.^{139,147,149,150,164,165,185,198} For trials that enrolled patients with cirrhosis, inclusion was restricted to trials in which the proportion of patients with cirrhosis was less than 20 percent, with the exception of one trial of grazoprevir / elbasvir that had a slightly higher proportion (22%).¹⁶⁶ All trials excluded patients with HBV infection. Five trials (in 4 publications) enrolled patients with a history of receiving methadone or buprenorphine for opioid use disorder.^{149,150,167,192} The other trials excluded patients with recent or current substance use or did not describe substance use.

Thirteen trials (in 11 publications) evaluated ombitasvir / paritaprevir / ritonavir / dasabuvir, with or without ribavirin, ^{137,149,151,155,162,186-189,191,192} ten trials ledipasvir / sofosbuvir,^{141,142,145,148,156,163,185,190,193,195} eight trials (in 6 publications) glecaprevir / pibrentasvir,^{143,167,194,196,197,199} seven trials (in 6 publications) sofosbuvir / velpatasvir,^{139,146,147,150,158,165} six trials elbasvir / grazoprevir,^{144,152,160,164,166,198} four trials daclatasvir / sofosbuvir.^{157,159,161,167} and three trials simeprevir / sofosbuvir.^{153,154,159} One trial compared a current DAA regimen versus placebo,¹³⁹ two trials (reported in one publication) compared a current DAA regimen versus a regimen with telaprevir,¹³⁷ and two trials (reported in one publication) compared a current DAA regimen versus an older, not currently recommended, DAA regimen.¹⁴⁷ Five other trials randomized patients to a DAA regimen versus placebo with delayed DAA therapy, but only reported SVR rates following active treatment.^{151,152,164,166,187} The other trials did not compare a current DAA regimen to placebo or an older antiviral regimen. The duration of treatment was 12 weeks in all trials except for seven trials (in 5 publications)^{143,167,196,197,199} which evaluated 8 or 12 weeks of glecaprevir / pibrentasvir and two trials which evaluated 8 or 12 weeks of ledipasvir / sofosbuvir.^{191,193} Fourteen trials (in 12 publications) evaluated the same DAA regimen with and without ribavirin;^{137,144,154,158,160,161,185,186,188,191,193,194} of these, six trials (in 4 publications^{137,186,188,191}) evaluated ombitasvir / paritaprevir / ritonavir / dasabuvir, two trials^{185,193} ledipasvir / sofosbuvir, two trials^{144,160} elbasvir / grazoprevir, and one trial each evaluated simeprevir / sofosbuvir,¹⁵⁴ sofosbuvir / velpatasvir,¹⁵⁸ glecaprevir / pibrentasvir,¹⁹⁴ and daclatasvir / sofosbuvir.¹⁶¹ Twentyone trials did not vary duration of treatment or use of ribavirin.^{141,142,145,146,148-150,153,155-157,159,162-} ^{165,189,190,192,195,198} Thirty-two trials (in 30 publications) enrolled treatment-naïve populations or reported results stratified according to prior treatment status, ^{137,141,142,144,146,149,151-153,155-157,159-} ^{167,185,188-191,193,195,198} five trials only enrolled treatment-experienced patients, ^{137,154,158,186,194} and 11 trials (in 10 publications) enrolled a mix of treatment-naïve and -experienced patients but did not stratify results according to treatment status.^{139,143,147,148,150,187,192,196,197,199} In trials of mixed populations, the proportion of treatment-naïve patients ranged from 52 to 95 percent. SVR was measured 12 weeks after the end of treatment in all trials except for one trial that assessed SVR at 14 weeks post-treatment¹⁶⁶ and four trials (in 3 publications) that reported 12- and 24-week post-treatment SVR rates.^{167,191,192} In the latter trials, 12- and 24-week SVR rates were identical or very similar.

Twenty-seven trials (in 24 publications^{137,139,144,146,147,151-154,158-161,166,167,185-191,193,194}) had multiple DAA treatment arms, and 22 trials (in 21 publications^{141-143,145,148-150,155-157,162-165,167,192,195-199}) were single-arm studies (**Appendix B Tables 10 and 11**). Among the trials with multiple treatment arms, 20 (in 18 publications^{137,144,146,147,153,154,158-161,167,185,186,189-191,193,194}) used an open-label design. In the open-label trials, treatment allocation was random in 11 trials (in 9 publications^{137,147,153,159,167,185,186,190,194}); in the other trials patients were allocated to treatment

based on genotype (4 trials^{144,146,160,161}), prior treatment status (1 trial¹⁹¹), or clinical characteristics (e.g., fibrosis stage).^{154,158,189,193} Thirteen trials were rated good quality,^{137,139,141,146,152,159,162,164,166,187-189,191} and the remainder were rated fair quality. Frequent methodological limitations included unclear randomization or enrollment methods (e.g., unclear if the trial enrolled consecutive patients meeting inclusion criteria, or a random sample). Loss to followup was low across all trials (range 0 to 3%). All of the trials were industry-funded.

SVR Rates in Comparative Trials

DAA regimen versus placebo. One randomized trial (n=706) compared sofosbuvir / velpatasvir versus placebo in persons with HCV infection (genotypes 1, 2, 4, 5, or 6; Table 11).¹³⁹ Genotype 1 infection was present in 53 percent of patients, 32 percent of patients had previously received interferon therapy, and 19 percent had cirrhosis at baseline. Sofosbuvir / velpatasvir was associated with an SVR rate of 99 percent (618/624), compared with no cases of SVR among 116 patients randomized to placebo (RR 231.6, 95% CI, 14.6 to 3680). Across genotypes, the SVR rate with sofosbuvir / velpatasvir ranged from 97 percent to 100 percent.

DAA regimen versus telaprevir-containing regimen. Two randomized trials (reported in one publication) compared ombitasvir / paritaprevir / ritonavir / dasabuvir (with or without ribavirin) for 12 weeks versus telaprevir (12 weeks) / pegylated interferon / ribavirin (24 or 48 weeks) for genotype 1 infection (Table 11).¹³⁷ One trial (n=311) enrolled treatment-naïve patients, and the other (n=148) enrolled patients previously treated with pegylated interferon and ribavirin. In treatment- naïve patients, ombitasvir / paritaprevir / ritonavir / dasabuvir was associated with increased likelihood of SVR versus telaprevir / pegylated interferon / ribavirin (98% vs. 80%, RR 1.22, 95% CI, 1.09 to 1.37). SVR rates were similar in genotype 1a patients who received ombitasvir / paritaprevir / dasabuvir with ribavirin (97%) and genotype 1b patients who received the same regimen with or without ribavirin (98 to 99%). In the other trial, ombitasvir / paritaprevir / pegylated interferon / ribavirin in treatment-experienced patients (99% vs. 66%, RR 1.50, 95% CI, 1.22 to 1.85). SVR rates were similar for genotype 1a (100%) and 1b (99%) infection.

DAA regimen versus non-recommended DAA regimen. Two randomized trials (reported in one publication) compared sofosbuvir / velpatasvir for 12 weeks versus sofosbuvir / ribavirin for 24 weeks.¹⁴⁷ One trial (n=269) enrolled patients with genotype 2 infection (14 to 15% prior interferon therapy, 14% cirrhosis) and one trial (n=280) enrolled patients with genotype 3 infection (26% prior interferon therapy and 29 to 30% cirrhosis; results reported for non-cirrhosis subgroup). Sofosbuvir / velpatasvir was associated with increased likelihood of SVR for genotype 2 infection (99% vs. 94%, RR 1.06, 95% CI, 1.01 to 1.11) and for genotype 3 infection (non-cirrhosis subgroup, 97% vs. 87%, RR 1.11, 95% CI, 1.05 to 1.18).

Pooled SVR Rates by Genotype

Genotype 1. Thirty-two trials (total N=6,055) reported SVR rates associated with seven different DAA regimens in persons with genotype 1 infection. $^{137,139,145,146,149,151-156,159-161,163-167,185-188,190-194,197,198}$ Across DAA regimens, the pooled SVR rate was 97.7 percent (95% CI, 96.6% to 98.4%; I²=82%; Figure 2). Although statistical heterogeneity was present, the SVR rate was 91

percent or higher in all of the trials. The most frequently evaluated regimen was ombitasvir / paritaprevir / ritonavir, with or without dasabuvir or ribavirin (11 trials).^{137,139,149,151,155,186,188,191,192} The pooled SVR rate with this regimen was 93.7 percent (95% CI, 89.0% to 96.5%; I^2 =77%) for genotype 1a infection (4 trials), 98.2 percent (95% CI, 96.4% to 99.1%; I^2 =68%) for genotype 1b infection (7 trials), and 93.2 percent (95% CI, 87.0% to 96.6%, I^2 =27%) for non-subtyped genotype 1 infection (2 trials). Ledipasvir / sofosbuvir was evaluated in six trials, ^{145,156,163,185,190,193} with a pooled SVR rate of 99.4 percent (95% CI, 95.2% to 99.9%, I^2 =89%), and elbasvir / grazoprevir was evaluated in five trials^{152,160,164,166,198} with pooled SVR rate of 96.7 percent (95% CI, 95.0% to 97.8%; I^2 =55%). Four other antiviral regimens were evaluated in two or three trials each; pooled SVR rates ranged from 95.7 percent to 99.0 percent for these regimens (Table 12).

Results were similar for trials rated good quality (pooled SVR 97.2%, 95% CI, 95.2% to 98.4%) or fair quality (pooled SVR 97.9%, 95% CI, 96.7% to 98.7%), for trials that excluded patients with cirrhosis (pooled SVR 97.1%, 95% CI, 95.7% to 98.1%) or included some (less than 20% of sample) patients with cirrhosis (pooled SVR 98.7%, 95% CI, 97.1% to 99.4%), and when the analysis was restricted to trials conducted in the United States and Canada (pooled SVR 96.7%, 95% CI, 93.1% to 98.4%) (**Table 12**). Results were also similar when the analysis was restricted to trials recommended or did not omit dasabuvir in combination with ombitasvir / paritaprevir / ritonavir (pooled SVR 98.3%, 95% CI, 97.4% to 98.9%) or when the analysis was restricted to treatment-naïve patients (pooled SVR 97.4%, 95% CI, 96.1% to 98.3%).

Genotype 2. Five trials (total N=526) reported SVR rates associated with two different DAA regimens in persons with genotype 2 infection (pooled SVR 98.9%, 95% CI, 97.5% to 99.5%; $I^2=4\%$; Figure 3).^{139,147,165,196,199} Three trials evaluated sofosbuvir / velpatasvir (pooled SVR 99.7%, 95% CI, 97.6% to %, $I^2=0\%$),^{139,147,164} and two trials evaluated glecaprevir / pibrentasvir (pooled SVR 97.9%, 95% CI, 95.0% to 99.1%, $I^2=0\%$).^{196,199} Estimates were similar when trials were stratified according to quality, geographic setting, or enrollment of some patients with cirrhosis (Table 12). SVR rates were also similar in trials that were restricted to treatment-experienced patients;^{139,147} one mixed population trial reported an SVR of 100% (95% CI, 95.4% to 100%) in the subgroup of treatment-naïve patients.¹³⁹

Genotype 3. Six trials (total N=742) reported SVR rates associated with three different DAA regimens in persons with genotype 3 infection (pooled SVR 95.5%, 95% CI, 91.6% to 97.7%; $I^2=66\%$; Figure 4).^{146,147,157,158,165,167} Estimates were similar for sofosbuvir / velpatasvir (4 trials; pooled SVR 95.6%, 95% CI, 87.1% to 98.6%; $I^2=82\%$)^{146,147,157,167} sofosbuvir / daclatasvir (2 trials; pooled SVR 96.4%, 95% CI, 93.0% to 98.2%, $I^2=0\%$),^{157,167} and glecaprevir / pibrentasvir (one trial, SVR 94.9%, 95% CI, 90.2% to 97.8%).¹⁶⁷

The SVR rate was higher in five trials that excluded patients with cirrhosis (pooled SVR 96.4%, 95% CI, 94.6% to 97.5%) than in one trial¹⁶⁵ that included some patients with cirrhosis (SVR 85.7%, 95% CI, 76.5% to 91.7%; p for interaction=0.01). Results were similar when trials were stratified according to study quality or when the analysis was restricted to trials conducted in the United States or Canada (Table 12). Results were also similar when the analysis excluded results

from one trial¹⁵⁸ of sofosbuvir / velpatasvir plus ribavirin (ribavirin is not required with this regimen; pooled SVR 95.2%, 95% CI, 91.4% to 97.3%) and when the analysis was restricted to treatment-naïve patients (pooled SVR 96.1%, 95% CI, 94.5% to 97.3%) (Table 12).

Genotype 4. Ten trials (total N=485) reported SVR rates associated with five different DAA regimens in persons with genotype 4 infection (pooled SVR 98.2%, 95% CI, 94.7% to 99.4%; $I^2=50\%$; Figure 5).^{139,142,144,162,164,166,189,195,196,198} Estimates were similar for elbasvir / grazoprevir (4 trials, pooled SVR 97.3%, 95% CI, 83.2% to 99.6%, $I^2=0\%$),^{138,144,164,166,198} ombitasvir / paritaprevir / ritonavir with ribavirin (2 trials, pooled SVR 98.7%, 95% CI, 72.7% to 99.95%; $I^2=88\%$),^{162,189} and ledipasvir / sofosbuvir (2 trials, pooled SVR 98.4%, 95% CI, 93.7% to 99.6%, $I^2=25\%$)^{142,195} (Table 12). One trial each evaluated sofosbuvir / velpatasvir (SVR 100%, 95% CI, 95.9% to 100%)¹³⁹ and glecaprevir / pibrentasvir (SVR 93.5%, 95% CI, 82.1% to 98.6%).¹⁹⁶

Results were similar when the analysis was restricted to trials that were rated good quality (pooled SVR 99.1%, 95% CI, 94.0% to 99.9%), when trials were stratified according to whether they were restricted to patients without cirrhosis (pooled SVR 98.3%, 95% CI, 94.4% to 99.5%) or included some patients with cirrhosis (pooled SVR 99.1%, 95% CI, 91.2% to 99.9%), and when trials were stratified according to geographic setting (Table 12). Results were also similar when the analysis was restricted to treatment-naïve patients (pooled SVR 98.3%, 95% CI, 94.5% to 99.5%).

Genotype 5. Four trials (total N=75) reported SVR rates associated with three different DAA regimens in patients with genotype 5 infection (pooled SVR 96.0%, 95% CI, 88.3% to 98.7%; $I^2=0\%$; Figure 6).^{139,141,143,196} Estimates were similar for glecaprevir / pibrentasvir (2 trials, pooled SVR 96.0%, 95% CI, 76.4% to 99.4%; $I^2=0\%$),^{143,196} ledipasvir / sofosbuvir (1 trial, SVR 95.2%, 95% CI, 76.2% to 99.9%),¹⁴¹ and sofosbuvir / velpatasvir (1 trial, SVR 96.6%, 95% CI, 82.2% to 99.9%).¹³⁹ Estimates were similar when trials were stratified according to study quality, inclusion of patients with cirrhosis, and geographic setting (Table 12). Results were also similar when the analysis was restricted to treatment-naïve patients (pooled SVR 95.6%, 95% CI, 83.9% to 98.9%).

Genotype 6. Five trials (total N=229) reported SVR rates associated with three different DAA regimens in persons with genotype 6 infection (pooled SVR 98.2%, 95% CI, 95.4% to 99.3%, $I^2=0\%$; Figure 7).^{139,143,148,165,196} Estimates were similar for glecaprevir / pibrentasvir (2 trials, pooled SVR 97.2%, 95% 89.4% to 99.3%; $I^2=42\%$),^{143,196} sofosbuvir / velpatasvir (2 trials, pooled SVR 99.2%, 95% CI, 94.9% to 99.9%; $I^2=0\%$)^{139,165} and ledipasvir / sofosbuvir (1 trial, SVR 96.0%, 95% CI, 79.6% to 99.9%).¹⁴⁸ Results were similar when analyses were stratified according to quality, enrollment of some patients with cirrhosis, and geographic setting (Table 12). Results were also similar when the analysis was restricted to treatment-naïve patients (pooled SVR 98.4%, 95% CI, 89.6% to 99.8%).

Mixed genotypes. Two trials (total N=108) reported SVR rates associated with sofosbuvir / velpatasvir in persons with mixed genotype 1 to 6 infections (pooled SVR 95.4%, 95% CI, 89.4% to 98.1%; $I^2=0\%$; Figure 8).^{146,150} Both trials were restricted to patients without cirrhosis. In one trial¹⁴⁶ patients were treatment-naïve, and in the other trial prior treatment status was not
reported.150

Subgroup analyses. Nineteen trials (in 18 publications) reported analyses stratified according to demographic characteristics.^{139,145,147,149,150,152,156,157,164-167,185-187,190,191,198} SVR rates were similar when patients were stratified according to age in 17 trials, according to sex in 17 trials, and according to race or ethnicity in 11 trials (**Table 13**). One trial found SVR rates were slightly higher in persons with body mass index (BMI) less than 30 kg/m² versus 30 kg/m² or more (97% vs. 92%) and in persons with diabetes versus no diabetes (100% vs. 96%).¹⁸⁷

Nine trials found SVR rates were similar when analyses were stratified according to whether patients were treatment-experienced or treatment-naïve.^{151-153,155,159,163,165,167,198} Five trials (in 4 publications)^{149,150,167,192} of patients with current or recent use of methadone or buprenorphine for opioid use disorder reported SVR rates ranging from 89 to 100 percent. The other trials excluded patients with current or recent opioid use or did not report opioid use status.

Adolescents

Seven trials evaluated the effects of DAA regimens on SVR in adolescents with HCV infection (Appendix B Tables 7 and 8).^{171,173,175,176,201-203} Sample sizes ranged from 30 to 100 (total N=348), mean age ranged from 12 to 15 years, and the proportion of female participants ranged from 35 to 63 percent. Four studies^{171,173,175,203} were multinational (primarily conducted in the United States, Europe and/or Australia), and three were conducted in Egypt.^{176,201,202} In the four multinational studies, the majority (75% or more) of participants were white.^{171,173,175,203} The three Egyptian studies^{176,201,202} enrolled genotype 4 patients, one multinational study enrolled patients with genotype 1,¹⁷⁵ and three multinational studies enrolled mixed genotypes.^{171,173,203} Patients with cirrhosis were excluded in two trials and cirrhosis/fibrosis stage inclusion criteria was not reported in a third trial. In the other four trials, enrollment of patients with cirrhosis was permitted, but two of these did not conduct liver biopsy or other testing for cirrhosis at baseline. Fibrosis stage was F0-F1 in 68 to 96 percent of the population in five studies;^{171,176,201-203} fibrosis stage was unknown in over half of participants in the other two studies. The proportion of treatment-naïve patients ranged from 66 to 100 percent. In the six trials that included treatmentexperienced patients, prior HCV treatment was interferon with or without ribavirin in three trials^{171,202,203} and was unclear in three trials.^{173,175,176} Four trials evaluated DAA regimens currently recommended and FDA-approved for use in adolescents: ledipasvir and sofosbuvir (2 trials),^{175,202}sofosbuvir and ribavirin (1 trial)¹⁷³ and glecaprevir / pibrentasvir (1 trial).¹⁷¹ Three trials evaluated DAA regimens currently recommended for adults but not FDA-approved for use in adolescents: sofosbuvir and daclatasvir (2 trials)^{176,201} and ombitasvir / paritaprevir / ritonavir / dasabuvir and weight-based ribavirin (one trial).²⁰³ One study was rated good quality,¹⁷⁶ and the others fair quality, primarily due to unclear patient enrollment methods (Appendix B Table 9).

SVR was assessed at 12-weeks post-treatment. Therapy was administered for 12 weeks in all trials with the exception of sofosbuvir / ribavirin which was administered for 12 (genotype 2) or 24 (genotype 3) weeks in one trial, and glecaprevir / pibrentasvir which was administered for 8 weeks for 94 percent of the study population in one trial.¹⁷¹ Across all studies, the rate of SVR ranged from 97 to 100 percent (**Table 14; Appendix B Tables 7** and **8**). Results were similar for specific genotypes (genotype 1 [N=31]: 98% to 100%; genotype 2 [N=13]: 100%; genotype 3 [N=39]: 97%; and, genotype 4 [N=7]: 98 to 100%), though the number of adolescents with

genotype 2 or 4 infection was very small. In two studies, SVR rates were 98 percent to 100 percent for both treatment-naïve and treatment-experienced patients.^{175,203}

Key Question 8. What Are the Harms of Currently Recommended Antiviral Treatments?

Summary

- The prior review found triple therapy with boceprevir or telaprevir plus pegylated interferon and ribavirin or dual therapy with pegylated interferon and ribavirin associated with high rates of adverse events:
 - Serious adverse events: Pooled rates 8.5 to 16 percent
 - Withdrawal due to adverse event: Pooled rates 12 to 15 percent
 - Fatigue: Pooled rates 51 to 64 percent
 - Influenza-like symptoms: Pooled rates 19 to 40 percent
 - Depression: Pooled rates 19 to 22 percent
 - Headache: Pooled rates 42 to 52 percent
 - Myalgia: Pooled rates 18 to 26 percent
- The prior review found triple therapy with boceprevir associated with increased risk of thrombocytopenia (3.8% vs. 1.4%, RR 3.2, 95% CI, 1.4 to 2.8) and neutropenia (33% vs. 18%, RR 1.8, 95% CI, 1.5 to 2.3) versus dual therapy, and telaprevir associated with increased risk of anemia (52% vs. 39%, RR 1.3, 95% CI, 1.1 to 1.3). Triple therapy with telaprevir was also associated with increased risk of rash versus dual therapy (49% vs. 35%, RR 1.4, 95% CI, 1.1 to 1.7) and boceprevir with increased risk of dysgeusia (35% vs. 13%, RR 2.5, 95% CI, 2.0 to 3.2).
- Four new randomized trials found current DAA regimens associated with slightly increased risk of any adverse event versus placebo (pooled RR 1.12, 95% CI, 1.02 to 1.24, I²=46%; adjusted risk difference [ARD] 8%, 95% CI, 2% to 15%) and nausea (pooled RR 1.42, 95% CI, 1.00 to 2.03, I²=10%, ARD 4%, 95% CI, -3% to 10%). There were no differences between DAA therapy versus placebo in risk of serious adverse events, withdrawal due to adverse events, diarrhea, fatigue, headache, or anemia.
- Two new randomized trials found ombitasvir / paritaprevir / ritonavir / dasabuvir with or without ribavirin associated with decreased risk of any adverse event (RR 0.65, 95% CI, 0.50 to 0.84, I²=87%; ARD -34%, 95% CI, -51% to -16%), serious adverse events (RR 0.08, 95% CI, 0.02 to 0.34, I²=0%; ARD -8%, 95% CI, -15% to -1%), withdrawal due to adverse events (RR 0.06, 95% CI, 0.01 to 0.29, I²=0%; ARD -9%, 95% CI, -14% to -3%), fatigue (RR 0.37, 95% CI, 0.21 to 0.63, I²=32%; ARD -18%, 95% CI, -27% to -10%), headache (RR 0.70, 95% CI, 0.52 to 0.95; ARD -0.10, 95% CI, -0.20 to -0.01), nausea (RR 0.31, 95% CI, 0.16 to 0.59, I²=65%; ARD -28%, 95% CI, -37% to -19%), anemia (RR 0.19, 95% CI, 0.04 to 0.23, I²=41%; ARD -37%, 95% CI, -46% to -28%), and rash (RR 0.19, 95% CI, 0.06 to 0.58, I²=48%; ARD -17%, 95% CI, -24% to -9%) versus telaprevir / pegylated interferon / ribavirin.
- Forty-nine new trials reported the proportion of patients on DAA regimens with adverse events:
 - Any adverse event (44 trials): 73.3 percent (95% CI, 68.0% to 78.1%, $I^2=95\%$)

- Serious adverse events (44 trials): 1.9 percent (95% CI, 1.5% to 2.4%, $I^2=31\%$)
- Withdrawal due to adverse events (44 trials): 0.4 percent (95% CI, 0.3% to 0.6%, $I^2=0\%$)
- Anemia (13 trials): 2.4 percent (95% CI, 0.9% to 6.3%, $I^2=85\%$)
- Fatigue (37 trials): 18.4 percent (95% CI, 15.6% to 21.7%, $I^2=90\%$)
- Headache (42 trials): 18.7 percent (95% CI, 15.6% to 22.2%, $I^2=90\%$)
- Insomnia (18 trials): 8.3 percent (95% CI, 6.8% to 10.1%, I²=58%)
- Nausea (36 trials): 11.1 percent (95% CI, 9.1% to 13.5%, $I^2=82\%$)
- Diarrhea (18 trials): 8.7 percent (95% CI, 6.9% to 11.0%, $I^2=70\%$)
- Vomiting (6 trials): 5.8 percent (95% CI, 3.4% to 9.7%, $I^2=43\%$)
- Rash (17 trials): 5.4 percent (95% CI, 4.1% to 7.1%, $I^2=70\%$)
- There was some variability by DAA regimens in adverse events estimates; estimates were generally higher for ombitasvir / paritaprevir / ritonavir / dasabuvir with ribavirin than without ribavirin.
- Adverse event estimates were generally similar when trials were stratified according to baseline cirrhosis status (excluded or included up to 20%) and prior antiviral therapy experience.

Evidence

Adults

The prior Agency for Healthcare Research and Quality review found no difference between triple therapy with boceprevir or telaprevir plus pegylated interferon and ribavirin versus dual therapy with pegylated interferon and ribavirin in risk of serious adverse events (pooled event rates ranged from 8.5% to 16%) or withdrawal due to adverse events (pooled event rates 12% to 15%).^{5,90} There were also no differences in rates of fatigue (pooled event rates 51% to 64%), influenza-like symptoms (pooled event rates 19% to 40%), depression (pooled event rates 19% to 22%), headache (pooled event rates 42% to 52%), or myalgia (pooled event rates 18% to 26%), but these adverse events occurred frequently with all regimens. Triple therapy was associated with increased risk of hematological adverse events versus dual therapy. Boceprevir was associated with increased risk of thrombocytopenia (3.8% vs. 1.4%, RR 3.2, 95% CI, 1.4 to 2.8) and neutropenia (33% vs. 18%, RR 1.8, 95% CI, 1.5 to 2.3), and telaprevir was associated with increased risk of anemia (52% vs. 39%, RR 1.3, 95% CI, 1.1 to 1.3). Triple therapy with telaprevir was also associated with increased risk of rash versus dual therapy (49% vs. 35%, RR 1.4, 95% CI, 1.1 to 1.7) and boceprevir with increased risk of dysgeusia versus dual therapy (35% vs. 13%, RR 2.5, 95% CI, 2.0 to 3.2).

Forty-nine new trials (in 44 publications) of DAA regimens without interferon reported the proportion of patients who experienced adverse events (**Table 15**; **Appendix B Tables 10 and 11**).^{137,139,141-167,185-199} One DAA trial¹⁵⁸ included in the SVR analysis was excluded from pooled analyses of adverse events because a high proportion of patients had cirrhosis (about 40%) and adverse event rates were not reported separately for persons without cirrhosis. Eleven trials (in 9 publications) of ombitasvir / paritaprevir / ritonavir / dasabuvir included ribavirin, which is recommended for treatment of genotype 1a and 4 infections.^{137,149,162,186-189,191,192} Regimens containing ribavirin were otherwise excluded from the adverse event analyses. Eight trials (in 6

publications) reporting adverse events compared a current DAA regimen versus placebo,^{139,151,164,187} triple therapy with telaprevir,¹³⁷ or an older DAA regimen.¹⁴⁷ Reporting of methods used to assess harms was suboptimal, with few details regarding use of active versus passive assessment or definitions of harms. Trial characteristics are described in more detail in KQ 7.

Adverse Events in Comparative Trials

DAA regimen versus placebo. Four randomized trials (total N=2,113) reported adverse events associated with current DAA regimens versus placebo.^{139,151,164,187} Each trial evaluated a different DAA regimen: sofosbuvir / velpatasvir (n=706),¹³⁹ ombitasvir / paritaprevir / ritonavir / dasabuvir with ribavirin (n=477),¹⁸⁷ ombitasvir / paritaprevir / dasabuvir (n=321),¹⁵¹ and elbasvir / grazoprevir (n=609)¹⁶⁴ (**Table 15; Appendix B Tables 10 and 11**). The trials of sofosbuvir / velpatasvir and elbasvir / grazoprevir enrolled people with mixed genotype (1, 2, 4, 5, and/or 6) infections, and the other trials enrolled persons with genotype 1 infection. One trial enrolled treatment-naïve patients;¹⁶⁴ in the remaining trials, approximately one-third of patients had previously received interferon therapy. In two trials,^{139,164} approximately 19 percent of patients had cirrhosis at baseline, and the other two trials restricted enrollment to persons without cirrhosis. All trials used a double-blind design.

DAA therapy was associated with slightly increased risk of any adverse event versus placebo that was of borderline statistical significance (4 trials, RR 1.12, 95% CI, 1.02 to 1.24, I²=46%; ARD 8%, 95% CI, 8% to 15%; Figure 9).^{139,151,164,187} Among patients randomized to DAA therapy, the proportion reporting any adverse event ranged from 47 percent to 86 percent. There were no differences between DAA therapy versus placebo in risk of serious adverse events (4 trials, RR 1.90, 95% CI, 0.73 to 4.95, I²=0%; Figure 10) or withdrawal due to adverse events (4 trials, RR 0.47, 95% CI, 0.14 to 1.58, $I^2=14\%$; Figure 11), though there were few events and estimates were imprecise.^{139,151,164,187} Among patients randomized to DAA therapy, the proportion with serious adverse events ranged from 2.0 percent to 3.3 percent, and the proportion who withdrew due to adverse events ranged from 0.2 percent to 0.9 percent. DAA therapy was associated with increased risk of nausea versus placebo (3 trials, RR 1.42, 95% CI, 1.00 to 2.03, $I^2=10\%$; ARD 4%, 95% CI, -3% to 10%; Figure 12).^{139,151,187} The point estimate was similar for diarrhea, but the difference was not statistically significant (two trials, RR 1.53, 95% CI, 0.88 to 2.68, I²=29%; Figure 13).^{139,187} There were no differences between DAA therapy versus placebo in risk of fatigue (3 trials, RR 1.05, 95% CI, 0.78 to 1.40; $I^2=32\%$; Figure 14)^{139,164,187} or headache (four trials, RR 1.12, 95% CI, 0.92 to 1.37, I²=0%; Figure 15).^{139,151,164,187} One trial¹³⁹ found no difference between sofosbuvir / velpatasvir versus placebo in risk of anemia (0.3% vs.)0%, RR 2.21, 95% CI, 0.11 to 46); no cases of anemia were reported in the other three trials.

DAA regimen versus telaprevir/pegylated interferon/ribavirin. Two randomized trials (reported in one publication) compared ombitasvir / paritaprevir / ritonavir / dasabuvir with or without ribavirin for 12 weeks versus triple therapy with telaprevir (12 weeks) / pegylated interferon / ribavirin (24 or 48 weeks) in patients with genotype 1 infection.¹³⁷ One trial (n=311) enrolled treatment-naïve patients, and one trial (n=148) enrolled patients previously treated with pegylated interferon and ribavirin. The DAA regimen was associated with decreased risk of any adverse event (RR 0.65, 95% CI, 0.50 to 0.84, I²=87%; ARD -34%, 95% CI, -51% to -16%;

Figure 16), serious adverse events (RR 0.08, 95% CI, 0.02 to 0.34, $I^2=0\%$; ARD -8%, 95% CI, -15% to -1%; **Figure 17**), withdrawal due to adverse events (RR 0.06, 95% CI, 0.01 to 0.29, $I^2=0\%$; ARD -9%, 95% CI, -14% to -3%; **Figure 18**), fatigue (RR 0.37, 95% CI, 0.21 to 0.63, $I^2=32\%$; ARD -18%, 95% CI, -27% to -10%; **Figure 19**), headache (RR 0.70, 95% CI, 0.52 to 0.95, $I^2=0\%$; ARD -0.10, 95% CI, -0.20 to -0.01; **Figure 20**), nausea (RR 0.31, 95% CI, 0.16 to 0.59, $I^2=65\%$; ARD -28%, 95% CI, -37% to -19%; **Figure 21**), anemia (RR 0.09, 95% CI, 0.04 to 0.23, $I^2=41\%$; ARD -37%, 95% CI, -46% to -28%; **Figure 22**), and rash (RR 0.19, 95% CI, 0.04 to 0.58, $I^2=48\%$; ARD -17%, 95% CI, -24% to -9%; **Figure 23**) versus the telaprevir regimen. The association between DAA therapy versus telaprevir and risk of any adverse event was less pronounced when ribavirin was included with DAA therapy (2 trials, RR 0.74, 95% CI, 0.65 to 0.84, $I^2=43\%$; **Figure 16**) than without ribavirin (1 trial, RR 0.50, 95% CI, 0.40 to 0.62; p for interaction=0.003). There was no interaction between prior antiviral treatment experience and risk estimates for any adverse event.

Pooled Adverse Event Rates for DAA Regimens

Any adverse event. Forty-four trials (in 41 publications, total N=8,045) reported the proportion of patients reporting any adverse event with eight different DAA regimens.^{137,139,141-156,159-167,185-190,192-199} Across regimens, the pooled rate for any adverse event was 73.3% (95% CI, 68.0% to 78.1%, I^2 =95%; **Figure 24**). Stratified by antiviral regimen, the rate of any adverse event ranged from 62.3% (95% CI, 56.1% to 68.1%) for glecaprevir / pibrentasvir (7 trials) to 82.7% (95% CI, 58.5% to 94.2%) for sofosbuvir / daclatasvir (2 trials). The rate of any adverse event was higher in trials of ombitasvir / paritaprevir / ritonavir / dasabuvir with ribavirin (10 trials [in 8 publications] pooled event rate 81.1%, 95% CI, 62.3% to 84.6%; I^2 =87%) than without ribavirin (6 trials, pooled event rate 75.1%, 95% CI, 62.3% to 84.6%; I^2 =92%) (**Table 16**). The proportion of patients with any adverse event was similar when trials were stratified according to whether they excluded patients with cirrhosis (24 trials, pooled event rate 75.5%, 95% CI, 69.0% to 81.1%) or included some patients with cirrhosis (19 trials, pooled event rate 72.4%, 95% CI, 64.6% to 79.0%; p for interaction=0.52), and there was no interaction between prior treatment experience status and rates of any adverse event (p for interaction=0.76).

Serious adverse events. Forty-four trials (in 40 publications, total N=8,070) reported the proportion of patients reporting serious adverse events with eight different DAA regimens.^{137,139,141-144,146-157,160-167,185-194,196-199} Across regimens, the pooled rate for serious adverse events was 1.9 percent (95% CI, 1.5% to 2.4%, I²=31%; **Figure 25**). Stratified by antiviral regimen, the rate of any adverse event ranged from 0.6 percent (95% CI, 0.1% to 4.1%, I²=0%) for simeprevir / sofosbuvir (2 trials) to 2.1 percent for elbasvir / grazoprevir (6 trials, 95% CI, 1.1% to 3.9%, I²=42%) and ombitasvir / paritaprevir / ritonavir / dasabuvir with ribavirin (11 trials, 95% CI, 1.5% to 3.0%, I²=26%) (**Table 16**). The rate of serious adverse events for ombitasvir / paritaprevir / dasabuvir without ribavirin. Estimates were similar when trials were stratified according to whether they excluded patients with cirrhosis (23 trials, pooled event rate 1.8%, 95% CI, 1.3% to 2.5%) or included some patients with cirrhosis (21 trials, pooled event rate 2.0%, 95% CI, 1.4% to 2.7%; p for interaction=0.69), and there was no interaction between prior treatment experience status and rates of serious adverse events (p for interaction=0.96).

Withdrawal due to adverse events. Forty-four trials (in 40 publications, total N=8,060) reported the proportion of patients who withdrew due to adverse events with eight different DAA regimens.^{137,139,141-156,160-167,185-194,196-199} Across regimens, there were a total of 35 withdrawals due to adverse events, with a pooled rate of 0.4 percent (95% CI, 0.3% to 0.6%, I^2 =0%; **Figure 26**). The proportion of patients who withdrew due to adverse events was less than or equal to 1 percent for all regimens (**Table 16**).

Anemia. Thirteen trials (in 9 publications, total N=1,555) reported the proportion of patients with anemia with five different DAA regimens.^{137,149,154,185,186,190-192,199} Across regimens, the pooled rate for anemia was 2.4 percent (95% CI, 0.9% to 6.3%, I²=85%; Figure 27). The rate of anemia was much higher in trials of ombitasvir / paritaprevir / ritonavir / dasabuvir with ribavirin (6 trials, pooled event rate 8.3%, 95% CI, 5.8% to 11.8%, I²=49%) than the same regimen without ribavirin (3 trials, pooled event rate 0.8%, 95% CI, 0.2% to 3.1%, I²=0%) or with other regimens (pooled event rates <0.5%) (Table 17).

Fatigue. Thirty-seven trials (in 33 publications, total N=7,571) reported the proportion of patients with fatigue with eight different DAA regimens. ^{137,139,141-150,153,155-157,159-162,164,167,185-192,194-196} Across regimens, the pooled rate for fatigue was 18.4 percent (95% CI, 15.6% to 21.7%, I^2 =90%; **Figure 28**). Stratified by antiviral regimen, rates of fatigue ranged from 10.9 percent (95% CI, 4.3% to 25.1%, I^2 =88%) for elbasvir / grazoprevir (3 trials) to 26.9 percent (95% CI, 20.5% to 34.4%, I^2 =88%) for ombitasvir / paritaprevir / ritonavir / dasabuvir with ribavirin (11 trials) (**Table 17**). The rate of fatigue was higher for ombitasvir / paritaprevir / ritonavir / dasabuvir with ribavirin than the same regimen without ribavirin (6 trials, pooled event rate 15.8%, 95% CI, 9.1% to 26.1%, I^2 =91%). Estimates were similar when trials were stratified according to whether they excluded patients with cirrhosis (18 trials, pooled event rate 16.7%, 95% CI, 13.1% to 21.2%; p for interaction=0.27) and there was no interaction between prior treatment status and rates of fatigue (p for interaction=0.54).

Headache. Forty-two trials (in 38 publications, total N=7,790) reported the proportion of patients with headache with 8 different DAA regimens.^{137,139,141-151,153,155-157,159-162,164,165,167,185-197,199} Across regimens, the pooled rate for headache was 18.7 percent (95% CI, 15.6% to 22.2%, I²=90%; **Figure 29**). Stratified by antiviral regimen, rates of headache ranged from 13.7 percent (95% CI, 8.4% to 21.5%, I²=85%) for ledipasvir / sofosbuvir (9 trials) to 27.6 percent (95% CI, 24.0% to 31.5%, I²=60%) for ombitasvir / paritaprevir / ritonavir / dasabuvir with ribavirin (11 trials) (**Table 17**). The rate of headache was higher for ombitasvir / paritaprevir / ritonavir / dasabuvir with ribavirin than the same regimen without ribavirin (7 trials, pooled event rate 20.7%, 95% CI, 15.6% to 26.9%, I²=83%). Estimates were similar when trials were stratified according to whether they excluded patients with cirrhosis (14 trials, pooled event rate 19.6%, 95% CI, 15.5% to 24.3%) or included some patients with cirrhosis (19 trials, pooled event rate 19.1%, 95% CI, 14.9% to 24.1%; p for interaction=0.89), and there was no interaction between prior treatment experience status and rates of headache (p for interaction=0.11).

Insomnia. Eighteen trials (in 17 publications, total N=3,517) reported the proportion of patients with insomnia with eight different DAA regimens.^{139,146,147,149,150,157,159-162,185,187,189,190,192,194,195} Across regimens, the pooled rate for insomnia was 8.3 percent (95% CI, 6.8% to 10.1%, I^2 =58%;

Figure 30). Stratified by antiviral regimen, rates of insomnia ranged from 6.0% (95% CI, 4.5 to 8.0%; $I^2=58\%$) for ledipasvir / sofosbuvir (3 trials) to 13.3% (95% CI, 11.1% to 15.9%; $I^2=0\%$) for ombitasvir / paritaprevir / ritonavir / dasabuvir with ribavirin (5 trials) (**Table 17**). The only trial of glecaprevir / pibrentasvir reported no cases of insomnia.¹⁶⁰ Estimates were similar when trials were stratified according to whether they excluded patients with cirrhosis (10 trials, pooled event rate 9.0%, 95% CI, 7.0% to 11.5%) or included some patients with cirrhosis (8 trials, pooled event rate 8.4%, 95% CI, 6.4% to 10.9%; p for interaction=0.70), and there was no interaction between prior treatment experience status and rates of insomnia (p for interaction=0.81).

Gastrointestinal adverse events. Thirty-six trials (in 34 publications, total N=6,145) reported the proportion of patients with nausea on eight different DAA regimens.^{137,139,142,144-151,153,157,159-^{162,167,185,186,188-196,199} Across regimens, the pooled rate for nausea was 11.1 percent (95% CI, 9.1% to 13.5%, I²=82%; **Figure 31**). Stratified by antiviral regimen, rates of nausea ranged from 6.5 percent (95% CI, 4.3% to 9.7%, I²=70%) for ombitasvir / paritaprevir / ritonavir / dasabuvir without ribavirin (7 trials) to 15.2 percent (95% CI, 9.6% to 23.2%, I²=90%) for ombitasvir / paritaprevir / ritonavir / dasabuvir with ribavirin (11 trials) (**Table 18**). Estimates were similar when trials were stratified according to whether they excluded patients with cirrhosis (21 trials, pooled event rate 10.6%, 95% CI, 8.2% to 13.5%) or included some patients with cirrhosis (14 trials, pooled event rate 12.9%, 95% CI, 9.6% to 17.1%; p for interaction=0.31), and there was no interaction between prior treatment experience status and rates of nausea (p for interaction=0.63).}

Eighteen trials (in 17 publications, total N=2,336) of six different DAA regimens reported the proportion of patients with diarrhea. ^{141,142,146,148,150,155,157,160,161,185-191,195} Across regimens, the pooled rate of diarrhea was 8.7 percent (95% CI, 6.9% to 11.0%, I²=70%; **Figure 32**). Stratified by antiviral regimen, rates of diarrhea ranged from 6.1 percent (95% CI, 3.4% to 10.8%, I²=50%) for sofosbuvir / velpatasvir (2 trials) to 11.6 percent (95% CI, 4.9% to 25.0%) for elbasvir / grazoprevir (1 trial) (**Table 18**). The rate of diarrhea was similar for ombitasvir / paritaprevir / ritonavir / dasabuvir with ribavirin (6 trials, pooled event rate 10.9%, 95% CI, 7.8% to 14.9%, I²=73%) and the same regimen without ribavirin (5 trials, pooled event rate 11.1%, 95% CI, 7.7% to 15.9%, I²=72%). Estimates were similar when trials were stratified according to whether they excluded patients with cirrhosis (10 trials, pooled event rate 10.1%, 95% CI, 7.9% to 12.8%) or included some patients with cirrhosis (5 trials, pooled event rate 8.0%, 95% CI, 5.5% to 11.6%; p for interaction=0.33), and there was no interaction between prior treatment experience status and rates of diarrhea (p for interaction=0.92).

Six trials (total N=444) of five different DAA regimens reported the proportion of patients with vomiting.^{148-150,159,161,192} Across regimens, the pooled rate of vomiting was 5.8 percent (95% CI, 3.4% to 9.7%, I²=43%; **Figure 33**). Stratified by antiviral regimen, rates of vomiting ranged from 1.9 percent (95% CI, 0.5% to 7.2%, I²=0%) for sofosbuvir / daclatasvir (2 trials) to 12.0 percent (2 trials, 95% CI, 7.4% to 18.9%; I²=0%) with ombitasvir / paritaprevir / ritonavir / dasabuvir with ribavirin.

Rash. Seventeen trials (in 15 publications, total N=2,256) reported the proportion of patients with rash on eight different DAA regimens.^{137,146,153,154,158-160,185-188,190,192,193,197} Across regimens,

the pooled rate for rash was 5.4 percent (95% CI, 4.1% to 7.1%, $I^2=70\%$; **Figure 34**). Stratified by antiviral regimen, rates of rash ranged from 1.5 percent (95% CI, 0.2% to 10.1%) for sofosbuvir / daclatasvir (1 trial) to 8.3 percent (95% CI, 4.9% to 13.7%, $I^2=45\%$) for sofosbuvir / velpatasvir (2 trials) (**Table 18**). The rate of rash was higher for ombitasvir / paritaprevir / ritonavir / dasabuvir with ribavirin (7 trials, pooled event rate 7.6%, 95% CI, 5.5% to 10.3%, $I^2=57\%$) than the same regimen without ribavirin (4 trials, event rate 2.6%, 95% CI, 1.0% to 6.7%, $I^2=66\%$). Estimates were similar when trials were stratified according to whether they excluded patients with cirrhosis (13 trials, pooled event rate 5.2%, 95% CI, 3.8% to 7.1%) or included some patients with cirrhosis (4 trials, pooled event rate 6.2%, 95% CI, 3.7% to 10.1%; p for interaction=0.56), and there was no interaction between prior treatment experience status and rates of rash (p for interaction=0.49).

HBV infection reactivation. All trials but one¹⁹⁵ excluded persons coinfected with HBV infection, and no cases of HBV reactivation were reported.

Adolescents

Seven trials of DAA regimens in adolescents reported harms (**Table 19; Appendix B Tables 7-9**).^{171,173,175,176,201-203} Study characteristics were described in detail in KQ 7; four trials evaluated regimens FDA-approved for use in adolescents (ledipasvir and sofosbuvir,^{175,202} sofosbuvir and ribavirin,¹⁷³ or glecaprevir / pibrentasvir¹⁷¹), and three trials evaluated DAA regimens recommended in adults but not approved in children (sofosbuvir and daclatasvir^{176,201} or ombitasvir / paritaprevir / ritonavir / dasabuvir²⁰³). Methods for reporting and assessing harms were generally not well described.

Five trials reported no withdrawals due to adverse events,^{171,173,175,176,203}, and one of five trials reported a single serious adverse event (a grade 3 joint injury) in adolescents treated with sofosbuvir plus ribavirin.¹⁷³ The rate of any adverse event was 27 percent in one study of sofosbuvir and daclatasvir (not FDA-approved for use in adolescents)¹⁷⁶ and 71 to 87 percent in four trials of other regimens.^{171,173,175,203} Rates of specific adverse events ranged from 3 to 48 percent for headache (7 trials),^{171,173,175,176,201-203} 5 to 53 percent for fatigue (7 trials),^{171,173,175,176,201-203} and 3 to 28 percent for gastrointestinal (nausea, vomiting, or diarrhea) adverse events (5 trials).^{173,175,176,201,202} One trial of ledipasvir and sofosbuvir reported insomnia in 23 percent (9/40) of participants.²⁰² Stratification by DAA regimen did not appear to explain the observed variability in adverse event estimates, though assessments were limited by the small number of trials and methodological limitations.

Key Question 9. What Is the Association Between Experiencing SVR Following Antiviral Treatment and Reduction in Risk of HCV-Related Adverse Health Outcomes?

Summary

• The prior review included 10 studies of patients in which less than 25 percent had

cirrhosis at baseline that found SVR after interferon-based antiviral therapy associated with decreased risk of all-cause mortality (7 studies, adjusted HR 0.12 to 0.71), liver-related mortality (5 studies, adjusted HR 0.04 to 0.22), and HCC (4 studies, adjusted HR 0.12 to 0.36) versus no SVR.

- Including studies published since the prior review, SVR after antiviral therapy was associated with decreased risk of all-cause mortality, liver mortality, cirrhosis, and HCC versus no SVR in studies in which less than 25 percent of the population had cirrhosis at baseline.
 - All-cause mortality (13 studies): Pooled adjusted HR 0.40 (95% CI, 0.28 to 0.56, $I^2=52\%$).^{69,168,204-214}
 - Liver mortality (4 studies): Pooled adjusted HR 0.11 (95% CI, 0.04 to 0.27, $I^2=0\%$).^{204,208,210,213}
 - Cirrhosis (4 cohorts reported in 3 studies): Pooled HR 0.36 (95% CI, 0.33 to 0.40; $I^2=0\%$).^{206,215,216}
 - HCC (20 studies): Pooled adjusted HR 0.29 (95% CI, 0.23 to 0.38; $I^2=19\%$).^{168,204,207,211,214,215,217-230}
- Estimates favored SVR in all studies and results were consistent when studies with potentially overlapping populations were excluded from analyses, when the analysis was restricted to studies that adjusted at a minimum for age, sex, genotype, and baseline fibrosis, and in stratified analyses based on duration of followup and geographic setting. For all-cause mortality, the effect of SVR was stronger in studies with followup longer than 5 years.
- All studies except for three evaluated SVR after interferon-based therapy; results were similar from two studies of SVR after DAA therapy, and estimates from a third study of SVR after DAA therapy were very imprecise.

Evidence

The prior review included 19 cohort studies that consistently found an SVR after interferonbased antiviral therapy associated with decreased risk of all-cause mortality (10 studies, adjusted HR ranged from 0.07 to 0.39), liver-related mortality (9 studies, adjusted HR 0.04 to 0.27), and HCC (11 studies, adjusted hazards ratios 0.12 to 0.71) versus no SVR after 3 to 9 years of followup. Six studies in the prior review evaluated populations of patients with cirrhosis at baseline, and in three other studies the proportion of patients with cirrhosis at baseline ranged from 30 to 70 percent. When results were restricted to 10 studies in which less than 25 percent of persons had cirrhosis at baseline, results also indicated an association between SVR after antiviral therapy and decreased risk of all-cause mortality (7 studies, adjusted HR 0.12 to 0.71), liver-related mortality (5 studies, adjusted HR 0.04 to 0.22), and HCC (4 studies, adjusted HR 0.12 to 0.36). The largest study (n=16,864), which also had the fewest methodological shortcomings, found SVR after antiviral therapy in a predominantly male, VA population associated with lower risk of all-cause mortality versus no SVR after a median of 3.8 years (adjusted HR 0.71, 95% CI, 0.60 to 0.86, 0.62, 95% CI, 0.44 to 0.87, and 0.51, 95% CI, 0.35 to 0.75, for genotypes 1, 2, and 3, respectively).⁶⁹

Thirty cohort studies (30 publications) reported associations between achieving SVR following antiviral treatment versus no SVR and clinical outcomes (**Appendix B Tables 14 and**

15).^{69,168,204-231} Nine of the studies were included in the prior report;^{69,204,208-211,213,214,222} nine other studies^{70,232-238} from the prior review were excluded because more than 25 percent of the populations had cirrhosis at baseline, and one study²³⁹ from the prior review was excluded because it did not report usable data.

Sample sizes ranged from 145 to 50,886 (total N=116,821), mean age ranged from 42 to 69 years, and the proportion of female participants ranged from 1.1 to 60 percent with five studies including samples that were less than 10 percent female.^{69,205,207,215,221} The proportion of patients with cirrhosis at baseline ranged from 0 percent to 21 percent. Seventeen studies were conducted in Japan, ^{204,208,210,211,213,217-220,222-224,226-230} seven in the United States, ^{69,205-207,212,215,221} two in South Korea,^{225,231} two in Taiwan,^{214,216} one in France,¹⁶⁸ and one in the United Kingdom.²⁰⁹ All of the U.S.-based studies except for one²¹² were conducted in VA populations. Several Japanese studies also appeared to evaluate overlapping or partially overlapping populations (**Table 20**; Appendix B Tables 14 and 15). None of the studies conducted in Asian countries reported race; among studies in the United States and the United Kingdom, white patients comprised 38 to 92 percent of the study population, black patients comprised 8 to 43 percent of the population, and Hispanic patients comprised 0.4 to 14 percent of the population. When genotype was reported, genotype 1 was generally the most common (36% to 99%), with genotype 2 the second most common (8% to 52%). One study reported that 52 percent of patients were genotype 2,²³¹ and two studies reported large proportions (54% and 55%) of 'non-genotype 1' patients, but did not otherwise specify genotype.^{209,214}

Three studies were prospective,^{168,218,224} and the others were either retrospective cohort studies or analyzed a prospectively collected dataset retrospectively. Twenty-six studies, including all of the studies carried forward from the prior USPSTF review, evaluated the association between SVR and clinical outcomes following treatment exclusively with interferon-based treatments.^{69,204,206-214,216-220,222-231} Three studies focused on DAAs,^{168,205,221} one study evaluated interferon-based treatments and DAAs,²²¹ and one study did not report what type of treatment was administered (likely primarily interferon-based therapies, given study date).²¹⁵ Average followup ranged from 1.5 to 10 years in all studies except for one study that described followup of at least a year.²³¹

Twenty studies evaluated the outcome HCC,^{168,204,207,211,214,215,217-230} thirteen studies all-cause mortality,^{69,168,204-214} seven liver-related mortality,^{204,207-210,213,214} four cohorts (in three publications) cirrhosis,^{206,215,216} and one study liver transplantation.²⁰⁷ Two studies evaluated composite outcomes related to mortality and liver outcomes,^{206,231} and one study assessed liver-related hospital episodes.²⁰⁹

All studies were rated fair quality (**Appendix B Table 16**). Although studies had to perform statistical analyses on potential confounders, 13 studies did not address all four pre-specified factors (age, sex, fibrosis stage, and genotype).^{206,208,210,213,216,218,220,222,224,226,227,230,231} No study matched patients who achieved SVR with patients who did not achieve SVR on potential confounders. Studies did not report baseline characteristics according to SVR status or reported large baseline differences between groups. Other methodological shortcomings included failure to report missing data or attrition and unclear masking of outcome assessors.

All-Cause Mortality

SVR after antiviral therapy was associated with decreased risk of all-cause mortality versus no SVR (13 studies, pooled HR 0.40, 95% CI, 0.28 to 0.56, I²=52%) (Figure 35).^{69,168,204-214} Estimates favored SVR in all studies, and HRs ranged from 0.11 to 0.66. Findings were similar when three studies²⁰⁶⁻²⁰⁸ with potentially overlapping populations were excluded from the analysis (pooled HR 0.37, 95% CI, 0.25 to 0.56). The estimate was slightly weaker in ten "fully adjusted" studies (defined as study methods controlled for age, sex, fibrosis stage, and genotype at a minimum; pooled HR 0.42, 95% CI, 0.29 to 0.62) than studies with partial adjustment (pooled HR 0.29, 95% CI, 0.15 to 0.55), but the difference was not statistically significant (p for interaction=0.34) (Table 21). Trials with longer duration of followup (more than 5 years) reported a stronger association between SVR after antiviral therapy and reduced risk of all-cause mortality (pooled HR 0.33, 95% CI, 0.24 to 0.46) than those with shorter followup (pooled HR 0.64, 95% CI, 0.56 to 0.74; p for interaction=0.003). In stratified analyses, there was no association between geographic setting (United States or Europe vs. Asia, p for interaction=0.10) or the proportion of patients with cirrhosis at baseline (more than 10% vs. 0 to 10%, p for interaction=0.58) and risk of all-cause mortality following SVR (Table 21). Patients received interferon therapy without a DAA in all studies, with the exception of one²⁰⁵ U.S. study conducted in a VA population and one French study¹⁶⁸ in which patients received DAA therapy. The VA study found an SVR after DAA therapy associated with decreased risk or mortality compared with no SVR (adjusted HR 0.57, 95% CI, 0.33 to 0.99), though duration of followup was relatively short (1.5 years);²⁰⁵ the estimate from the French study was very imprecise (adjusted HR 1.36, 95% CI, 0.15 to 12.35).¹⁶⁸

Liver Mortality

SVR after antiviral therapy was associated with decreased risk of liver mortality versus no SVR (4 studies, pooled HR 0.11, 95% CI, 0.04 to 0.27, $I^2=0\%$) (**Figure 36**).^{204,208,210,213} Estimates favored SVR in all studies, and HRs ranged from 0.05 to 0.13. All of the studies were conducted in Asia in patients who received interferon therapy without a DAA with duration of followup longer than 5 years. Estimates were very similar when studies were stratified according to whether they were fully or partially adjusted or whether the proportion of patients with cirrhosis at baseline was 0 to 10 percent or over 10 percent, with HR estimates ranging from 0.10 to 0.13 (**Table 21**).

Cirrhosis

SVR after antiviral therapy was associated with decreased risk of cirrhosis versus no SVR (4 cohorts reported in 3 studies, pooled HR 0.36, 95% CI, 0.33 to 0.40; $I^2=0\%$) (**Figure 37**).^{206,215,216} Estimates favored SVR in all studies, and HRs ranged from 0.29 to 0.40. Three cohorts were from the United States and one²¹⁶ from Asia. All patients received treatment with interferon therapy without a DAA, or the antiviral regimen was not reported²¹⁵ but likely to be interferon therapy based on the study date. Estimates were very similar when studies were stratified according to whether they were fully or partially adjusted or the proportion of patients with cirrhosis at baseline (**Table 21**).

Hepatocellular Carcinoma

SVR after antiviral therapy was associated with decreased risk of HCC versus no SVR (20 studies, pooled HR 0.29, 95% CI, 0.23 to 0.38; I²=19%) (Figure 38).^{168,204,207,211,214,215,217-230} Estimates favored SVR in all studies, and HRs ranged from 0.06 to 0.41. Findings were similar when four studies with potentially overlapping populations^{207,215,219,222} were excluded from the analysis (pooled HR 0.25, 95% CI, 0.19 to 0.35). Pooled estimates were similar for four studies conducted in the United States and Europe (pooled HR 0.32, 95% CI, 0.28 to 0.36)^{168,207,215,221} and 16 studies conducted in Asia (pooled HR 0.24, 95% CI, 0.18 to 0.33; p for interaction=0.37). Pooled estimates were also very similar when studies were stratified according to whether they were fully or partially adjusted, the duration of followup (longer or shorter than 5 years), or the proportion of patients with cirrhosis at baseline (greater or less than 10%) (Table 21). Patients received or were likely to have received interferon therapy without a DAA in all studies except for one VA study²²¹ of DAA-only therapy, DAA plus interferon, or interferon-only therapy and one French study¹⁶⁸ of DAA-only therapy. Like the other studies, the VA study found SVR after antiviral therapy associated with decreased risk of HCC versus no SVR (adjusted HR 0.39, 95% CI, 0.35 to 0.43). Estimates were similar when the analysis was stratified according to receipt of a DAA-only regimen (adjusted HR 0.29, 95% CI, 0.23 to 0.37), a DAA plus interferon (adjusted HR 0.48, 95% CI, 0.32 to 0.73), or interferon-only (adjusted HR 0.32, 95% CI, 0.28 to 0.37). The French study was also consistent with an association between SVR after DAA therapy and decreased risk of HCC, though the estimate was imprecise and not statistically significant (adjusted HR 0.22, 95% CI, 0.03 to 1.76).¹⁶⁸

Contextual Question 1. Based on Population Level Estimates, What Are Recent Trends in the Epidemiology, Prevalence, and Incidence of HCV Infection in the United States, Including in Primary Care Settings, Over the Past 5 to 10 Years?

The incidence of HCV infection increased 3.5-fold from 2010 to 2016, rising each year during that period.²⁰ The annual increase was 20 percent from 2012 to 2013, 2.6 percent in 2014, 11 percent in 2015, and 22 percent in 2016. An estimated 41,200 new HCV infections occurred in 2016.

The increase in HCV incidence in the United States has primarily been concentrated among young persons and PWID.²⁰ From 2004 to 2010, the proportion of cases of acute HCV infection reporting injection drug use in each year ranged from 59 percent to 72 percent; since 2011, the proportion has been at least 75 percent in each year (84% in 2014).²⁴⁰ Acute HCV incidence in persons 18 to 29 years of age increased from 0.4 cases per 100,000 in 2004 to 2.0 cases per 100,000 in 2014 and in persons 30 to 39 years of age from 0.4 cases per 100,000 to 1.7 cases per 100,000 over the same time period.²⁴⁰ Among persons 40 to 49 years of age, the incidence of acute HCV infection increased slightly from 0.5 to 0.7 cases per 100,000, and in persons 50 to 59 years of age incidence was unchanged at 0.2 cases per 100,000. The increase in acute HCV incidence in young persons was greater in nonurban counties (13% annually) than in urban counties (5% annually).²⁴¹ Similar trends in acute HCV incidence have been reported in specific

regions in the United States. One study found a 364 percent increase between 2006 and 2012 in HCV infection among persons 12 to 29 years of age living in the Appalachian region of the United States.^{21,22} Another study found that new cases of HCV infection among persons 15 to 24 years of age in Massachusetts nearly doubled from 2002 to 2009.²³

Recent trends towards increased HCV prevalence among reproductive aged (15 to 44 years) females have also been observed.^{24,25} Analyses of national laboratory databases (reasons for testing not available) estimate that the number of reproductive aged females with acute and past or present HCV infection doubled from 2006 to 2014,²⁵ with an increase of 22 percent from 2011 to 2014.²⁴ Among pregnant females who underwent testing from 2011 to 2014, 0.73 percent had HCV infection.²⁵ Over the same time period there was a 68 percent increase (from 0.19% to 0.32%) in the proportion of infants born to HCV-infected females.²⁴ Similar trends have been observed in several states. For example, in Kentucky, the rate of HCV detection among females of childbearing age increased 21 percent from 2011 to 2014 (from 139 to 169 per 100,000), and the proportion of infants born to HCV-infected females increased from 0.71 percent to 1.59 percent.²⁴ In Wisconsin Medicaid recipients, the prevalence of HCV infection increased from 0.27 percent in 2011 to 0.52 percent in 2015.²⁴² Nationally, 29,000 females with HCV infection are estimated to give birth each year, resulting in 1,700 infected infants.²⁵ Within the United States., there are geographic variations in trends regarding incidence and prevalence of HCV infection. From 2004 to 2014, six states (Kansas, Maine, New Jersey, Wisconsin, Ohio, and Massachusetts) reported increases in HCV incidence of 1,000 percent or higher.²⁴⁰ A positive correlation was observed between increases in acute HCV infection incidence at the state level and increases in the proportion of treatment admissions reporting opioid injection drug use. Nine states (California, Texas, Florida, New York, Pennsylvania, Ohio, Michigan, Tennessee, and North Carolina) account for over half (51.9%) of persons living with HCV infection; five of these states are in the Appalachian region.²⁴³

Population level estimates of HCV prevalence based on the 2013 to 2016 NHANES data of noninstitutionalized civilians in the United States and incorporating estimates from four additional populations not included in NHANES (incarcerated persons, unsheltered homeless persons, active duty military personnel, and nursing home residents) indicate approximately 4.1 (range 3.4 to 4.9) million persons positive for HCV antibody and 2.4 (range 2.0 to 2.8) million persons chronically infected.¹⁶ This is lower than an earlier estimate of total HCV prevalence that used 2003 to 2010 NHANES data (4.6 million positive for HCV antibody and 3.5 with chronic infection),¹³ but there were differences in estimation methods, making it difficult to assess time trends. Based on NHANES data alone, the prevalence of chronic HCV infection decreased slightly in 2013 to 2016 to 0.84 percent (95% CI, 0.75% to 0.96%) from 1.0 percent (95% CI, 0.8 to 1.2%) in 2003 to 2010.¹⁸ Factors influencing the observed trends include declines in prevalence due to mortality primarily in the 1945 to 1965 birth cohort and use of more effective antiviral therapies, offset by the higher incidence of acute HCV infection in younger persons primarily related to injection drug use. Data to determine how recent trends in the epidemiology of acute HCV infection among young white persons have impacted the epidemiology of chronic HCV infection are not yet available.

Contextual Question 2. What Are the Effects of Different Risk- or Prevalence-Based Methods for Screening for HCV Infection in Modeling Studies?

The USPSTF previously reviewed two modeling studies that found birth-cohort screening of all persons in the United States born between 1945 and 1965 to be cost-effective compared with risk-based screening.^{8,9} Although one analysis assumed rates of progression to cirrhosis and mortality substantially higher than observed in longitudinal cohorts,⁸ the other study utilized more conservative estimates consistent with natural history data.⁹ Several other cost-effective compared with risk based screening alone.²⁴⁴⁻²⁴⁶ All of these analyses were based on treatment with outdated antiviral regimens (i.e., no all DAA regimens), reducing relevance to current practice, and did not compare expanded screening strategies versus currently recommended screening (risk-based plus birth cohort screening).

Five studies published since the prior USPSTF modeled the cost-effectiveness of HCV screening in U.S. settings based on use of DAA regimens (**Table 22**). Two studies evaluated cost-effectiveness of screening in the general adult population,^{247,248} one focused on screening persons 15 to 30 years of age,²⁴⁹ and two evaluated cost-effectiveness of prenatal HCV screening.^{250,251} The analyses generally found expanded HCV screening strategies associated with incremental cost-effectiveness ratios of less than \$50,000/quality adjusted life year (QALY), though there was variability in the screening strategies compared and cost-effectiveness estimates, due in part to differences in the assumptions used in each model.

One analysis by Barocas et al. of HCV screening in the general adult population utilized the Hepatitis C Cost-Effectiveness (HEP-CE) model, an individual-based, stochastic Monte Carlo simulation model with an embedded Markov state transition matrix.²⁴⁷ It compared one time "standard of care" birth cohort screening of all U.S. persons born between 1945 and 1965 versus one time screening of all persons at least 18, at least 30, or at least 40 years of age. All screening strategies included targeted screening of high-risk persons. The model assumed that all cases of incident HCV infection were related to injection drug use (12 cases per 100 person-years), with background (not related to screening) testing rates of 33 percent in PWID and 2.6 percent to 27 percent in other persons. Treatment was based on sofosbuvir / velpatasvir at a cost of \$23,026 per month (\$0 to \$38,000 in sensitivity analyses), with an SVR rate in persons without cirrhosis of 99 percent (50 to 99% in sensitivity analyses) and in persons with cirrhosis of 93 percent (93 to 96% in sensitivity analyses).

The model estimated that compared with birth cohort screening, the 18 and over strategy would identify 256,000 additional cases of HCV infection and lead to 280,000 additional cures and 4,400 fewer cases of HCC over the cohort lifetime, with an incremental cost-effectiveness ratio of \$28,193/QALY. More cures than additional cases of HCV infection occurred in the model because of reinfections. Among persons with HCV infection, the 18 and over strategy was associated with an average increase in life expectancy of 0.68 years (0.63 QALY) compared with standard of care screening. The 18 and over strategy dominated (less costly and more effective or lower incremental cost-effectiveness ratio) the 30 and over or 40 and over strategies in the base

analysis and remained associated with incremental cost-effectiveness ratios of less than \$40,000/QALY in one-way sensitivity analyses that assumed reduced linkage to care, absence of mortality benefit from SVR, higher HCV treatment costs (\$130,000), lower HCV prevalence, or greater restrictions on HCV treatment (i.e., restricting treatment to persons with more advanced fibrosis), compared with the base case assumptions. The 18 and over strategy was less costeffective in scenarios in which antiviral treatment was assumed to be half as effective (\$53,500/QALY), when fibrosis progression was assumed to be half as rapid (\$65,500/QALY), and when testing was assumed to be twice as inefficient (i.e., need to screen twice as many patients to identify the same number of HCV-infected persons, \$44,100/QALY). In some sensitivity analyses (e.g., high treatment costs, less rapid fibrosis progression, lower HCV prevalence, lower rates of linkage to care, greater treatment restrictions), the 30 and over strategy was more cost-effective than the 18 and over strategy. The 30 and over strategy performed best relative to the 18 and over strategy in the decreased fibrosis (\$42,800/OALY vs. \$65,500/OALY) and inefficient testing (\$33,900/QALY vs. \$44,100/QALY) scenarios. The 40 and over strategy was dominated in all sensitivity analyses. An analysis of screening in the general adult population by Eckman et al. compared one-time screening of all persons 18 years or older with screening of persons born between 1945 to 1965 (birth cohort screening) or no screening in a 2stage Markov simulation model.²⁴⁸ Unlike the cost-effectiveness analysis by Barocas et al.,²⁴⁷ screening strategies did not include risk-based screening. The Eckman et al. analysis also assumed lower utilities for chronic HCV infection without cirrhosis (0.79, compared with 0.94 in the other analysis), lower costs of DAA therapy (\$24,270 vs. \$69,078 for a full 12 week course), and higher rates of linkage to care (100% vs. 18% to 29%). It did not model HCV incidence (including reinfection) following successful treatment with antiviral therapy. Despite these differences, the Eckman et al. analysis also found expanded HCV screening to be cost-effective compared with birth cohort screening.

In the Eckman et al. analysis, screening all persons 18 years of age and older was associated with an average gain of 0.0022 QALYs compared with birth cohort screening, and 0.0101 QALYs compared with no screening. The incremental cost-effectiveness of the 18 and older strategy versus birth cohort screening was \$11,378/QALY, and the 18 and older strategy dominated no screening. In sensitivity analysis, the incremental cost-effectiveness ratio of the 18 and older strategy versus birth cohort screening exceeded \$50,000/QALY when the HCV prevalence in the non-birth cohort was less than 0.07 percent (base case 0.29%) or when the monthly cost of antiviral therapy exceeded \$28,000. Cost-effectiveness estimates were also sensitive to the age at time of HCV infection (older age at acquisition associated with lower cost-effectiveness).

An analysis based on the HEP-CE model (used in the study by Barocas et al.) estimated effects of nine one-time screening strategies in U.S. persons, focusing on the population 15 to 30 years of age.²⁴⁹ The screening strategies differed on three factors: 1) routine (screen all persons) versus expanded targeted testing (validated HCV screening checklist used to identify high-risk persons) versus current practice (risk-based testing in persons perceived to be at high risk, without the checklist), 2) rapid finger stick versus venipuncture, and 3) screening ordered by physician versus by counselor or tester using standing orders. Testing rates were assumed to be lower with physician ordering and receipt of results higher with rapid testing. Current practice screening rates were assumed to be 5 percent in PWID and 3 percent otherwise. The model was based on treatment with sofosbuvir / ledipasvir or sofosbuvir / velpatasvir with the cost of a course of

treatment ranging from \$71,950 to \$137,820 and SVR rates of 93 percent to 99 percent, depending on cirrhosis status and genotype.

The model found that strategies involving rapid testing dominated strategies involving venipuncture testing. Compared with current practice, counselor-initiated, routine rapid testing identified more cases (20% vs. 5%), resulted in a greater number of patients achieving SVR (18% vs. 2%), and resulted in fewer HCV-related deaths (34% to 31%), with an incremental cost-effectiveness ratio of \$71,000/QALY. In probabilistic sensitivity analyses, the incremental cost-effectiveness ratio with this strategy remained below \$100,000/QALY unless the prevalence of injection drug use was less than 0.59 percent, the HCV prevalence in PWID was less than 16 percent, the reinfection rate was more than 26 cases per 100 person-years, or reflex confirmatory testing was performed following all reactive venipuncture tests. Although physician-ordered, counselor-performed, expanded targeted rapid testing (\$40,000/QALY) and counselor-initiated, routine rapid testing (\$44,000/QALY) were more cost-effective than counselor-initiated, routine rapid testing, average gains in QALYs were lower with these strategies than with the counselor-initiated, routine rapid testing strategy (incremental differences 0.0008 to 0.0011 QALYs).

Two studies focused on prenatal HCV screening.^{250,251} An analysis by Tasillo et al. evaluated prenatal screening using the HEP-CE model.²⁵¹ The analysis compared universal one-time screening during pregnancy versus current practice (14% screened during pregnancy); both strategies lifetime testing that occurred following pregnancy. The model assumed that therapy with a DAA regimen would be offered 6 months postpartum, with a base cost of \$39,600 for glecaprevir / pibrentasvir (for persons without cirrhosis) and \$68,773 for sofosbuvir / velpatasvir (for persons with cirrhosis). The analysis did not include neonatal outcomes in cost-effectiveness estimates or model the lifetime of neonates born with HCV infection, but estimated the proportion of neonates identified as exposed to HCV infection. HCV prevalence in pregnancy was assumed to be 0.38 percent; assumptions regarding HCV incidence, utilities associated with HCV infection, and rates of linkage to care were similar to the study by Barocas et al. on HCV screening in the general adult population.

The Tasillo et al. analysis found prenatal screening associated with earlier diagnosis and time to cure of HCV infection, with 27 percent of cases achieving SVR within 5 years and 36 percent within 10 years (compared with 16% and 37%, respectively, with current practice). Prenatal screening was associated with a 16 percent reduction in HCV-attributable mortality over the lifetime of the cohort, and average gains of 0.002 QALYs in the entire cohort and 0.0.5 QALYs in HCV-infected persons compared with current practice, with an incremental cost-effectiveness ratio of \$41,000/QALY. The incremental cost-effectiveness ratio was \$83,000/QALY when prevalence was half (0.18%) of the base case assumption (0.18%) and less than or equal to \$50,000/QALY when HCV testing rates were higher (50%) in PWID, when treatment initiation rates were lower (64.5%), and when neonatal testing costs were considered. The incremental cost-effectiveness ratio was \$168,000/QALY when the rate of fibrosis progression was reduced by half (average time to cirrhosis, 70 years) and \$137,000/QALY when HCV infection before cirrhosis had no associated cost or decrease in quality of life. Prenatal screening increased the identification of neonates exposed to HCV at birth from 44 percent to 92 percent.

An analysis by Chaillon et al. also evaluated prenatal screening versus risk-based screening, using a closed cohort Markov model.²⁵⁰ The analysis assumed antiviral treatment after pregnancy with a DAA regimen (base cost \$25,000 for a full treatment course) and a background testing and linkage rate of 5 percent per year; it did not model costs or effects on the neonate. Compared with the analysis by Tasillo et al., base case assumptions in Chaillon et al. included higher HCV prevalence (0.73% vs. 0.38%), lower antiviral treatment costs (\$25,000 vs. \$39,600 in persons with cirrhosis and \$68,773 in persons without cirrhosis), and lower utilities for F1 to F3 fibrosis in HCV-infected persons (0.83-0.86 vs. 0.94). In addition, the model appeared to assume that all persons diagnosed with HCV infection would be linked to care and receive treatment.

In the Chaillon et al. analysis, prenatal screening was estimated to result in the detection and treatment of 7,000 additional females, with an average gain of 0.019 QALY and an incremental cost-effectiveness ratio of \$2,826/QALY, compared with risk-based screening. Incremental cost-effectiveness ratios remained below \$5,000/QALY in sensitivity analyses based on alternative treatment eligibility scenarios, lower HCV prevalence rates (0.03% to 0.04%), lower fibrosis progression rates (21% cirrhosis at 35 years), lower SVR (85%), higher baseline rates of diagnosis and linkage to care (40%), higher loss to followup (50% per year), and higher background testing (20% per year). Screening was estimated to result in detection and treatment of an estimated 300 children born to mothers infected by HCV.

Identification and treatment of HCV infection prior to pregnancy could result in the additional benefit of reducing the risk of mother-to-child transmission following successful treatment.²⁵² However, we identified no study on the cost-effectiveness of screening strategies aimed at women prior to pregnancy.

Contextual Question 3. What Is the Effect of Antiviral Treatments on Behavioral Outcomes?

No trial of DAA therapy included in this report reported behavioral outcomes. Two open-label studies of HCV-infected PWID found receipt of interferon-based therapy associated with reductions in some self-reported drug and substance use behaviors.^{253,254} A non-randomized study (n=124) found interferon-based therapy associated with reduced likelihood of injection drug use equipment sharing (adjusted OR 0.85, 95% CI, 0.74 to 0.99) compared with no treatment at median followup of 1.8 years after adjusting for age, sex, housing status, education level, employment status, and social functioning level, but no effect on injection drug use in the last 30 days (adjusted OR 1.06, 95% CI, 0.93 to 1.21).²⁵⁴ A before-after analysis of persons with current or past injection drug use (n=93) found decreased likelihood of injection drug use (unadjusted OR 0.89, 95% CI, 0.83 to 0.95) and alcohol use (unadjusted OR 0.56, 95% CI, 0.40 to 0.77) 24 weeks after completing interferon-based therapy compared with prior to therapy, but no difference in likelihood of injection drug use equipment sharing (unadjusted OR 0.87, 95% CI, 0.70 to 1.07).²⁵³

Chapter 4. Discussion

Summary of Review Findings

This report updates prior reviews on HCV screening and treatments in adults, and interventions to prevent mother-to-child transmission.^{2,3,90} It expands upon the prior reviews by adding evidence on adolescents and addressing the benefits and harms of currently recommended all-oral, direct acDAA regimens. As in the prior USPSTF review,² we found no direct evidence on the clinical benefits of screening for HCV versus not screening or on the yield of repeat screening. We also found no new evidence to better evaluate harms of screening; the prior review included studies suggesting potential negative psychological and social effects of screening, but the quality of the evidence was poor. Other evidence reviewed for this update is summarized in **Table 23**.

Since the prior USPSTF recommendation, there has been a major shift in antiviral therapy to use of all-oral DAA regimens without interferon.⁷⁴ At the time of the prior review, standard antiviral therapy for HCV infection for genotype 1 infection was transitioning to boceprevir or telaprevir with pegylated interferon and ribavirin (SVR rates 68% to 72%); for genotypes 2 and 3 standard therapy was pegylated interferon plus ribavirin (SVR rates 68% to 78%).⁹⁰ New evidence indicates that SVR rates with currently recommended all-oral DAA regimens are substantially higher than with prior therapies. Pooled SVR rates ranged from 95.5 percent to 98.9 percent across genotypes; for the three most common genotypes in the United States (1, 2, and 3), pooled SVR rates ranged from 95.5 percent to 98.9 percent. Evidence was most robust for genotype 1 infection (32 trials), the most frequent genotype in the United States (approximately 75%), followed by genotype 4 infection (10 trials); data were limited for other genotypes (4 to 6 trials each). SVR estimates generally exceeded 95 percent when analyses were stratified according to DAA regimen, study quality, inclusion of patients with cirrhosis at baseline (with the exception of genotype 3 infection, which was associated with a lower SVR rate in one trial that included patients with cirrhosis),¹⁶⁵ geographic setting, prior experience with older antiviral regimens, and use of ribavirin. Few trials directly compared a current DAA regimen versus placebo or an older antiviral regimen, but those available supported high DAA regimen effectiveness. In one trial of patients with mixed genotype infection, the SVR rate was 99 percent with sofosbuvir / velpatasvir and 0 percent with placebo,¹³⁹ and in two trials of patients with mixed genotype infection the SVR rate was 98 percent to 99 percent with ombitasvir / paritaprevir / ritonavir / dasabuvir (with or without ribavirin) and 66 percent to 80 percent with telaprevir / pegylated interferon / ribavirin.¹³⁷ Evidence on DAA regimens in adolescents is limited but indicates SVR rates similar to those observed in adults (97% to 100%).^{171,173,175,176,201-203} Some trials of DAA regimens in adolescents evaluated regimens that are not FDA-approved for use in adolescents but that are recommended in adults.

Evidence also indicates that current DAA regimens are associated with fewer harms than older interferon-containing therapies; the duration of treatment is also shorter at 12 weeks (8 weeks for glecaprevir / pibrentasvir or ledipasvir / sofosbuvir in persons with genotype 1 infection who are non-black, HIV-uninfected, and whose HCV RNA level is under 6 million IU/mL)⁷⁴ compared with prior interferon-containing regimens (24 to 48 weeks). The prior review found therapies

with interferon associated with rates of serious adverse events of 8.5 percent to 16 percent and withdrawal due to adverse events of 12 percent to 15 percent.⁹⁰ Interferon-based therapies were also associated with high rates of fatigue (51% to 64%), depression (19% to 22%), influenza-like symptoms (19% to 40%), and other adverse events. Boceprevir and telaprevir containing regimens were associated with increased risk of hematological adverse events compared with pegylated interferon plus ribavirin. Four new randomized trials found DAA regimens associated with slightly increased risk of any adverse event (ARD 8%, for a number needed to harm [NNH] of approximately 13) and nausea (ARD 4%, for a NNH of approximately 25) versus placebo, with no difference in risk of serious adverse events, withdrawal due to adverse events, or specific adverse events (e.g., diarrhea, fatigue, headache, or anemia).^{139,151,164,187} Two trials found DAA regimens associated with decreased risk of any adverse event versus triple therapy with telaprevir (ARD -34%, for a number needed to avoid harm [NNAH] of approximately 3), serious adverse events (ARD -8%, NNAH approximately 12), withdrawal due to adverse events (ARD -9%, NNAH approximately 11), and specific adverse events (NNAH for fatigue, nausea, anemia, and rash ranged from approximately 3 to 6).¹³⁷ Across DAA trials, the pooled rate of any adverse event was relatively high at 73.3 percent, but rates of serious adverse events and withdrawal due to adverse events were low (1.9% and 0.4%, respectively) relative to older interferon-containing regimens. Pooled rates of specific adverse events ranged from 2.4 percent for anemia to 18.4 percent for headache, also lower than observed with interferon-containing therapies. Ombitasvir / paritaprevir / ritonavir / dasabuvir with ribavirin was generally associated with increased rates of adverse events compared with the same regimen without ribavirin, with a marked increase in risk of anemia (pooled rates 8.3% vs. 0.8%). All DAA trials in this report excluded patients with HBV coinfection, and no cases of HBV reactivation were reported. One cohort study of VA patients with HCV infection treated with a DAA regimen (n=34,632) that did not meet inclusion criteria reported an HBV reactivation rate of 30.0 per 1,000 person-years.²⁵⁵ Eleven percent of patients in this cohort were surface antigen of HBV-positive at baseline. The HBV reactivation rate with DAA therapy was similar to the reactivation rate with pegylated interferon plus ribavirin (25.4 per 1,000 person-years, p=0.8).

Direct evidence on the effects of antiviral therapy on clinical outcomes is limited. Although several randomized trials found interferon therapy associated with decreased risk of HCC compared with no antiviral therapy, they did not meet inclusion criteria for this report because they focused on patients with cirrhosis at baseline or used a non-standard (i.e. indefinite duration of treatment) regimen.¹²⁷⁻¹³⁴ Trials of DAA therapies were not designed to assess effects on mortality or other long-term clinical outcomes. Ten DAA trials reported improvements in some quality of life and functional outcomes following treatment compared with prior to treatment, but differences were small, studies were open-label, and there was no non-DAA comparison group, making it difficult to interpret more subjective outcomes like these.¹³⁵⁻¹³⁷ Large cohort studies conducted on a large national VA database in which approximately 20 percent of patients had cirrhosis at baseline found DAA therapy associated with reduced risk of cardiovascular events, HCC, and mortality versus no therapy after adjusting for potential confounders, with effects similar to or stronger than interferon-based therapy.^{169,170} A French study found no association between DAA therapy versus no antiviral therapy in risk of all-cause mortality or HCC in the subgroup of patients without cirrhosis at baseline, but there were few events, and estimates were imprecise.¹⁶⁸ In this study, when patients with cirrhosis (approximately 33% of the population) were included in the analysis, DAA therapy was associated with decreased risk of all-cause

mortality (adjusted HR 0.48, 95% CI, 0.33 to 0.70), liver-related mortality (adjusted HR 0.39, 95% CI, 0.21 to 0.71), and HCC (adjusted HR 0.66, 95% CI, 0.46 to 0.93).

No study evaluated effects of DAA therapies on behaviors associated with HCV acquisition. There was limited evidence that interferon-based therapies are not associated with increased injection drug use behaviors, and may be associated with reductions in some behaviors.^{253,254} No study evaluated effects of DAA therapy on HCV transmission.²⁵⁶ Such studies would be difficult to design and carry out, but assessments of potential transmission effects could be informed by modeling studies.^{257,258} One study that modeled effects on transmission risk estimated that among PWID, decreasing HCV prevalence in half within 15 years would require increasing the proportion of patients treated 2- to 15-fold, depending on the baseline HCV prevalence.²⁵⁹

In lieu of limited direct evidence on the effects of antiviral therapy on clinical outcomes, cohort studies of SVR after antiviral therapy versus no SVR may help to understand potential clinical effects. Our findings of a consistent association between SVR after antiviral therapy and improved clinical outcomes were consistent with the prior review.⁹⁰ Moreover, our findings may be more applicable to screening because we excluded previously utilized studies in which a high proportion of patients had cirrhosis at baseline. SVR after antiviral therapy (primarily interferon-based therapy) was associated with decreased risk of all-cause mortality (pooled adjusted HR 0.40, 95% CI, 0.28 to 0.56), liver mortality (pooled adjusted HR 0.11, 95% CI, 0.04 to 0.27), cirrhosis (pooled adjusted HR 0.36, 95% CI, 0.33 to 0.40), and HCC (pooled adjusted HR 0.29, 95% CI, 0.23 to 0.38). Evidence was most robust for all-cause-mortality and HCC (reported in 13 and 20 studies, respectively), and less robust for liver mortality and cirrhosis (reported in 4 studies each). Findings were consistent when studies were stratified according to how well they adjusted for potential confounders, duration of followup, and geographic setting (United States or Europe vs. Asia), though effects on mortality were stronger in studies with longer followup.

Although most studies on the association between SVR after antiviral therapy and clinical outcomes evaluated interferon-based therapy, results were similar in two studies of SVR after DAA therapy,^{205,221} with one study showing similar effects of DAA and interferon regimens on HCC risk. Estimates from a third study of SVR after DAA therapy were very imprecise. This is consistent with a recent systematic review that found no evidence for differential hepatocellular occurrence or recurrence risk following SVR from DAA or interferon-based therapy, though most studies in that review evaluated patients with cirrhosis or a history of HCC.²⁶⁰

Our findings regarding the benefits and harms of current DAA regimens were consistent with a recent systematic review that also reported high SVR rates (greater than 95%) in patients with HCV genotype 1 infection without cirrhosis, high SVR rates but limited evidence for other HCV genotypes, low rates of serious adverse events and treatment discontinuation rates, and higher adverse event rates with ribavirin.⁷³ Our results are also consistent with a systematic review that found insufficient evidence from clinical trials to determine effects of DAA regimens on HCV-related mortality and morbidity;²⁶¹ unlike that review, we also evaluated the indirect chain of evidence linking DAA therapy with clinical outcomes. Our review is consistent with prior reviews that found a consistent association between an SVR after antiviral therapy and reduced risk of mortality and HCC.^{72,260,262-264} Our review differs from prior reviews in focusing on populations more likely to be identified by screening, by excluding studies in which a high

proportion of patients had cirrhosis, and by restricting inclusion to currently recommended DAA regimens. One review on effects of antiviral therapy on extrahepatic manifestations of HCV infection found SVR after antiviral therapy associated with increased likelihood of cryoglobulinemia vasculitis remission and malignant B-cell lymphoproliferative disease response, outcomes not considered in our review because they relate to symptomatic and uncommon conditions.²⁶² It also found attaining SVR associated with reduced risk of insulin resistance and a protective effect on diabetes incidence; we restricted analysis of the association between SVR versus no SVR to mortality and long-term hepatic outcomes and did not identify any studies on the effects of DAA therapy versus no therapy on diabetes.

New evidence on interventions to reduce the risk of mother-to-infant transmission of HCV was limited and did not change the conclusion from the prior review that no intervention has been clearly demonstrated to reduce risk.³ All studies were observational; in addition, we excluded most of the studies in the prior review because they were poor quality and did not conduct multivariate analyses. Studies on the effects of cesarean versus vaginal delivery (5 studies, 1 new)¹⁰⁷ and breastfeeding versus no breastfeeding (3 studies, 1 new)¹⁰⁷ continued to show inconsistent effects on risk of mother-to-child transmission. Although use of internal fetal monitoring and prolonged rupture of membranes were both associated with markedly increased risk of mother-to-child transmission, each was evaluated in only 1 study.¹⁰⁴

Evidence to determine the yield of alternative screening strategies remains limited. Although one new study found that risk-based screening would identify slightly more HCV cases and require testing of fewer patients than birth cohort screening, this was based on a retrospective analysis and the assumption of perfect implementation of risk-based testing, which has not been attained in clinical practice.⁹⁹ Modeling studies suggest that expanded screening strategies may be costeffective in the general population as well as in pregnant females. Two studies found expanded screening of all persons 18 years and older associated with incremental cost-effectiveness ratios under \$30,000/OALY compared with birth cohort screening, despite different assumptions regarding utilities associated with chronic HCV virus infection states, costs of DAA therapy, and rates of linkage to care. In most sensitivity analyses, incremental cost-effectiveness ratios remained less than \$50,000/QALY.^{247,248} Another study found routine HCV screening of persons 15 to 30 years of age associated with incremental cost-effectiveness ratios less than \$50,000/QALY under certain scenarios.²⁴⁹ Two modeling studies found routine prenatal screening associated with incremental cost-effectiveness ratios of \$50,000/QALY versus current practice, though there was more variability in estimates (\$2,826/QALY and \$41,000/QALY).^{250,251} Both studies assumed that antiviral treatment was withheld until after childbirth and did not attempt to model effects on neonatal costs or outcomes. A factor complicating interpretation of the cost-effectiveness analyses are marked differences in base-case assumptions regarding costs of DAA therapy (range approximately \$25,000 [similar to the current cost of a full course of therapy with a generic DAA regimen]²⁶⁵ to over \$100,000), though expanded HCV screening appeared cost-effective even in analyses that assumed high DAA therapy costs. Costs of DAA therapy are expected to decline further, ²⁶⁶⁻²⁶⁸ which would further enhance the cost-effectiveness of expanded screening strategies.

Limitations

Our report has potential limitations. Because there were few trials of current DAA regimens versus placebo or older antiviral therapies, we utilized non-randomized trials of DAA therapies, including trials without a non-DAA therapy comparison group. Pooled SVR rates derived from such trials were considered highly informative because SVR rates are very objective, and SVR rates without treatment are close to zero. However, more subjective outcomes such as quality of life, function, and adverse events are more difficult to interpret in the absence of randomization or a comparison group. SVR is a well-established marker for sustained viral clearance (HCV infection cure) but is an intermediate (non-clinical) outcome. There was little evidence directly evaluating effects of antiviral therapies versus no antiviral therapy on clinical outcomes, due in part to the long duration required to evaluate effects on mortality and other long-term sequelae of HCV infection and ethical considerations related to withholding recommended treatment in randomized trials. Therefore, we included cohort studies on the association between SVR versus antiviral therapy versus no SVR and effects on clinical outcomes. Because such studies are susceptible to residual confounding if other factors associated with achieving an SVR also predict better outcomes, we restricted inclusion to studies that reported multivariate risk estimates and performed stratified analyses based on the degree to which studies adjusted for potential confounders.²⁶⁹ No trial of DAA therapy was conducted in screen-detected patients, and few trials reported presence or severity of baseline symptoms. In order to evaluate effectiveness of DAA therapies in populations likely to be identified by screening, we focused on studies in which patients with cirrhosis, who are more likely to be symptomatic, were excluded, or in which the proportion with cirrhosis was small. Although we included trials of patients previously treated with interferon-based therapies or boceprevir or telaprevir with pegylated interferon and ribavirin, who would not be identified by screening, such patients may be asymptomatic or mildly asymptomatic, and SVR rates were similar in treatment-naïve and -experienced patients. Trials of DAA therapy could overestimate SVR rates compared with typical clinical practice. However, observational studies, including a study of difficult to treat persons in a safety net health system, report SVR rates of 90 percent, or only modestly lower than observed in the trials.^{270,271} We did not assess effects of counseling or immunizations on clinical outcomes in persons diagnosed with HCV infection, though prior reviews found no evidence to estimate effects,⁹¹ and no study evaluated effects of DAA treatments on HCV transmission. We excluded studies of patients coinfected with HBV or HIV and with advanced renal disease since management of these conditions was determined to be outside the scope of screening. We excluded non-English language articles, which could result in language bias, though we identified no non-English language studies that would have met inclusion criteria. We did not search for studies published only as abstracts. We did not formally assess for publication bias using graphical or statistical methods to detect small sample effects due to the small number of randomized trials meeting inclusion criteria; the usefulness of such methods when assessing event rates (rather than risk estimates) is uncertain.

Emerging Issues/Next Steps

All DAA regimens currently recommended were approved by the FDA since the prior review. DAA regimens continue to evolve and treatment guidelines are regularly updated.⁷⁴ Several

newer DAA regimens are pangenotypic, meaning that they are effective across all genotypes,⁷⁵ and most currently recommended DAA regimens do not require use of ribavirin. Although three pangenotypic regimens (glecaprevir / pibrentasvir, sofosbuvir / velpatasvir, and sofosbuvir / velpatasvir / voxilaprevir) have been approved by the FDA, one regimen (sofosbuvir / velpatasvir / voxilaprevir) was developed for use in previously treated persons with resistant virus.²⁷² Advantages of pangenotypic regimens include elimination of the need for genotyping and simplified selection of therapy. Costs of current DAA regimens has been a barrier to treatment but competition and negotiated pricing have reduced prices.^{266,267} Another issue is the shift towards management of HCV infection in primary care settings rather than in specialty settings, potentially facilitating access to treatment. Initial studies indicate that treatment in primary care settings is associated with similar outcomes as treatment in specialty settings, though more data are needed.^{78,79}

Relevance for Priority Populations, Particularly Racial/Ethnic Minorities and Older Adults

In the 2003 to 2010 NHANES survey, persons 40 to 49 years of age (OR 6.0, 95% CI, 3.2 to 11.1) and those 50 to 59 years of age (OR 9.5, 95% CI, 5.3 to 16.8) were more likely to have HCV infection than persons 20 to 39 years of age.¹⁸ Subgroup analyses from trials of currently recommended DAA therapies indicate similar effectiveness in older (over 55 or over 65 years of age) versus younger adults (**Table 13**). Older patients who acquired HCV infection as a young adult are more likely to have more advanced disease due to longer duration of infection, and the HCV-related mortality rate is highest in persons 55 to 74 years of age. Therefore, antiviral therapy may have greater impact on clinical outcomes in older patients.²⁷³

Subgroup analyses from trials of current DAA therapies also indicate similar effectiveness among different racial and ethnic groups. An analysis of the national VA ERCHIVES database (n=21,095) that did not meet inclusion criteria found that SVR rates with DAA regimens were similar in black patients (90%), Hispanic patients (86%), white patients (90%), and Asian/Pacific Islander/American Indian/Alaska Native patients (91%).²⁷¹ However, black patients and Hispanic patients were less likely to achieve SVR than white patients after adjusting for baseline characteristics (OR 0.77, p<0.001 and OR 0.76, p<0.007, respectively).

Most trials of DAA therapies have excluded persons with current drug use or those receiving treatment for opioid use disorder. However, five trials included in this report of persons with current or recent use of methadone or buprenorphine for opioid use disorder reported SVR rates that ranged from 90 to 100 percent.^{149,150,167,192} This is consistent with a systematic review that included observational studies, which found a pooled SVR rate of DAA treatment of almost 90 percent among patients with current or recent injection drug use.²⁷⁴ A systematic review of 57 studies found a 5-year HCV reinfection rate of 10.67 percent in PWID following SVR, compared with 0.95 percent in non-PWID, indicating the need for followup after treatment in this population.²⁷⁵ Current guidelines do not consider ongoing injection drug use a contraindication to DAA therapy.⁷⁴

Although DAA therapy appears similarly effective in adolescents and adults, only three antiviral therapies (ledipasvir / sofosbuvir, sofosbuvir / ribavirin, and glecaprevir / pibrentasvir) are FDA-approved for use in adolescents. Though DAA treatment options in this population are currently limited, a number of trials of DAA regimens in adolescents are ongoing.²⁷⁶

Antiviral therapy is currently not recommended in pregnancy. However, prenatal screening could identify HCV-infected women who could benefit from treatment following pregnancy, facilitate testing of infants, and potentially prevent HCV transmission during subsequent pregnancies. Identification of HCV-infected women prior to pregnancy in order to initiate antiviral therapy could be a strategy to reduce risk of mother-to-child transmission, but has not yet been studied.

Future Research

Research is needed to better understand the association between use of current DAA therapy and clinical outcomes. Long-term randomized trials of treatment versus no treatment would be ethically challenging and difficult to carry out. Rather, large cohort studies that measure important confounders could be highly informative for addressing this question. Trials and cohort studies that measure effects on quality of life, function, and extrahepatic effects of HCV infection (e.g., renal function, cardiovascular effects, or diabetes) would also be helpful for understanding effects of DAA regimens on shorter-term clinical outcomes. Studies on the association between SVR after DAA therapy and clinical outcomes would help to verify the link between SVR and clinical outcomes with current therapies. Additional studies would be helpful for confirming the effectiveness of DAA regimens in adolescents and to identify additional regimens that could be used in this population.²⁷⁶ Studies are also needed to understand risks of HCV reinfection following DAA therapy and optimal treatment strategies. Research is also needed to identify labor management practices (e.g., prolonged rupture of membranes or use of internal fetal monitoring) and other strategies (e.g., identification and treatment of HCV infection prior to pregnancy) on risk of mother-to-child transmission. Well-designed prospective studies are needed to understand the effects of different HCV screening strategies, including repeat screening, on diagnostic yield.

Conclusions

The USPSTF previously determined that HCV screening is highly accurate. Currently recommended all-oral DAA regimens are associated with very high SVR rates (95.5% to 98.9% across genotypes) and few harms relative to older antiviral therapies. An SVR after antiviral therapy is associated with improved clinical outcomes compared with no SVR after adjusting for potential confounders. Direct evidence on the benefits of HCV screening remains unavailable; direct evidence on the effects of antiviral therapy on clinical outcomes remains limited but indicates improved long-term outcomes.

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- 1a. Does screening for hepatitis C virus (HCV) infection in pregnant and nonpregnant adolescents and adults without known abnormal liver enzyme levels reduce HCV-related mortality and morbidity or affect quality of life?
- 1b. Does prenatal screening for HCV infection reduce risk of vertical transmission of HCV infection?
- 2. What is the effectiveness of different risk- or prevalence-based methods for screening for HCV infection on clinical outcomes?
- 3. What is the yield (number of new diagnoses per tests performed) of one-time versus repeat screening or alternative screening strategies for HCV infection, and how does the screening yield vary in different risk groups?
- 4. What are the harms of screening for HCV infection (e.g., anxiety and labeling)?
- 5. What are the effects of interventions during labor and delivery or the perinatal period on risk of vertical transmission of HCV infection?
- 6. What is the effectiveness of currently recommended antiviral treatments in improving health outcomes in patients with HCV infection?
- 7. What is the effectiveness of currently recommended antiviral treatments in achieving a sustained virologic response in patients with HCV infection?
- 8. What are the harms of currently recommended antiviral treatments?
- 9. What is the association between experiencing sustained virologic response following antiviral treatment and reduction in risk of HCV-related adverse health outcomes?

Abbreviations: HCV = hepatitis C virus; SVR = sustained virologic response.

Note: The numbers in the figure correspond to the numbers of the Key Questions.

^{*} Includes persons without abnormal laboratory values. Adolescents are defined as those ages 12 to 17 years. Excludes persons living with HIV, transplant recipients, and patients with renal failure.

[†] Defined as HCV antibody testing with confirmatory HCV RNA testing as indicated.

[‡] Includes interventions that may affect vertical transmission of HCV, such as cesarean delivery, amniocentesis, fetal monitoring, management of ruptured membranes, breastfeeding, and antiviral treatment.

				Ge	enotype 1			
Author Year	Country	Age	% Female	Cirrhosis %	Tx naïve	Events/Total		Proportion (95% CI)
edipasvir/sofosbuvir tidahl 2014 cowdley 2014a awitz 2014b Chuang 2016 im 2016 Wei 2018 Subtotal (I ² = 88.7%, p =	Multinational U.S. U.S. Taiwan Korea China = 0.000)	52 53 48 55 54 47	41 41 38 58 43 50	0% 0% ≤20% ≤20% ≤20%	Yes Yes Yes Mixed Yes No	357/357 408/431 58/60 83/85 46/46 206/206		1.000 (0.990, 1.000) 0.947 (0.921, 0.966) 0.967 (0.885, 0.996) 0.976 (0.918, 0.997) 1.000 (0.922, 1.000) 1.000 (0.982, 1.000) 0.994 (0.952, 0.999)
Simeprevir/sofosbuvir awitz 2014a Śwo 2016 Pott-Junior 2019 Subtotal (l² = 0.0%, p =	U.S. Canada & U.S. Brazil 0.503)	56 56 53	30 47 48	0% 0% 0%	Mixed Mixed Mixed	61/64 150/155 56/60		0.953 (0.869, 0.990) 0.968 (0.926, 0.989) 0.933 (0.838, 0.982) 0.957 (0.926, 0.975)
Sofosbuvir/velpatasvir Everson 2015 Feld, 2015 Vei 2019b Subtotal (l² = 26.6%, p =	U.S. Multinational Multinational = 0.256)	49 54 45	39 60 47	0% 0% ≤20%	Yes Mixed No	28/28 251/255 129/129		■ 1.000 (0.877, 1.000) ■ 0.984 (0.960, 0.996) ■ 1.000 (0.972, 1.000) ■ 0.990 (0.954, 0.998)
Elbasvir/grazoprevir Sulkowski 2015 euzem 2015 Sperl 2016/Ng 2018 Kumada 2017 Wei 2019a Subtotal (l² = 54.6%, p =	Multinational Multinational Japan Multinational = 0.066)	51 52 48 61 48	51 46 57 62 56	0% ≤20% ≤20% ≤20%	Yes Yes Mixed Mixed Yes	122/129 273/288 122/123 219/227 422/432		0.946 (0.891, 0.978) 0.948 (0.916; 0.971) ■ 0.992 (0.956; 1.000) 0.965 (0.932; 0.985) 0.977 (0.958; 0.989) 0.967 (0.950; 0.978)
Sofosbuvir/daclatasvir Sulkowski 2014 Pott-Junior 2019 Subtotal (l² = 45.3%, p =	U.S. Brazil = 0.176)	55 56	51 53	≤20% 0%	Yes Mixed	80/82 65/65	-	0.976 (0.915, 0.997) ■ 1.000 (0.945, 1.000) ■ 0.986 (0.947, 0.997)
Slecaprevir/pibrentasvir Poordad 2017 Chayama 2018 Zeuzem 2018 Subtotal (l² = 77.9%, p =	U.S. Japan Multinational = 0.011)	58 64 53	18 64 51	0% Unclear 0%	No Mixed No	46/50 128/129 663/667		0.920 (0.808, 0.978) 0.992 (0.958, 1.000) 0.994 (0.985, 0.998) 0.986 (0.941, 0.997)
Ombitasvir/paritaprevir/ri alezari 2015 Grebely 2018b Subtotal (l² = 26.7%, p =	itonavir/dasabuvir U.S. Multinational = 0.243)	48 48	34 23	0% 0%	Mixed Yes	37/38 73/80		+ 0.974 (0.862, 0.999) 0.913 (0.828, 0.964) 0.932 (0.870, 0.966)
Ombitasvir/paritaprevir/ri eld 2014 erenci 2014 PIV Kowdley 2014 Ore 2016 M1 Ore 2016 M2 Subtotal (l² = 77.2%, p =	itonavir/dasabuvir - ge Multinational Multinational Multinational Multinational Multinational = 0.002)	enotype 49 51 50 46 47	= 1a 43 35 42 39 46	Unclear 0% 0% 0% 0%	Yes Yes Mixed Mixed Mixed	307/322 282/305 183/212 67/69 19/19		0.953 (0.924, 0.974) 0.925 (0.889; 0.952) 0.863 (0.809; 0.906) 0.971 (0.899; 0.996) 1.000 (0.824; 1.000) 0.937 (0.890; 0.965)
Ombitasvir/paritaprevir/ri Indreone 2014 erenci 2014 PIII Sowdley 2014 Kumada 2015 awitz 2015 Dore 2016 M2 Dore 2016 M2 Subtotal (I ² = 68.5%, p =	tonavir/dasabuvir - ge Multinational Multinational Multinational Japan Multinational Multinational Multinational Multinational = 0.002)	enotype 54 49 48 50 61 55 46 47	= 1b 45 43 54 42 63 51 54 46	0% Unclear 0% 0% 0% 0% 0%	No Yes Mixed Mixed Mixed Mixed Mixed	176/179 148/151 416/419 113/113 204/215 76/82 164/167 81/82		0 983 (0 952 0 997) 0 980 (0 943; 0 996) 0 993 (0 979 0 999) 1 000 (0 968, 1 000) 0 949 (0 910, 0 974) 0 927 (0 848 0 9973) 0 982 (0 948, 0 996) 0 988 (0 934, 1 000) 0 988 (0 944, 0 991)
leterogeneity between g Overall (l² = 81.623%, p	groups: p = 0.005 = 0.000)						 	0.977 (0.966, 0.984)

Abbreviations: CI = confidence interval; NR = not reported; SVR = sustained virologic response; Tx = treatment; U.S. = United States.



Figure 3. Key Question 7: Direct Acting Antiviral Regimens and Pooled Sustained Virologic Response Rates, Genotype 2

Abbreviations: CI = confidence interval; SVR = sustained virologic response; Tx = treatment; U.S. = United States.

Genotype 3											
Author Year	Country	Age	% Female	Cirrhosis %	Tx naïve	Events/Total	Proportion (95% CI				
Sofosbuvir/velpata	svir										
Everson 2015	U.S.	50	37	0%	Yes	25/27	0.926 (0.757, 0.991				
Foster 2015	U.S.	49	39	0%	Mixed	191/197	0.970 (0.935, 0.989				
Pianko 2015	Australia, NZ, U.S.	55	34	0%	No	53/53	⊢ ∎ 1.000 (0.933, 1.000				
Wei 2019b	Multinational	45	47	≤20%	No	72/84	0.857 (0.764, 0.924				
Subtotal (I ² = 82.5	3%, p = 0.001)						0.956 (0.871, 0.986				
Sofosbuvir/daclate	asvir										
Nelson 2015	U.S.	55	41	0%	Mixed	105/109	- 0.963 (0.909, 0.990				
Zeuzem 2018 E3	Multinational	49	55	0%	Yes	111/115	0.965 (0.913, 0.990				
Subtotal (I ² = 0.0	%, p = 0.935)						0.964 (0.930, 0.982				
Glecaprevir/pibrer	ntasvir										
Zeuzem 2018 E3	Multinational	47	41	0%	Yes	149/157	0.949 (0.902, 0.978				
Untergraphic bot	$a_{1000} = 0.79$	4									
	ween groups. $p = 0.78$	4									
Overall (1* = 65.6	u4‰, p = 0.008)						Ψ 0.955 (0.916, 0.977				

Figure 4. Key Question 7: Direct Acting Antiviral Regimens and Pooled Sustained Virologic Response Rates, Genotype 3

Abbreviations: CI = confidence interval; NZ = New Zealand; SVR = sustained virologic response; Tx = treatment; U.S. = United States.

					SVR						
				(Genotype	4					
Author Year	Country	Age	% Female	Cirrhosis %	TX naïve	Events/Total				Proportio	n (95% CI)
Ledipasvir/sofosbuvir										1	
Abergel 2016a	France	52	50	≤20%	Yes	21/22				■ 0.955 (0.)	772. 0.999)
Ahmed 2018	Egypt	51	35	Unclear	Yes	99/100				- 0.990 (0.	946, 1.000)
Subtotal (I ² = 24.6%,	p = 0.250)									0.984 (0.	937, 0.996)
Sofosbuvir/velpatasvir											
Feld 2015	Multinational	54	60	0%	Mixed	89/89				■ 1.000 (0.5	959, 1.000)
Elbasvir/grazoprevir											
Zeuzem 2015	Multinational	52	46	≤20%	Yes	18/18				1.000 (0.	815, 1,000)
Sperl 2016/Ng 2018	Multinational	48	57	≤20%	Mixed	6/6				1.000 (0.	541, 1,000)
Brown 2018	Multinational	52	58	0%	Yes	9/10				0.900 (0.	555, 0.997)
Wei 2019a	Multinational	48	56	≤20%	Yes	3/3				1 000 (0.	292 1 000)
Subtotal (I ² = 0.0%, p	= 0.610)								•	0.973 (0.	832, 0.996)
Glecaprevir/pibrentasv	vir									-	
Asselah 2018	Multinational	48	36	0%	Mixed	43/46				0.935 (0.3	821, 0.986)
Ombitasvir/paritaprevi	r/ritonavir/dasa	buvir									
Hezode 2015	Multinational	48	29	0%	Mixed	91/91				1.000 (0.	960, 1.000)
Waked 2016	Egypt	49	30	≤20%	Yes	94/100			-1	0.940 (0.	874, 0.978)
Subtotal (I ² = 87.5%,	p = 0.005)									Q 0.987 (0.	727, 0.9995)
Heterogeneity betwee	n groups: p =	0.137									
Overall (I ² = 50.294%)	, p = 0.034)									0.982 (0.9	947, 0.994)
								1	1	1	
					Droportion	(.25	.5	.75	1	

Abbreviations: CI = confidence interval; SVR = sustained virologic response; Tx = treatment.

					Genotype	5	1		
Author Year	Country	Age	% Female	Cirrrhosis %	Tx naïve	Events/Total			Proportion (95% CI)
Ledipasvir/sofos	buvir								
Abergel 2016b	France	61	48	≤20% cirrhosis	Yes	20/21			0.952 (0.762, 0.999)
Sofosbuvir/velpa	ıtasvir								
Feld 2015	Multinational	54	60	0% cirrhosis	Mixed	28/29			0.966 (0.822, 0.999)
Glecaprevir/pibr	entasvir								
Asselah 2018	Multinational	48	36	0% cirrhosis	Mixed	2/2			1.000 (0.158, 1.000)
Asselah 2019	Multinational	68	57	≤ 0% cirrhosis	Mixed	22/23			0.957 (0.781, 0.999)
Subtotal (I ² = 0	.000%, p = 0.86	0)						<	0.960 (0.764, 0.994)
Heterogeneity b	etween groups:	p = 0.	956						
Overall (I ² = 0.0	000%, p = 0.989)						<	0.960 (0.883, 0.987)
							1	 	1 1

Figure 6. Key Question 7: Direct Acting Antiviral Regimens and Sustained Virologic Response Rates, Genotype 5

Abbreviations: CI = confidence interval; SVR = sustained virologic response; Tx = treatment.

				C	Genotype	6				
Author Year	Country	Age	% Female	Cirhosis %	Tx naïve	Events/Total				Proportion (95% CI)
Ledipasvir/sofo	osbuvir								1	
Gane 2015	New Zealand	51	36	Unclear	Mixed	24/25		_	-	0.960 (0.796, 0.999)
Sofosbuvir/velp	oatasvir									
Feld 2015	Multinational	54	60	0% cirrhosis	Mixed	35/35			- •	1.000 (0.900, 1.000)
Wei 2019b	Multinational	45	47	≤20% cirrhosis	No	97/98			-	0.990 (0.944, 1.000)
Subtotal (I² =	0.0%, p = 0.698	3)							ŧ	0.992 (0.949, 0.999)
Glecaprevir/pib	orentasvir									
Asselah 2018	Multinational	48	36	0% cirrhosis	Mixed	9/10	_		•	0.900 (0.555, 0.997)
Asselah 2019	Multinational	54	52	≤20% cirrhosis	Mixed	60/61			-4	0.984 (0.912, 1.000)
Subtotal (I ² =	42.1%, p = 0.18	39)							4	0.972 (0.894, 0.993)
Heterogeneity	between group	s: p =	0.373							
Overall (I ² = 0	.000%, p = 0.42	27)							0	0.982 (0.954, 0.993)
							1		-	

Figure 7. Key Question 7: Direct Acting Antiviral Regimens and Sustained Virologic Response Rates, Genotype 6

Abbreviations: CI = confidence interval; SVR = sustained virologic response; Tx = treatment.

SVR Mixed Genotypes Author Year Age % Female Cirrhosis % Tx naïve Events/Total Proportion (95% CI) Country 1 Sofosbuvir/velpatasvir Everson 2015 U.S. 0% 21/22 0.955 (0.772, 0.999) 54 32 Yes Grebely 2018a Multinational 48 28 0% Unclear 82/86 0.953 (0.885, 0.987) Subtotal (I² = 0.0%, p = 0.829) 0.954 (0.894, 0.981) .25 .5 .75 0 1 Proportion

Figure 8. Key Question 7: Direct Acting Antiviral Regimens and Sustained Virologic Response Rates, Mixed Genotypes

Abbreviations: CI = confidence interval; SVR = sustained virologic response; Tx = treatment; U.S. = United States.

Figure 9. Key Question 8: Direct Acting Antiviral Regimens Versus Placebo, Any Adverse Events

	DAA reg	imen	Place	bo		Risk Ratio	Risk Ratio
Study or Subgroup	Events	Total	Events	Total	Weight	M-H, Random, 95% Cl	M-H, Random, 95% Cl
1.1.1 Ombitasvir / parita	aprevir / ri	tonavir					
Kumada 2015	148	215	60	106	17.9%	1.22 [1.01, 1.47]	
Subtotal (95% CI)		215		106	17.9%	1.22 [1.01, 1.47]	•
Total events	148		60				
Heterogeneity: Not appli	icable						
l est for overall effect: Z :	= 2.02 (P =	= 0.04)					
1.1.2 Ombitasvir / parita	aprevir / ri	tonavir	/ dasabu	vir / rib	avirin		
Feld 2014	414	473	116	158	35.3%	1.19 [1.08, 1.32]	• • • • • • • • • • • • • • • • • • •
Subtotal (95% CI)		473		158	35.3%	1.19 [1.08, 1.32]	•
Total events	414		116				
Heterogeneity: Not appl	icable						
Test for overall effect: Z	= 3.45 (P =	= 0.0006	i)				
1.1.3 Sofosbuvir / velpa	tasvir						
Feld 2015	485	624	89	116	33.0%	1 01 0 91 1 131	•
Subtotal (95% CI)		624		116	33.0%	1.01 [0.91, 1.13]	•
Total events	485		89				
Heterogeneity: Not appl	icable						
Test for overall effect: Z	= 0.23 (P =	: 0.82)					
1 1 A Elbaevir / grazonr	ovir						
	220	301	50	100	12.000	1 1 2 10 00 1 401	_
Subtotal (95% CI)	230	400	52	123	13.8%	1.12 [0.89, 1.40]	•
Total events	230		52				ſ
Heterogeneity: Not appli	icable						
Test for overall effect: Z:	= 0.97 (P =	= 0.33)					
							L.
Total (95% CI)		1798		503	100.0%	1.12 [1.02, 1.24]	•
Total events	1277		317				
Heterogeneity: Tau ² = 0.	00; Chi ² =	5.56, df	= 3 (P =	0.14); P	•=46%		0.01 0.1 1 10 100
Test for overall effect: Z:	= 2.36 (P =	÷U.UZ) i≊ – 6.61	df = 2 /0	0 - 0 1 /	1 12 - 16	504	Favors DAA Favors placebo

Figure 10. Key Question 8: Direct Acting Antiviral Regimens Versus Placebo, Serious Adverse Events

	DAA regi	men	Place	bo		Risk Ratio	Risk Ratio
Study or Subgroup	Events	Total	Events	Total	Weight	M-H, Random, 95% Cl	M-H, Random, 95% Cl
1.2.1 Ombitasvir / parita	previr / rit	onavir					
Kumada 2015	7	215	2	106	38.0%	1.73 [0.36, 8.16]	
Subtotal (95% CI)		215		106	38.0%	1.73 [0.36, 8.16]	
Total events	7		2				
Heterogeneity: Not applic	cable						
Test for overall effect: Z =	: 0.69 (P =	0.49)					
1.2.2 Ombitasvir / parita	previr / rit	onavir	/ dasabu	vir / rib	avirin		
Feld 2014	10	473	0	158	11.5%	7.04 [0.42, 119.53]	
Subtotal (95% CI)		473		158	11.5%	7.04 [0.42, 119.53]	
Total events	10		0				
Heterogeneity: Not applic	cable						
Test for overall effect: Z =	: 1.35 (P =	0.18)					
1.2.3 Sofosbuvir / velpat	asvir						
Feld 2015	15	624	Ο	116	11.6%	5 80 (0 35 96 32)	
Subtotal (95% CI)	10	624		116	11.6%	5.80 [0.35, 96.32]	
Total events	15		0				
Heterogeneity: Not appli	cable						
Test for overall effect: Z =	: 1.23 (P =	0.22)					
124 Elbasvir / grazonre	vir						
Wei 2019 (C-CORAL)		496	2	172	20.0%	1 01 0 22 4 71	
Subtotal (95% CI)	0	400	2	123	38.9%	1.01 [0.22, 4.71]	
Total events	8		2				
Heterogeneity: Not appli	cable		-				
Test for overall effect: Z =	0.02 (P =	0.99)					
Total (95% CI)		1798		503	100.0%	1.90 [0.73, 4.95]	
Total events	40		4				, , , ,
Heterogeneity: Tau ² = 0.0	JU; Chi≝ = : 	2.33, df	= 3 (P =	0.51); P	*=0%		0.01 0.1 1 10 100
i est for overall effect: Z =	: 1.31 (P =	0.19)					Equare DAA Equare placebo

Risk Ratio DAA regimen Placebo Risk Ratio Total Events Total Weight M-H, Random, 95% CI M-H, Random, 95% CI Study or Subgroup Events 1.3.1 Ombitasvir / paritaprevir / ritonavir Kumada 2015 2 215 0 106 14.9% 2.48 [0.12, 51.14] 2.48 [0.12, 51.14] Subtotal (95% CI) 215 106 14.9% 2 0 Total events Heterogeneity: Not applicable Test for overall effect: Z = 0.59 (P = 0.56) 1.3.2 Ombitasvir / paritaprevir / ritonavir / dasabuvir / ribavirin Feld 2014 3 473 1 158 25.1% 1.00 [0.10, 9.56] Subtotal (95% CI) 473 158 25.1% 1.00 [0.10, 9.56] 3 Total events 1 Heterogeneity: Not applicable Test for overall effect: Z = 0.00 (P = 1.00) 1.3.3 Sofosbuvir / velpatasvir Feld 2015 624 1 22.7% 0.09 [0.01, 1.02] 2 116 Subtotal (95% CI) 624 116 22.7% 0.09 [0.01, 1.02] 2 Total events 1 Heterogeneity: Not applicable Test for overall effect: Z = 1.95 (P = 0.05) 1.3.4 Elbasvir / grazoprevir Wei 2019 (C-CORAL) 3 486 37.2% 0.38 [0.06, 2.25] 2 123 Subtotal (95% CI) 486 123 37.2% 0.38 [0.06, 2.25] Total events 3 2 Heterogeneity: Not applicable Test for overall effect: Z = 1.07 (P = 0.29) Total (95% CI) 1798 503 100.0% 0.47 [0.14, 1.58] Total events 9 5 Heterogeneity: Tau² = 0.22; Chi² = 3.48, df = 3 (P = 0.32); l² = 14% 0.01 0.1 10 100 Test for overall effect: Z = 1.23 (P = 0.22) Favors DAA Favors placebo Test for subgroup differences: $Chi^2 = 3.41$, df = 3 (P = 0.33), $l^2 = 11.9\%$

Figure 11. Key Question 8: Direct Acting Antivirals Regimens Versus Placebo, Withdrawals Due to Adverse Events

	DAA reg	imen	Place	bo		Risk Ratio		Risk Ratio	
Study or Subgroup	Events	Total	Events	Total	Weight	M-H, Random, 95% Cl		M-H, Random, 95% Cl	
1.7.1 Ombitasvir / par	ritaprevir /	ritonav	ir						
Kumada 2015	9	215	4	106	9.2%	1.11 [0.35, 3.52]			
Subtotal (95% CI)		215		106	9.2%	1.11 [0.35, 3.52]			
Total events	9		4						
Heterogeneity: Not ap	plicable								
Test for overall effect:	Z = 0.18 (F	^o = 0.86)						
1.7.2 Ombitasvir / par	ritaprevir	ritonav	ir / dasal	buvir / i	ribavirin				
Feld 2014	112	473	21	158	55.0%	1.78 [1.16, 2.74]			
Subtotal (95% CI)		473		158	55.0%	1.78 [1.16, 2.74]		-	
Total events	112		21						
Heterogeneity: Not ap	plicable								
Test for overall effect:	Z = 2.63 (F	P = 0.00	8)						
1.7.3 Sofosbuvir / vel	oatasvir								
Feld 2015	75	674	13	116	35.8%	1 07 0 62 1 871			
Subtotal (95% CI)		624		116	35.8%	1.07 [0.62, 1.87]		•	
Total events	75		13			,		Ť	
Heterogeneity: Not an	nlicable								
Test for overall effect:	Z = 0.25 (F	P = 0.80)						
	(0.00	,						
Total (95% CI)		1312		380	100.0%	1.42 [1.00, 2.03]		◆	
Total events	196		38						
Heterogeneity: Tau ² =	0.01; Chi ^a	²= 2.23,	df = 2 (P	= 0.33)); I ^z = 10%	6			100
Test for overall effect:	Z = 1.94 (ł	P = 0.05)				0.01 0	Eavors DAA Eavors placebo	100
Test for subgroup diff	erences: (>hi ² = 2.	22, df = 2	(P = 0.	.33), I ² = 1	0.0%			

Abbreviations: CI = confidence interval; DAA = direct acting antiviral; df = degrees of freedom; M-H = Mantel-Haenszel test.

	DAA reg	imen	Place	bo		Risk Ratio		Risk Ratio
Study or Subgroup	Events	Total	Events	Total	Weight	M-H, Random, 95% Cl		M-H, Random, 95% Cl
1.5.1 Ombitasvir / pa	ritaprevir	/ ritonav	ir / dasal	buvir / I	ribavirin			
Feld 2014	65	473	11	158	55.8%	1.97 [1.07, 3.64]		
Subtotal (95% CI)		473		158	55.8%	1.97 [1.07, 3.64]		◆
Total events	65		11					
Heterogeneity: Not ap	plicable							
Test for overall effect:	Z=2.17 (P = 0.03)					
1.5.2 Sofosbuvir / vel	patasvir							
Feld 2015	48	624	8	116	44.2%	1.12 [0.54, 2.30]		
Subtotal (95% CI)		624		116	44.2%	1.12 [0.54, 2.30]		•
Total events	48		8					
Heterogeneity: Not ap	plicable							
Test for overall effect:	Z = 0.30 (P = 0.77)					
Total (95% CI)		1097		274	100.0%	1.53 [0.88, 2.68]		•
Total events	113		19					
Heterogeneity: Tau ² =	: 0.05; Chi ^a	² = 1.40,	df = 1 (P	= 0.24)); I ² = 29%)		
Test for overall effect:	Z=1.50 (I	P = 0.13)				0.01	Eavors DAA Eavors placebo
Test for subgroup diff	ferences: (Chi² = 1.	40, df = 1	(P = 0.	24), I ² = 2	8.3%		

Abbreviations: CI = confidence interval; DAA = direct acting antiviral; df = degrees of freedom; M-H = Mantel-Haenszel test.

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Figure 15. Key Question 8: Direct Acting Antivirals Versus Placebo, Headache

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Figure 16. Key Question 8: Direct Acting Antiviral Regimens Versus Telaprevir, Any Adverse Events

	DAA	1	Telapre	evir		Risk Ratio		Risk Ratio
Study or Subgroup	Events	Total	Events	Total	Weight	M-H, Random, 95% Cl		M-H, Random, 95% Cl
2.1.1 Ribavirin								
Dore 2016 (MALACHITE-1)	115	153	37	38	36.6%	0.77 [0.69, 0.86]		•
Dore 2016 (MALACHITE-2)	63	101	43	47	33.0%	0.68 [0.57, 0.81]		
Subtotal (95% CI)		254		85	69.6%	0.74 [0.65, 0.84]		•
Total events	178		80					
Heterogeneity: Tau ² = 0.00; C	hi² = 1.77	', df = 1	(P = 0.18	i); I ² = 4	3%			
Test for overall effect: Z = 4.58	6 (P ≤ 0.0I	0001)						
2.1.2 No ribavirin								
Dore 2016 (MALACHITE-1)	41	83	37	37	30.4%	0.50 [0.40, 0.62]		+
Subtotal (95% CI)		83		37	30.4%	0.50 [0.40, 0.62]		•
Total events	41		37					
Heterogeneity: Not applicable	!							
Test for overall effect: Z = 6.18	8 (P < 0.0)	0001)						
								•
Total (95% CI)		337		122	100.0%	0.65 [0.50, 0.84]		•
Total events	219		117					
Heterogeneity: Tau ^z = 0.04; C	hi² = 14.9	8, df =	2 (P = 0.0	1006); P	²=87%			
Test for overall effect: Z = 3.29	9 (P = 0.0)	010)					0.01 0.1	avors DAA Eavors telaprevir
Test for subgroup differences	: Chi ^z = 8	.83, df	= 1 (P = 0	1.003),	l ^z = 88.7%	6		atore bitt i atore telapreni

Figure 17. Key Question 8: Direct Acting Antiviral Regimens Versus Telaprevir, Serious Adverse Events

	DAA	1	Telapro	evir		Risk Ratio	Risk Ratio
Study or Subgroup	Events	Total	Events	Total	Weight	M-H, Random, 95% Cl	M-H, Random, 95% Cl
2.2.1 Ribavirin							
Dore 2016 (MALACHITE-1)	1	153	5	38	42.9%	0.05 [0.01, 0.41]	←
Dore 2016 (MALACHITE-2)	1	101	2	47	34.1%	0.23 [0.02, 2.50]	
Subtotal (95% CI)		254		85	77.1%	0.10 [0.02, 0.48]	
Total events	2		7				
Heterogeneity: Tau ² = 0.00; C	hi² = 0.91	, df = 1	(P = 0.34)	l); l² = 0)%		
Test for overall effect: Z = 2.88	8 (P = 0.0	04)					
2.2.2 No ribavirin							
Dore 2016 (MALACHITE-1)	0	83	4	37	22.9%	0.05 [0.00, 0.91]	<
Subtotal (95% CI)		83		37	22.9%	0.05 [0.00, 0.91]	
Total events	0		4				
Heterogeneity: Not applicable	9						
Test for overall effect: Z = 2.02	2 (P = 0.0	4)					
Total (95% CI)		337		122	100.0%	0.08 [0.02, 0.34]	
Total events	2		11				
Heterogeneity: Tau ² = 0.00; C	hi² = 1.07	', df = 2	(P = 0.59	3); I ≥ = 0)%		
Test for overall effect: Z = 3.49	9 (P = 0.0	005)					Eavors DAA Eavors telaprevir
Test for subgroup differences	: Chiř = C).16, df	= 1 (P = 0).69), l ^z	= 0%		r atoro britt i atoro totapitum

Figure 18. Key Question 8: Direct Acting Antiviral Regimens Versus Telaprevir, Withdrawal Due to Adverse Events

	DAA	1	Telapre	evir		Risk Ratio	Risk Ratio
Study or Subgroup	Events	Total	Events	Total	Weight	M-H, Random, 95% Cl	M-H, Random, 95% Cl
2.3.1 Ribavirin							
Dore 2016 (MALACHITE-1)	1	153	3	37	45.8%	0.08 [0.01, 0.75]	←
Dore 2016 (MALACHITE-2)	0	101	5	47	27.7%	0.04 [0.00, 0.76]	←
Subtotal (95% CI)		254		84	73.5%	0.06 [0.01, 0.37]	
Total events	1		8				
Heterogeneity: Tau ² = 0.00; C	hi = 0.13	l, df = 1	(P = 0.72)	?); I z = 0	1%		
Test for overall effect: Z = 3.08	6 (P = 0.0)	02)					
2.3.2 No ribavirin							
Dore 2016 (MALACHITE-1)	0	83	3	38	26.5%	0.07 [0.00, 1.25]	←
Subtotal (95% CI)		83		38	26.5%	0.07 [0.00, 1.25]	
Total events	0		3				
Heterogeneity: Not applicable	l						
Test for overall effect: Z = 1.81	(P = 0.0)	7)					
Total (95% CI)		337		122	100.0%	0.06 [0.01, 0.29]	
Total events	1		11				
Heterogeneity: Tau ^z = 0.00; C	hi ^z = 0.12	, df = 2	(P = 0.94	l); I ^z = 0	1%		
Test for overall effect: Z = 3.58	6 (P = 0.0	004)					Eavors DAA Eavors telaprevir
Test for subgroup differences	: Chi ² = 0	1.00, df	= 1 (P = 0).98), I <mark>²</mark>	= 0%		

Figure 19. Key Question 8: Direct Acting Antiviral Regimens Versus Telaprevir, Fatigue

	DAA	1	Telapr	evir		Risk Ratio	Risk Ratio
Study or Subgroup	Events	Total	Events	Total	Weight	M-H, Random, 95% Cl	M-H, Random, 95% Cl
2.5.1 Ribavirin							
Dore 2016 (MALACHITE-1)	21	153	12	38	43.9%	0.43 [0.24, 0.80]	
Dore 2016 (MALACHITE-2)	12	101	12	47	36.0%	0.47 [0.23, 0.96]	
Subtotal (95% CI)		254		85	79.9%	0.45 [0.28, 0.71]	◆
Total events	33		24				
Heterogeneity: Tau ² = 0.00; C	hi² = 0.02	2, df = 1	(P = 0.89	9); I ≥ = 0)%		
Test for overall effect: Z = 3.37	' (P = 0.0	007)					
2.5.2 No ribavirin							
Dore 2016 (MALACHITE-1)	4	83	11	37	20.1%	0.16 [0.06, 0.48]	
Subtotal (95% CI)		83		37	20.1%	0.16 [0.06, 0.48]	
Total events	4		11				
Heterogeneity: Not applicable	!						
Test for overall effect: Z = 3.31	(P = 0.0	009)					
T							
Total (95% CI)		337		122	100.0%	0.37 [0.21, 0.63]	-
Total events	37		35				
Heterogeneity: Tau ² = 0.07; C	hi² = 2.94	l, df = 2	(P = 0.23	3); I 2 = 3	32%		
Test for overall effect: Z = 3.67	' (P = 0.0	002)					Eavors DAA Eavors telaprevir
Test for subgroup differences	: Chi ² = 2	2.87, df	<u>= 1 (P = (</u>).09), I <mark>*</mark>	= 65.2%		

Figure 20. Key Question 8: Direct Acting Antiviral Regimens Versus Telaprevir, Headache

	DAA	1	Telapr	evir		Risk Ratio	Risk Ratio
Study or Subgroup	Events	Total	Events	Total	Weight	M-H, Random, 95% Cl	M-H, Random, 95% CI
2.4.1 Ribavirin							
Dore 2016 (MALACHITE-1)	41	153	12	38	32.0%	0.85 [0.50, 1.45]	
Dore 2016 (MALACHITE-2)	29	101	21	47	47.0%	0.64 [0.41, 1.00]	
Subtotal (95% CI)		254		85	79.0%	0.72 [0.51, 1.01]	▲
Total events	70		33				
Heterogeneity: Tau ² = 0.00; C	hi² = 0.62	?, df = 1	(P = 0.43)	3); I 2 = 0)%		
Test for overall effect: Z = 1.89) (P = 0.0	6)					
2.4.2 No ribavirin							
Dore 2016 (MALACHITE-1)	16	83	11	37	21.0%	0.65 [0.33, 1.26]	
Subtotal (95% CI)		83		37	21.0%	0.65 [0.33, 1.26]	
Total events	16		11				
Heterogeneity: Not applicable	!						
Test for overall effect: Z = 1.28) (P = 0.2	0)					
		227		400	400.0%	0.70 (0.62, 0.05)	
Total (95% CI)		221		122	100.0%	0.70 [0.52, 0.95]	
Total events	86		44				
Heterogeneity: Tau ² = 0.00; C	hi² = 0.69	l, df = 2	(P = 0.71)	1); I ^z = 0)%		
Test for overall effect: Z = 2.27	' (P = 0.0	2)					Favors DAA Favors telaprevir
Test for subgroup differences	: Chi ² = 0	1.07, df	<u>= 1 (P = (</u>),79), l ^z	= 0%		·

Figure 21. Key Question 8: Direct Acting Antiviral Regimens Versus Telaprevir, Nausea

	DAA	1	Telapro	evir		Risk Ratio		Risk Ratio
Study or Subgroup	Events	Total	Events	Total	Weight	M-H, Random, 95% Cl		M-H, Random, 95% Cl
2.6.1 Ribavirin								
Dore 2016 (MALACHITE-1)	32	153	15	38	39.1%	0.53 [0.32, 0.87]		
Dore 2016 (MALACHITE-2)	10	101	20	47	32.7%	0.23 [0.12, 0.46]		
Subtotal (95% CI)		254		85	71.8%	0.36 [0.16, 0.82]		\bullet
Total events	42		35					
Heterogeneity: Tau ² = 0.26; C	hi = 3.78	8, df = 1	(P = 0.05)	5); I 2 = 7	4%			
Test for overall effect: Z = 2.44	4 (P = 0.0	1)						
2.6.2 No ribavirin								
Dore 2016 (MALACHITE-1)	7	83	15	37	28.2%	0.21 [0.09, 0.47]		
Subtotal (95% CI)		83		37	28.2%	0.21 [0.09, 0.47]		
Total events	7		15					
Heterogeneity: Not applicable)							
Test for overall effect: Z = 3.80) (P = 0.0	001)						
Total (95% CI)		337		122	100.0%	0.31 [0.16, 0.59]		-
Total events	49		50					
Heterogeneity: Tau ² = 0.21; C	hi² = 5.78	3, df = 2	(P = 0.06	6); I² = 6	65%			
Test for overall effect: Z = 3.57	7 (P = 0.0	004)					0.01	Eavors DAA Eavors telaprevir
Test for subgroup differences	<u>:: Chi² = 0</u>).90, df	= 1 (P = 0).34), I ^z	= 0%			Tarele Britt Tarele telaprovil

Figure 22. Key Question 8: Direct Acting Antiviral Regimens Versus Telaprevir, Anemia

	DAA	1	Telapro	evir		Risk Ratio	Risk Ratio
Study or Subgroup	Events	Total	Events	Total	Weight	M-H, Random, 95% Cl	M-H, Random, 95% Cl
2.7.1 Ribavirin							
Dore 2016 (MALACHITE-1)	10	153	17	38	52.4%	0.15 [0.07, 0.29]	
Dore 2016 (MALACHITE-2)	3	101	16	47	32.1%	0.09 [0.03, 0.28]	
Subtotal (95% CI)		254		85	84.5%	0.13 [0.07, 0.23]	◆
Total events	13		33				
Heterogeneity: Tau ² = 0.00; C	hi² = 0.59), df = 1	(P = 0.44)	4); I≊ = 0)%		
Test for overall effect: Z = 6.72	2 (P ≤ 0.0	0001)					
2.7.2 No ribavirin							
Dore 2016 (MALACHITE-1)	1	83	17	37	15.5%	0.03 (0.00, 0.19)	← ■
Subtotal (95% CI)		83		37	15.5%	0.03 [0.00, 0.19]	
Total events	1		17				
Heterogeneity: Not applicable	9						
Test for overall effect: Z = 3.61	l (P = 0.0	003)					
Total (95% CI)		337		122	100.0%	0.09 [0.04, 0.23]	\bullet
Total events	14		50				
Heterogeneity: Tau ² = 0.25; C	hi² = 3.38), df = 2	(P = 0.18	3); ² = 4	11%		
Test for overall effect: Z = 5.31	I (P ≤ 0.0	0001)					Eavors DAA Eavors telaprevir
Test for subgroup differences	s: Chi ² = 2	2.26, df	= 1 (P = 0).13), <mark>I</mark> ≊	= 55.7%		ratio bitt ratio trapfom

Figure 23. Key Question 8: Direct Acting Antiviral Regimens Versus Telaprevir, Rash

	DAA	1	Telapro	evir		Risk Ratio	Risk Ratio
Study or Subgroup	Events	Total	Events	Total	Weight	M-H, Random, 95% Cl	M-H, Random, 95% Cl
2.8.1 Ribavirin							
Dore 2016 (MALACHITE-1)	12	153	9	38	51.7%	0.33 [0.15, 0.73]	
Dore 2016 (MALACHITE-2)	3	101	8	47	35.9%	0.17 [0.05, 0.63]	
Subtotal (95% CI)		254		85	87.6%	0.28 [0.14, 0.54]	◆
Total events	15		17				
Heterogeneity: Tau ² = 0.00; C	hi² = 0.72	?, df = 1	(P = 0.40))); I ≃ = 0)%		
Test for overall effect: Z = 3.74	4 (P = 0.0	002)					
2.8.2 No ribavirin							
Dore 2016 (MALACHITE-1)	Ο	83	8	37	12.4%	0.03 (0.00. 0.45)	← ■
Subtotal (95% CI)	Ŭ	83	Ŭ	37	12.4%	0.03 [0.00, 0.45]	
Total events	0		8				
Heterogeneity: Not applicable	e						
Test for overall effect: Z = 2.51	1 (P = 0.0	1)					
Total (95% CI)		337		122	100.0%	0.19 [0.06, 0.58]	
Total events	15		25				
Heterogeneity: Tau ² = 0.44; C	:hi² = 3.82	2, df = 2	(P = 0.15	5); I² = 4	18%		
Test for overall effect: Z = 2.95	5 (P = 0.0	03)					Favors DAA Favors telaprevir
Test for subgroup differences	<u>s: Chi² = 2</u>	<u>2.50, df</u>	<u>= 1 (P = 0</u>).11), I ^z	= 60.1%		

				Δ	Any Adve	rse Event	oc			
ior year	Country	Age	% Female	Fibrosis stage	Tx naïve	Events/Total	Genotype			Proportion (95%
pasvirisofos buvir ini 2014 diev 2014 diev 2014 tz 2040 2016 2016 e 2015 rgel 2016a rgel 2016a rgel 2016a rgel 2016a rgel 2016b	Multinational U.S. U.S. Korea Talwan China New Zealand France Egypt France	52 53 47 47 47 47 51 51	41 36 43 59 50 42 50 35 48	≤20% cirrhosis 0% cirrhosis 20% cirrhosis 420% cirrhosis 420% cirrhosis 420% cirrhosis 420% cirrhosis 420% cirrhosis 420% cirrhosis 420% cirrhosis	Yes Yes Yes Yes Mixed Mixed Yes Yes Yes	169/214 355/431 17/39 46/93 51/60 120/206 46/50 31/44 26/100 33/41	1 1 1 3 or 6 4 5	- <u>-</u>		0.790 (0.729, 0.8 0.824 (0.784, 0.8 0.436 (0.278, 0.6 0.495 (0.389, 0.6 0.850 (0.734, 09 0.583 (0.512, 0.6 0.920 (0.808, 0.9 0.705 (0.548, 0.8 0.260 (0.177, 0.3 0.805 (0.654, 0.9 0.6544, 0.548, 0.3
eprevir/sofos buvir Itz 2014a 2016 Iotal (I² = 0.0%, p = 0.399)	U.S. Canada & U.S.	56 56	42 47	0% cirrhosis 0% cirrhosis	No Mixed	11/14 103/155	1	_	~	0.786 (0.492, 0.9 0.665 (0.584, 0.7 0.675 (0.600, 0.7
sbuvir/velpatasvir ier 2015 A2 ier 2015 A3 ison 2015 (Part A) i 2015 bely 2018a 2019b iotal (I ² = 96.4%, p = 0.000)	U.S. U.S. U.S. Multinational Multinational Multinational	57 49 49 54 48 45	36 39 39 50 28 47	≤20% cirrhosis ≤20% cirrhosis 0% cirrhosis ≤20% cirrhosis ≤20% cirrhosis ≤20% cirrhosis	Mixed Mixed Mixed NR Mixed	92/134 245/277 54/77 485/524 85/103 189/375	2 Mixed Mixed Mixed Mixed	+		0.687 (0.601, 0.7 0.884 (0.841, 0.9 0.701 (0.586, 0.8 0.777 (0.743, 0.4 0.825 (0.738, 0.4 0.504 (0.452, 0.5 0.746 (0.635, 0.5
isvirlgrazoprevir ovveki 2015 zem 2015 ri 2016 vn 2018 2018 2019a total (i² = 98.0%, p = 0.000)	Multinational Japan Multinational Multinational Multinational Multinational	52 51 52 48 52 48	48 52 46 57 58 56	0% cirrhosis 0% cirrhosis ≤20% cirrhosis ≤20% cirrhosis 0% cirrhosis ≤20% cirrhosis	Yes Mixed Yes Mixed Yes Yes	24/43 219/227 175/246 57/129 15/19 230/486	1 Mixed Mixed Mixed Mixed			0.558 (0.399, 0. 0.965 (0.932, 0. 0.711 (0.650, 0. 0.519 (0.430, 0. 0.729 (0.544, 0. 0.473 (0.4226, 0. 0.791 (0.500, 0.
isbuvir/diaciatasvir owski 2014 zem 2018 E3 total (l² = 90.5%, p = 0.001)	U.S. Multinational	55 49	51 55	≤20% cirrhosis 0% cirrhosis	Yes Yes	38/41 80/115	1 3		~	0.927 (0.801, 0 0.696 (0.603, 0 0.827 (0.585, 0
sapreurinpibre ntasvir rolad 2017 zema 2018 Etem 2018 E1 obla 2018 E3 stah 2018 E3 stah 2018 stah 2019 otal (12 - 77.7%, p = 0.000)	U.S. Japan Multinational Japan Multinational Multinational Multinational	59 54 57 48 58	25 54 51 53 45 52 54	0% cirrhosis 0% cirrhosis 0% cirrhosis 0% cirrhosis 0% cirrhosis 0% cirrhosis ≤20% cirrhosis	No Mixed Mixed Mixed Yes Mixed Mixed	23/28 74/129 4507703 43/90 275/390 128/203 45/84	1 1 2 3 Mixed Mixed	 		0.821 (0.631, 0 0.574 (0.484, 0 0.640 (0.603, 0 0.478 (0.371, 0 0.705 (0.657, 0 0.631 (0.560, 0 0.548 (0.435, 0 0.623 (0.561, 0
oltasviriparita previri/riton avir/da reone 2014 enci 2014 Pill enci 2014 Pill adda, 2015 Iz 2015 e 2016 M1 total (I ² = 91.6%, p = 0.000)	sabuvir Multinational Multinational Japan Múltinational Multinational	54 51 55 47	40 59 37 53 51 52	0% cirrhosis 0% cirrhosis 0% cirrhosis 0% cirrhosis 0% cirrhosis 0% cirrhosis	No Yes Mixed Mixed Yes	74/95 140/209 169/205 148/215 76/82 ¢1/83		_•		0.779 (0.682. 0 0.670 (0.602. 0 0.824 (0.765. 0 0.688 (0.622. 0 0.927 (0.848. 0 0.494 (0.382. 0 0.751 (0.623. 0
oftasviripanita previriniton a viri/da enol 2014 Pili enol 2014 Pili zani 2015 e 2016 M1 e 2016 M1 e 2016 M2 e 2015 e 2015 e 2015 da 2015 cold (i ² = 87.2%, p = 0.000)	sabuvir + ribavirin Mutinational Mutinational Mutinational U.S. Mutinational Mutinational Mutinational Egyot Mutinational	54 48 48 48 48 49 49	50 49 30 33 49 45 32 9 30 43	0% cirrhosis 0% cirrhosis 0% cirrhosis 0% cirrhosis 0% cirrhosis ∞20% cirrhosis ∞20% cirrhosis ∞20% cirrhosis	No Yes Mixed Yes No Yes Mixed Yes Yes	72/91 168/210 92/100 35/38 115/153 53/101 53/87 50/101 50/100 414/473	1 1 1 1 1 4 Mixed	=		0.791 (0.593, 0. 0.800 (0.739, 0. 0.920 (0.348, 0. 0.920 (0.756, 0. 0.752 (0.575, 0. 0.629 (0.522, 0. 0.809 (0.749, 0. 0.800 (0.748, 0. 0.875 (0.842, 0. 0.875 (0.842, 0. 0.811 (0.742, 0.
erogeneity between groups: p rall (i ² = 94.712%, p = 0.000)	- 0.003								- 0 -	0.733 (0.680, 0
							() .25 .5	.75	l 1

Abbreviations: CI = confidence interval; Tx = treatment; U.S. = United States.

				A	cross differer	nt genotypes			
uthor year	Country	Age	% Female	Fibrosis stage	Tx naïve	Events/Total	Genotype		Proportion (95% CI)
edipasvirisofosbuvir awitz 2014b (oxodley 2014a ddahl 2014 im 2016 Jeine 2016 Jane 2016 bergel 2016b bergel 2016b	U.S. U.S. Multinational Korea China New Zealand France France	47 53 52 54 47 47 52 61	36 41 43 50 42 50 48	0% cirrhosis ≤20% cirrhosis ≤20% cirrhosis ≤20% cirrhosis ≤20% cirrhosis ≤20% cirrhosis ≤20% cirrhosis	Yes Yes Yes Mixed Mixed Yes Yes	1/39 9/431 1/214 3/93 3/206 5/50 0/44 1/41	1 1 1 3 or 6 4 5		0.026 (0.001, 0.135) 0.021 (0.010, 0.039) 0.005 (0.000, 0.028) 0.032 (0.007, 0.081) 0.032 (0.007, 0.081) 0.000 (0.030, 0.218) 0.000 (0.030, 0.218) 0.000 (0.000, 0.080) 0.024 (0.001, 0.129)
subtotal (I*= 47.0%, p = 0.067) Simeprevir/sofosbuvir awitz 2014a Swo 2018 Subtotal (I*= 0.0%, p = 0.808)	U.S. Canada & U.S.	56 56	42 47	0% cirrhosis 0% cirrhosis	No Mixed	0/14 1/155	1 1	0	0.020 (0.010, 0.039) 0.000 (0.000, 0.232) 0.006 (0.000, 0.035) 0.006 (0.001, 0.041)
Sofosbuvir/velpatasvir verson 2015 (Part A) oster 2015 A3 eld, 2015 srebely 2015 83 Vel 2015b Vel 2019b Jubtotal (I ² = 57.0%, p = 0.040)	U.S. U.S. U.S. Multinational Multinational Multinational	49 57 49 54 48 45	39 36 39 60 28 47	0% cirrhosis ≰20% cirrhosis ≼20% cirrhosis ≼20% cirrhosis ≰20% cirrhosis ≰20% cirrhosis	Yes Mixed Mixed NR Mixed	1/77 1/134 6/277 15/824 7/103 3/375	1 3 Mixed Mixed Mixed		$\begin{array}{c} 0.013 \left(0.000, 0.070 \right) \\ 0.007 \left(0.000, 0.047 \right) \\ 0.024 \left(0.014, 0.039 \right) \\ 0.024 \left(0.014, 0.039 \right) \\ 0.026 \left(0.028, 0.135 \right) \\ 0.008 \left(0.022, 0.023 \right) \\ 0.010 \left(0.001, 0.041 \right) \end{array}$
:Ibasvirigrazoprevir Sulkowski 2015 Umrada 2017 Jeuzem 2015 Isrown 2018 Vei 2019a Subtotal (I ² = 41.6%, p = 0.128)	Multinational Japan Multinational Multinational Multinational Multinational	52 81 52 48 52 48	48 82 46 57 58 56	0% cirrhosis 0% cirrhosis ≤20% cirrhosis 0% cirrhosis 0% cirrhosis ≤20% cirrhosis	Yes Mixed Yes Mixed Yes Yes	0/44 11/227 7/246 1/129 0/19 8/486	1 Mixed Mixed Mixed Mixed		0.000 (0.000.0.080) 0.048 (0.024.0.085) 0.028 (0.012.0.085) 0.008 (0.000.0.042) 0.000 (0.000.0.176) 0.016 (0.07.0.032) 0.021 (0.011.0.039)
Nofosbuvir/daclatasvir Sulkowski 2014 Jelson 2015 Jeuzem 2018 Subtotal (I ² = 0.0%, p = 0.521)	U.S. U.S. Multinational	55 53 49	51 43 55	≤20% cirrhosis ≤20% cirrhosis 0% cirrhosis	Yes Yes Yes	1/41 1/152 2/115	1 3 3		0.024 (0.001, 0.129) 0.007 (0.000, 0.036) 0.017 (0.002, 0.061) 0.013 (0.005, 0.034)
Biecaprevi/pibrentssvir Voordad 2017 Unayame 2018 uzzem 2018 E1 Voyoda 2018 duzzem 2018 E3 duzzem 2018 sealah 2018 sealah 2018 ubotal (I = 43.5%, p = 0.088)	U.S. Japan Multinational Japan Multinational Multinational Multinational Multinational	59 64 52 57 48 47 58 52	25 84 51 63 48 41 54 52	0% cirrhosis 0% cirrhosis 0% cirrhosis 0% cirrhosis 0% cirrhosis 20% cirrhosis ≰20% cirrhosis	No Mixed Mixed Yes Yes Mixed Mixed	1/28 0/129 9/703 2/90 5/233 3/157 5/84 2/203	1 1 2 3 Mixed Mixed		0.036(0.001.0.183) 0.001(0.000.0.028) 0.013(0.006.0.024) 0.022(0.003.0.078) 0.022(0.003.0.078) 0.021(0.007.0.049) 0.019(0.004.0.055) 0.060(0.020.0.133) 0.010(0.001.0.035) 0.010(0.011.0.035)
Imbitasvir/paritaprevir/ritonavir/dasabuvir erend 2014 Pill Lovadley 2014b erend 2014 PIV awitz 2015 Limada, 2015 Limada, 2015 Utottal (1 ² = 31.2%, p = 0.190)	Multinational Multinational Multinational Multinational Japan Multinational	49 54 48 51 55 81 47	59 40 43 37 51 63 52	0% cirrhosis 0% cirrhosis 0% cirrhosis 0% cirrhosis 0% cirrhosis 20% cirrhosis \$20% cirrhosis	Yes No Yes Yes Mixed Mixed Yes	4/209 3/95 2/79 1/205 2/82 7/215 0/83	1 1 1 1 1 1	- +	0.019(0.005,0.048) 0.032(0.07,0.090) 0.025(0.03,0.068) 0.054(0.001,0.027) 0.054(0.001,0.027) 0.034(0.010,0.027) 0.030(0.010,0.049) 0.000(0.000,0.043) 0.019(0.012,0.032)
Mbliskriparitaprevir/ritonavir/dasabuvir- erenci 2014 P(II eld, 2014 P) errenci 2014 P) errenci 2014 P more 2016 M1 preschi 2016 Mitter 2016 Wated 2016 Wated 2016 Wated 2016	+ribavirin Multinational Multinational Multinational Multinational U.S. Multinational Multinational Multinational Multinational Multinational Egypt	48 49 50 52 54 48 47 46 48 48 48 49	49 43 44 30 30 34 39 23 29 30	0% cirrhosis 0% cirrhosis 0% cirrhosis 0% cirrhosis 20% cirrhosis 20% cirrhosis 20% cirrhosis 20% cirrhosis 20% cirrhosis	Yes Yes Yes No Mixed No Yes Yes Mixed Yes	4/210 10/473 3/100 2/91 3/38 1/101 1/153 5/87 0/91 2/100	1 1 1 1 1 1 1 4 4		0.019(0.005.0.048) 0.021(0.010.0.038) 0.031(0.000,0.049) 0.030(0.006,0.049) 0.022(0.003,0.077) 0.079(0.017,0.214) 0.010(0.000,0.044) 0.007(0.000,0.044) 0.007(0.000,0.044) 0.007(0.000,0.044) 0.007(0.000,0.044) 0.007(0.000,0.044) 0.007(0.000,0.044) 0.007(0.000,0.044) 0.007(0.000,0.044) 0.007(0.000,0.044) 0.007(0.000,0.044) 0.007(0.000,0.044) 0.007(0.000,0.044) 0.000(0.014,0.0146) 0.000(0.000,0.044) 0.000(0.014,0.0146) 0.000(0.000,0.044) 0.00
leterogeneity between groups: p = 0.939 Sverall (l ² = 31.434%, p = 0.019)								• •	0.019(0.015, 0.024)
									1 .3 .4



Across different genotypes												
uthor year	Country	Age	% Female	Fibrosis stage	Tx naïve	Events/Total	Genotype		Proportion (95% CI)			
edipasvir/sofosbuvir tishi 2014 switz 2014b owoley 2014a m 2016 00:16 ei 2018 sne 2015 bergel 2016a bergel 2016a bergel 2016a	Multinational U.S. U.S. Korea Taiwan China China Mew Zealand France France	52 47 53 54 54 47 47 52 61	41 38 41 43 69 50 42 50 42	£20% cirrhosis 0% cirrhosis 20% cirrhosis £20% cirrhosis £20% cirrhosis £20% cirrhosis £20% cirrhosis £20% cirrhosis £20% cirrhosis	Yes Yes Yes Mixed Mixed Yes Yes	0/214 0/39 2/431 1/93 1/85 0/206 1/50 0/44 0/41	1 1 1 3 or 6 4 5		$\begin{array}{c} 0.000 \left(0.000, 0.017 \right) \\ 0.000 \left(0.000, 0.090 \right) \\ 0.005 \left(0.001, 0.017 \right) \\ 0.015 \left(0.011, 0.017 \right) \\ 0.011 \left(0.000, 0.056 \right) \\ 0.000 \left(0.000, 0.018 \right) \\ 0.020 \left(0.000, 0.018 \right) \\ 0.000 \left(0.000, 0.080 \right) \\ 0.000 \left(0.000, 0.080 \right) \\ 0.004 \left(0.002, 0.010 \right) \end{array}$			
meprevir/sofosbuvir witz 2014a vo 2018 ubtotal (I ² = 0.0%, p= 0.510)	U.S. Canada & U.S.	56 56	42 47	0% cirrhosis 0% cirrhosis	No Mixed	0/14 0/155	1 1	¢	0.000 (0.000, 0.232) 0.000 (0.000, 0.024) 0.000 (0.000, 0.218)			
ofosbuvir/velpatasvir verson 2015 (Part A) oster 2015 A2 oster 2015 A3 ela 2015 rebely 2018 ei 2019 bubotal (I ² = 0.0%, p = 0.432)	U.S. U.S. Multinational Multinational Multinational	49 57 49 54 48 45	39 36 39 60 28 47	0% cirrhosis ≤20% cirrhosis ≤20% cirrhosis ≤20% cirrhosis ≤20% cirrhosis ≤20% cirrhosis	Yes Mixed Mixed NR Mixed	0/77 1/134 0/277 1/624 1/103 0/375	1 2 3 Mixed Mixed Mixed	F	$\begin{array}{c} 0.000 \left(0.000, 0.047 \right) \\ 0.07 \left(0.000, 0.047 \right) \\ 0.000 \left(0.000, 0.043 \right) \\ 0.000 \left(0.000, 0.013 \right) \\ 0.02 \left(0.000, 0.053 \right) \\ 0.010 \left(0.000, 0.053 \right) \\ 0.000 \left(0.000, 0.010 \right) \\ 0.002 \left(0.001, 0.000 \right) \end{array}$			
basvir/grazoprevir ulkowski 2015 suzem 2015" suzem 2015" perl 2016 ti 2019 si 2019 si 2019 subtotal (l ² = 0.0%, p = 0.830)	Multinational Japan Multinational Multinational Multinational Multinational	52 81 52 48 52 48	48 62 46 57 58 58	0% cirrhosis 50% cirrhosis 420% cirrhosis 0% cirrhosis 0% cirrhosis 420% cirrhosis	Yes Mixed Yes Mixed Yes Yes	0/44 3/227 2/246 1/129 1/19 3/486	1 Mixed Mixed Mixed Mixed		$\begin{array}{c} 0.000 \left(0.000, 0.080 \right) \\ 0.013 \left(0.003, 0.038 \right) \\ 0.008 \left(0.001, 0.029 \right) \\ 0.008 \left(0.001, 0.029 \right) \\ 0.005 \left(0.001, 0.042 \right) \\ 0.005 \left(0.001, 0.260 \right) \\ 0.006 \left(0.001, 0.260 \right) \\ 0.009 \left(0.005, 0.018 \right) \end{array}$			
ofosbuvir/daclatasvir ulkowski 2014 euzem 2018 E3 ubtotal (I ² = 0.0%, p = 0.700)	U.S. Multinational	55 49	51 55	≤20% cirrhosis 0% cirrhosis	Yes Yes	0/41 1/115	1 3	<u> </u>	0.000 (0.000, 0.088) 0.009 (0.000, 0.047) 0.006 (0.001, 0.044)			
lecaprevi/pibrentasvir oordad 2017 hayama 2018 auzem 2018 E1 oyoda 2018 auzem 2018 E3 aselah 2018 biototal (I ^e = 0.0%, p = 0.565)	U.S. Japan Multinational Japan Multinational Multinational Multinational	59 64 52 57 48 52 58	25 84 51 53 45 52 54	0% cirrhosis 0% cirrhosis 0% cirrhosis 0% cirrhosis 0% cirrhosis 0% cirrhosis ≤20% cirrhosis	No Mixed Mixed Yes Mixed Mixed	0/28 0/129 1/703 1/90 3/390 0/203 0/84	1 1 2 3 Mixed Mixed		$\begin{array}{c} 0.000 \; (0.000, 0.123) \\ 0.000 \; (0.000, 0.023) \\ 0.001 \; (0.000, 0.026) \\ 0.001 \; (0.000, 0.000) \\ 0.008 \; (0.002, 0.022) \\ 0.000 \; (0.000, 0.018) \\ 0.000 \; (0.000, 0.018) \\ 0.003 \; (0.001, 0.043) \\ 0.003 \; (0.001, 0.043) \\ \end{array}$			
$eq:started_st$	Multinational Multinational Multinational Multinational Japan Multinational Multinational	54 49 51 48 61 55 47	40 59 37 43 63 51 52	0% cirrhosis 0% cirrhosis 0% cirrhosis 0% cirrhosis 0% cirrhosis 0% cirrhosis 20% cirrhosis ≤20% cirrhosis	No Yes Yes Mixed Mixed Yes	0/95 0/209 0/205 0/79 2/215 0/82 0/83	1 1 1 1 1 1 1		$\begin{array}{c} 0.000 (0.000, 0.038) \\ 0.000 (0.000, 0.018) \\ 0.000 (0.000, 0.016) \\ 0.000 (0.000, 0.046) \\ 0.000 (0.000, 0.046) \\ 0.000 (0.000, 0.044) \\ 0.000 (0.000, 0.044) \\ 0.000 (0.000, 0.044) \\ 0.000 (0.000, 0.044) \\ 0.001 (0.000, 0.044) \\ 0.001 (0.000, 0.044) \end{array}$			
mbitasvir(paritaprevir/ittonavir/dasabuvir- send 2014 PUI bid 2014 send 2014 PUI ordeone 2014 over 2014 bid over 2016 bid over 2016 bid rebely 2018 bid sold 2016 bidtal (1 = 11.2%, p = 0.337)	+ ribavirin Multinational Multinational Multinational U.S. Multinational Multinational Multinational Multinational Multinational Egypt	52 9 48 54 54 46 7 48 46 48 49	30 43 50 44 39 46 23 29 30	0% cirrhosis 0% cirrhosis 0% cirrhosis 0% cirrhosis 0% cirrhosis 200% cirrhosis 220% cirrhosis 0% cirrhosis 220% cirrhosis	Yes Yes No Yes Mixed Yes No Yes Mixed Yes	0/100 0/210 2/91 2/79 1/38 1/153 0/81 0/81 0/91 0/100	1 1 1 1 1 1 1 1 1 4		$\begin{array}{c} 0.000 \left(0.000, 0.036 \right) \\ 0.006 \left(0.001, 0.018 \right) \\ 0.006 \left(0.000, 0.017 \right) \\ 0.022 \left(0.003, 0.077 \right) \\ 0.022 \left(0.003, 0.078 \right) \\ 0.021 \left(0.003, 0.086 \right) \\ 0.007 \left(0.000, 0.036 \right) \\ 0.000 \left(0.000, 0.036 \right) \\ 0.000 \left(0.000, 0.046 \right) \\ 0.000 \left(0.000$			
eterogeneity between groups: $p = 0.194$ verall ($l^2 = 0.000\%$, $p = 0.653$)									0.004 (0.003, 0.008)			

Abbreviations: CI = confidence interval; NR = not reported; Tx = treatment; U.S. = United States.

						Anemia			
					Across di	ifferent genotype	s		
Author year	Country	Age	% Female	Fibrosis tage	Tx naïve	Events/Total	Genotype		Proportion (95% CI)
Ledipasvir/sofosb	uvir								
Afdahl 2014	Multinational	52	41	≤20% cirrhosis	Yes	0/214	1	•	0.000 (0.000, 0.017)
Kowdley 2014a	U.S.	53	41	0% cirrhosis	Yes	3/431	1	•	0.007 (0.001, 0.020)
Subtotal (I ² = 43.	9%, p = 0.182)								0.005 (0.002, 0.014)
Simeprevir/sofost	ouvir								
Lawitz 2014a	U.S.	56	42	0% cirrhosis	No	0/14	1	• •	0.000 (0.000, 0.232)
Glecaprevir/pibre	ntasvir						_	11	
Toyoda 2018	Japan	57	53	0% cirrhosis	Mixed	0/90	2	₽†:	0.000 (0.000, 0.040)
Ombitasvir/paritap	previr/ritonavir/da	sabuvir							
Andreone 2014	Multinational	54	40	0% cirrhosis	No	0/95	1	₽ -+	0.000 (0.000, 0.038)
Kowdley 2014b	Multinational	48	43	0% cirrhosis	Yes	1/79	1	= :	0.013 (0.000, 0.069)
Dore 2016 M2	Multinational	47	52	≤20% cirrhosis	Yes	1/83	1	<u>.</u>	0.012 (0.000, 0.065)
Subtotal (I ² = 0.0	%, p = 0.439)							2	0.008 (0.002, 0.031)
Ombitasvir/paritap	orevir/ritonavir/da	sabuvir	+ ribavirin						
Andreone 2014	Multinational	54	50	0% cirrhosis	No	10/91	1	:	0.110 (0.054, 0.193)
Kowdley 2014b	Multinational	50	44	0% cirrhosis	Yes	7/79	1	l:	0.089 (0.036, 0.174)
Lalezari 2015	U.S.	48	34	0% cirrhosis	Mixed	4/38	1	;	0.105 (0.029, 0.248)
Dore 2016 M1	Multinational	46	39	≤20% cirrhosis	Yes	10/153	1	!	0.065 (0.032, 0.117)
Dore 2016 M2	Multinational	47	46	≤20% cirrhosis	No	3/101	1		0.030 (0.006, 0.084)
Grebely 2018a	Multinational	48	23	≤20% cirrhosis	Yes	12/87	1	! <u></u> ■	0.138 (0.073, 0.229)
Subtotal (I ² = 48.	6%, p = 0.083)								0.083 (0.058, 0.118)
Heterogeneitv be	tween groups: p	= 0.000							
Overall (1 ² = 85.1	94%, p = 0.000)							\diamond	0.024 (0.009, 0.063)
								!	
									4
						Proportion		5 .1 .Z .0	

Abbreviations: CI = confidence interval; Tx = treatment; U.S. = United States.
				A	Across diffe	erent genoty	oes		
Author year	Country	Age	% Female	Fibrosis stage	Tx naïve	Events/Total	Genotype		Proportion (95% CI)
edipasvirisofoobuvir tidani 2014 ixdani 2014 im 2016 ihuang, 2016 ibergel 2016 ibergel 2016 ibergel 2018 ubergel 2018	Multinational U.S. Korea Taiwan New Zealand France Egypt France	52 53 54 47 52 51 81	41 41 69 42 50 35 48	≤20% cirrhosis ≤20% cirrhosis ≤20% cirrhosis ≤20% cirrhosis ≤20% cirrhosis Undear/NR ≤20% cirrhosis	Yes Yes Yes Mixed Yes Yes Yes	44/214 94/431 7/93 8/85 11/50 9/44 18/100 4/41	1 1 3 or 6 4 5		0 206 (0 154, 0 266) 0 218 (0 180, 0 260) 0 075 (0 031, 0 149) 0 094 (0 042, 0 177) 0 220 (0 115, 0 380) 0 206 (0 198, 0 383) 0 180 (0 110, 0 289) 0 098 (0 027, 0 231) 0 162 (0 122, 0 210)
Simeprevir/sofosbuvir (wo 2016 ?ott-Junior 2019 Subtotal (I ² = 88.2%, p = 0.007)	Canada & U.S. Brazil	56 53	47 48	0% cirrhosis 0% cirrhosis	Mixed Mixed	19/155 17/80	1 1		0.123 (0.075, 0.185) 0.283 (0.175, 0.414) 0.184 (0.098, 0.318)
iofosbuvir/velpatasvir iverson 2015 (Part A) ioster 2015 A2 ioster 2015 A3 eld 2015 isebely 2018a iubtotal (I ² = 44.5%, p = 0.125)	U.S. U.S. U.S. Multinational Multinational	49 57 49 54 48	39 38 39 60 28	0% cirrhosis ≤20% cirrhosis ≤20% cirrhosis ≤20% cirrhosis ≤20% cirrhosis	Yes Mixed Mixed NR	14/77 20/134 71/277 128/624 23/103	1 2 3 Mixed Mixed		0.182 (0.103.0.286) 0.149 (0.094, 0.221) 0.256 (0.206, 0.312) 0.205 (0.171, 0.236) 0.223 (0.147, 0.316) 0.228 (0.179, 0.240)
Elbasvir/grazoprevir Sulkowski 2015 Brown 2018 Vei 2019a Subtotal (I ² = 87.9%, p = 0.000)	Multinational Multinational Multinational	52 52 48	48 58 56	0% cirrhosis 0% cirrhosis ≤20% cirrhosis	Yes Yes Yes	10/43 3/19 22/486	1 Mixed Mixed		0.233 (0.118, 0.386) 0.158 (0.034, 0.396) 0.045 (0.029, 0.088) 0.109 (0.043, 0.251)
Sofosbuvir/daclatasvir Sulkowski 2014 ott.Junior 2019 Jelson 2015 Jeuzem 2018 E3 Subtotal (I ² = 72.3%, p = 0.013)	U.S. Brazil U.S. Multinational	55 58 53 49	51 53 43 55	≤20% cirrhosis 0% cirrhosis ≤20% cirrhosis 0% cirrhosis	Yes Mixed Yes Yes	18/41 15/65 29/152 16/115	1 1 3 3		0.390 (0.242, 0.555) 0.231 (0.135, 0.352) 0.191 (0.132, 0.282) 0.139 (0.822, 0.216) 0.217 (0.149, 0.307)
Slecaprevir/pibrentasvir 'oordad 2017 leuzem 2018 E1 leuzem 2018 E3 sselah 2018 sselah 2018a Subtotal (I ² = 54.1%, p = 0.069)	U.S. Multinational Multinational Multinational Multinational	59 52 48 58 52	25 51 45 54 52	0% cirrhosis 0% cirrhosis 0% cirrhosis ≤20% cirrhosis 0% cirrhosis	No Mixed Yes Mixed Mixed	5/28 74/703 64/390 11/84 28/203	1 1 3 Mixed Mixed		0.179 (0.061.0.369) 0.105 (0.084.0.130) 0.164 (0.129.0.205) 0.131 (0.067.0.222) 0.138 (0.064.0.193) 0.138 (0.084.0.193)
Ombitasvir/paritaprevir/ritonavir/dasabu erenci 2014 PIII erenci 2014 PIV undreone 2014 Gwadley 2014b a.witz 2015 Jore 2016 M2 Subtotal (I ² = 90.9%, p = 0.000)	uvir Multinational Multinational Multinational Multinational Multinational Multinational	49 51 54 48 55 47	59 37 40 43 51 52	0% cirrhosis 0% cirrhosis 0% cirrhosis 0% cirrhosis 0% cirrhosis ≤20% cirrhosis	Yes Yes No Yes Mixed Yes	48/209 72/205 15/95 16/79 6/82 4/83	1 1 1 1 1		0.230 (0.174, 0.293) 0.361 (0.286, 0.421) 0.168 (0.091, 0.247) 0.230 (0.102, 0.308) 0.073 (0.027, 0.162) 0.048 (0.013, 0.119) 0.158 (0.091, 0.261)
Imbilisari/iparitapreviri/itonaviri/dasabu ondraone 2014 ieranci 2014 PIII iowaley 2014 ieranci 2014 PIV alezari 2016 owe 2016 M2 ieranci 2015 iezode 2015 vatotale 2016 vatotale 2016 vatotale 16 = 87.7%, p = 0.000)	vir + ribavirin Multinational Multinational Multinational Multinational U.S. Multinational Multinational Multinational Multinational Multinational Egypt	54 48 50 49 52 48 47 48 48 48 49	50 49 44 30 34 46 39 23 29 30	0% cirrhosis 0% cirrhosis 0% cirrhosis 0% cirrhosis 0% cirrhosis 20% cirrhosis 20% cirrhosis 20% cirrhosis 20% cirrhosis 20% cirrhosis	No Yes Yes Yes Mixed No Yes Mixed Yes	29/91 45/210 22/79 184/473 48/100 18/38 12/101 21/153 25/87 14/91 35/100	1 1 1 1 1 1 1 4 4		0 319 (0 225, 0 425) 0 214 (0 161, 0 276) 0 278 (0 183, 0.381) 0 347 (0 334, 0 332) 0 460 (0 360, 0 563) 0 119 (0 060 0 1932) 0 119 (0 060 0 1932) 0 227 (0 195, 0 334) 0 154 (0 087, 0 245) 0 360 (0 257, 0 452) 0 326 (0 225, 0 344)
leterogeneity between groups: p = 0.00 Overall (l ² = 89.508%, p = 0.000)	05								0.184 (0.156, 0.217)
								0 .2 .4 .	6 .8



				٨	Hea cross diffo	dache	05		
uthor Year	Country	Age	% Female	Fibrosis stage	TX naïve	Events/Total	Genotype	1	Proportion (95% CI)
dipasvir/sofosbuvir switz 2014b dahl 2014 wadiey 2014a m 2016 ang 2016 ang 2016 bergel 2016 bergel 2016 bergel 2016 bottari (1- 85.3%, p = 0.000)	U.S. Multinational U.S. Korea Taiwan New Zealand France Egypt France	47 52 53 54 47 52 52 51 81	36 41 43 69 42 50 35 48	0% cirrhosis \$20% cirrhosis \$20% cirrhosis \$20% cirrhosis \$20% cirrhosis \$20% cirrhosis Unclear/NR \$20% cirrhosis	Yes Yes Yes Yes Mixed Yes Yes Yes	2/39 53/214 63/431 7/93 12/85 12/50 11/44 2/100 11/41	1 1 1 3 or 6 4 5		$\begin{array}{c} 0.051 \left(0.006, 0.173 \right) \\ 0.248 \left(0.191, 0.311 \right) \\ 0.146 \left(0.114, 0.183 \right) \\ 0.075 \left(0.031, 0.149 \right) \\ 0.240 \left(0.131, 0.149 \right) \\ 0.240 \left(0.131, 0.382 \right) \\ 0.250 \left(0.132, 0.403 \right) \\ 0.202 \left(0.002, 0.070 \right) \\ 0.288 \left(0.142, 0.429 \right) \\ 0.137 \left(0.084, 0.215 \right) \end{array}$
meprevir/sofosbuvir vo 2018 ott-Junior 2019 ubtotal (I ² = 81.4%, p = 0.020)	Canada & U.S. Brazil	56 53	47 48	0% cirrhosis 0% cirrhosis	Mixed Mixed	22/155 17/80	1 1		0.142(0.091,0.207) 0.283(0.175,0.414) 0.195(0.117,0.308)
ofosbuvir/ve[patasvir verson 2015 (Part A) sster 2015 A2 eld, 2015 rebely 2018a ei 2019b ubtotal (l ² = 96.4%, p = 0.000)	U.S. U.S. U.S. Multinational Multinational Multinational	49 57 49 54 48 45	39 36 39 60 28 47	0% cirrhosis ≤20% cirrhosis ≤20% cirrhosis ≤20% cirrhosis ≤20% cirrhosis ≤20% cirrhosis	Yes Mixed Mixed NR Mixed	14/77 24/134 90/277 182/824 19/103 18/375	1 2 3 Mixed Mixed Mixed		0.182(0.103,0.286) 0.179(0.118,0.285) 0.325(0.270,0.384) 0.292(0.286,0.329) 0.184(0.116,0.273) 0.048(0.029,0.075) 0.180(0.108,0.285)
basvir/grazoprevir Jlkowski 2015 rown 2018 ei 2019a Jubtotal (l²= 93.5.%, p = 0.000)	Multinational Multinational Multinational	52 52 48	48 58 58	0% cirrhosis 0% cirrhosis ≤20% cirrhosis	Yes Yes Yes	15/43 5/19 27/486	1 Mixed Mixed		0.349 (0.210, 0.509) 0.263 (0.091, 0.512) 0.056 (0.037, 0.080) 0.171 (0.061, 0.395)
ofosbuvir/daclatasvir ulkovski 2014 tit-Junior 2019 elson 2015 euzem 2018 E3 ubtotal (I ² = 41.2%, p = 0.184)	U.S. Brazil U.S. Multinational	55 56 53 49	51 53 43 55	≤20% cirrhosis 0% cirrhosis ≤20% cirrhosis 0% cirrhosis	Yes Mixed Yes Yes	14/41 10/85 30/152 23/115	1 1 3 3		0.341 (0.201, 0.506) 0.154 (0.076, 0.265) 0.197 (0.137, 0.270) 0.200 (0.131, 0.285) 0.206 (0.168, 0.251)
lecaprevir/pibrentasvir oordad 2017 sayama 2018 tuzem 2018 E1 syoda 2018 suzem 2018 E3 selah 2018 selah 2018 bottoti (f= #8.7%, p = 0.000)	U.S. Japan Multinational Japan Multinational Multinational Multinational	59 64 52 57 48 52 57 48 58	25 84 51 53 45 52 54	0% cirrhosis 0% cirrhosis 0% cirrhosis 0% cirrhosis 0% cirrhosis 0% cirrhosis ≤20% cirrhosis	No Mixed Mixed Yes Mixed Mixed	9/28 6/129 130/703 6/90 91/390 37/203 11/84	1 1 2 3 Mixed Mixed		0 321 (0 159 0 524) 0 047 (0 017 0 068) 0 168 (0 157 0 276) 0 057 (0 025 0 139) 0 230 (192 0 279) 0 152 (0 132 0 242) 0 131 (0 057 0 222) 0 147 (0 094 0 222)
mbitasvir/paritaprevir/ritonavir/dasab drieone 2014 erenci 2014 PIV erenci 2014 PIV erenci 2014 PIII worley 2014 PI writz 2016 mada 2016 total (I= 83.2%, p = 0.000)	buvir Multinational Multinational Multinational Multinational Japan Multinational	54 51 49 48 55 61 47	40 37 59 43 51 63 52	0% cirrhosis 0% cirrhosis 0% cirrhosis 0% cirrhosis 0% cirrhosis 0% cirrhosis ≤20% cirrhosis	No Yes Yes Mixed Mixed Yes	22/95 58/205 49/209 15/79 24/82 19/215 16/83	1 1 1 1 1 1	-+++ +++-+0 +-+0	$\begin{array}{c} 0.232(0.151,0.329)\\ 0.283(0.222,0.380)\\ 0.234(0.179,0.298)\\ 0.159(0.110,0.294)\\ 0.239(0.102,0.294)\\ 0.088(0.004,0.135)\\ 0.139(0.141,0.294)\\ 0.230(0.146,0.266)\\ 0.207(0.166,0.266)\\ \end{array}$
mbilasvi/parilaprevir/itonavir/dasab wolley 2014 Hd 2014 Erend 2014 PIII drecene 2014 Serend 2014 PIV Drecene 2016 PIV Drecene 2016 M1 recele 2018 Scode 2015 Scode 2015 Scode 2015 Scode 2015	suvir + ribavirin Mutiinational Mutiinational Mutinational Mutinational Mutinational Mutinational Mutinational Mutinational Egypt	50 48 54 48 48 47 48 48 48 48 48 49	44 43 49 30 34 46 39 23 29 30	0% cirrhosis 0% cirrhosis 0% cirrhosis 0% cirrhosis 0% cirrhosis 420% cirrhosis 420% cirrhosis 420% cirrhosis 420% cirrhosis 420% cirrhosis	Yes Yes No Yes Nixed No Yes Yes Mixed Yes	21/79 153/473 51/210 22/91 25/100 12/38 29/101 41/153 12/87 28/91 41/100	1		$\begin{array}{c} 0.266 (0.173, 0.377)\\ 0.323 (0.281, 0.368)\\ 0.243 (0.186, 0.307)\\ 0.243 (0.186, 0.307)\\ 0.245 (0.169, 0.347)\\ 0.255 (0.169, 0.347)\\ 0.256 (0.169, 0.347)\\ 0.286 (0.200, 0.346)\\ 0.288 (0.200, 0.346)\\ 0.288 (0.200, 0.346)\\ 0.308 (0.274, 0.229)\\ 0.316 (0.312, 0.216)\\ 0.2176 (0.240, 0.316)\\ 0.216 (0.240, 0.316)\\ 0.216 (0.240, 0.316)\\ 0.216 (0.240, 0.316)\\ 0.216 (0.240, 0.316)\\ 0.216 (0.240, 0.316)\\ 0.216 (0.240, 0.316)\\ 0.216 (0.240, 0.316)\\ 0.216 (0.240, 0.316)\\ 0.216 (0.240, 0.316)\\ 0.216 (0.240, 0.316)\\ 0.216 (0.240, 0.316)\\ 0.216 (0.240, 0.316)\\ 0.216 (0.240, 0.316)\\ 0.216 (0.240, 0.216)\\ 0.216 (0.240, $
eterogeneity between groups: p = 0.0 verall (l ² = 90.362%, p = 0.000)	004							↓	0.187 (0.156, 0.222)
					Pro	portion		0 .2 .4	.6

Abbreviations: CI = confidence interval; NR = not reported; Tx = Treatment; U.S. = United States.

					Insor	mnia			
				Acr	oss differe	nt genotypes		1	
Author year	Country	Age	% Female	Fibrosis stage	Tx naïve	Events/Total	Genotype		Proportion (95% CI)
Ledipasvir/sofosbuvir Afdahl 2014 Kowdley 2014a Ahmed 2018 Subtotal (l² = 58.2%, p = 0	Multinational U.S. Egypt .091)	52 53 51	41 41 35	≤20% cirrhosis 0% cirrhosis Unclear/NR	Yes Yes Yes	17/214 26/431 2/100	1 1 4	-++ -++	0.079 (0.047, 0.124) 0.060 (0.040, 0.087) 0.020 (0.002, 0.070) 0.060 (0.045, 0.080)
Simeprevir/sofosbuvir Pott-Junior 2019	Brazil	53	48	0% cirrhosis	Mixed	6/60	1	<u>-</u>	0.100 (0.038, 0.205)
Sofosbuvir/velpatasvir Everson 2015 (Part A) Foster 2015 A2 Foster 2015 A3 Feld 2015 Grebely 2018a Subtotal (I ² = 32.4%, p = 0	U.S. U.S. U.S. Multinational Multinational .206)	49 57 49 54 48	39 36 39 60 28	0% cirrhosis ≤20% cirrhosis ≤20% cirrhosis ≤20% cirrhosis ≤20% cirrhosis	Yes Mixed Mixed Mixed NR	5/77 6/134 31/277 50/624 9/103	1 2 3 Mixed Mixed		$\begin{array}{c} 0.065 & (0.021, \ 0.145) \\ 0.045 & (0.017, \ 0.095) \\ 0.112 & (0.077, \ 0.155) \\ 0.080 & (0.060, \ 0.104) \\ 0.087 & (0.041, \ 0.159) \\ 0.083 & (0.067, \ 0.102) \end{array}$
Elbasvir/grazoprevir Sulkowski 2015	U.S.	55	51	≤20% cirrhosis	Yes	3/43	1		0.070 (0.023, 0.195)
Sofosbuvir/daclatasvir Sulkowski 2014 Nelson 2015 Pott-Junior 2019 Poordad 2017 Subtotal (I ² = 0.0%, p = 0.8	U.S. Multinational Brazil U.S. 300)	59 55 56 53	18 51 53 43	0% cirrhosis 0% cirrhosis 0% cirrhosis ≤20% cirrhosis	No Mixed Mixed Yes	4/41 9/152 4/65 0/6	1 1 1 3 1		0.098 (0.027, 0.231) 0.059 (0.027, 0.109) 0.062 (0.017, 0.150) 0.000 (0.000, 0.459) 0.064 (0.040, 0.101)
Glecaprevir/pibrentasvir Poordad 2017	U.S.	59	50	0% cirrhosis	No	0/22	1		0.000 (0.000, 0.154)
Ombitasvir/paritaprevir/ritor Feld 2014 Lalezari 2015 Waked 2016 Grebely 2018b Hezode 2015 Subtotal (I ² = 0.0%, p = 0.6 Heterogeneity, between gro	navir/dasabuvir + rib. Multinational Canada & U.S. Multinational Multinational Multinational 604) ups: p = 0.000	avirin 49 56 52 48 48	43 47 48 23 29	0% cirrhosis 0% cirrhosis 0% cirrhosis ≤20% cirrhosis 0% cirrhosis	Yes Mixed Yes Yes Mixed	66/473 7/38 9/100 11/87 12/91	1 1 1 4		0.140 (0.110, 0.174) 0.184 (0.077, 0.343) 0.090 (0.042, 0.164) 0.126 (0.065, 0.215) 0.132 (0.070, 0.219) 0.133 (0.111, 0.159)
Overall (l ² = 58.287%, p =	0.001)							¢	0.083 (0.068, 0.101)
					_			0.2.4	.6

Abbreviations: CI = confidence interval; Tx = treatment; U.S. = United States.

					Across diff	erent genotv	pes		
uthor year	Country	Age	% Female	Fibrosis stage	Tx naïve	Events/Total	Genotype	1	Proportion (95% CI)
edipasvir/sofosbuvir	M. 16	50			Mar	04/044			0.440 (0.070.0.480)
Kowdley 2014a	U.S.	53	41	0% cirrhosis	Yes	39/431	1		0.090 (0.065, 0.122)
awitz 2014b	U.S.	47	36	0% cirrhosis	Yes	3/39	1		0.077 (0.016, 0.209)
Sane 2015	New Zealand	54 47	42	≤20% cirrhosis ≤20% cirrhosis	Yes Mixed	5/85 9/50	3 or 6		0.180 (0.086, 0.314)
Abergel 2016a	France	52	50	≤20% cirrhosis	Yes	4/44	4	I_ 	0.091 (0.025, 0.217)
Ahmed 2018 Subtotal (1 ² = 60.0%, p = 0.020)	Egypt	51	35	Undear/NR	Yes	2/100	4		0.020 (0.002, 0.070) 0.084 (0.057, 0.121)
50515121 (1 = 55.5 %, p = 5.525)								<u> </u>	0.004(0.007,0.121)
Simeprevir/sofosbuvir (wo. 2018	Canada & U.S.	58	47	0% cirrhosis	Mixed	23/155	1	<u> </u>	0 148 (0 096 0 214)
Pott-Junior 2019	Brazil	53	48	0% cirrhosis	Mixed	8/60	1		0.133 (0.059, 0.246)
Subtotal (I ² = 0.0%, p = 0.831)								\sim	0.144 (0.103, 0.198)
Bofosbuvir/velpatasvir								1 2	
Everson 2015 Foster 2015 A2	U.S. U.S	49	39	0% cirrhosis ≾20% cirrhosis	Yes	8/77 14/134	1		0.104 (0.046, 0.194)
Foster 2015 A3	U.S.	49	39	≤20% cirrhosis	Mixed	46/277	3		0.166 (0.124, 0.215)
Feld 2015	Multinational	54	60	≤20% cirrhosis	Mixed	75/624	Mixed		0.120 (0.096.0.148)
Subtotal (I ² = 13.1%, p = 0.330)	Mutunational	40	20	A20% Calmosis	INITS.	14/103	INIXED		0.129 (0.110, 0.15)
- Ibasvir/grazoprevir								1	
Bulkowski 2015	Multinational	52	48	0% cirrhosis	Yes	7/43	1		0.163 (0.068, 0.307)
Brown 2018	Multinational	52	58	0% cirrhosis	Yes	1/19	Mixed		0.053 (0.001. 0.260)
Subtotal (I* = 19.4%, p = 0.200)									0.129 (0.066, 0.237)
Sofosbuvir/daclatasvir			E1	COOK simbosis	Vee	0/44			0.105 (0.088, 0.249)
Pott-Junior 2019	Brazil	56	53	0% cirrhosis	Mixed	4/85	1		0.062(0.017, 0.150)
Velson 2015	U.S.	53	43	≤20% cirrhosis	Yes	18/152	3		0.118 (0.072, 0.181)
Subtotal (I ² = 31.9%, p = 0.221)	Morunational	45	00	0% cimosis	1 45	10/110	3	8	0.121 (0.091, 0.158)
3lecaprevir/pibrentasvir								1	
Poordad 2017	U.S.	59	25	0% cirrhosis	No	5/28	1		0.179 (0.061, 0.369)
Zeuzem 2018 E1	Multinational	52	51	0% cirrhosis	Mixed	48/703	1	1.7	0.068 (0.051, 0.090)
Toyoda 2018 Zeuzem 2018 E3	Japan Multinational	48	45	0% cirrhosis	Yes	51/390	3		0.033 (0.007, 0.094) 0.131 (0.099, 0.168)
Asselah 2018a	Multinational	52	52	0% cirrhosis	Mixed	23/203	Mixed		0.113 (0.073. 0.165)
Subtotal (I= / 9.2%, p = 0.001)									0.053 (0.004, 0.134)
Ombitasvir/paritaprevir/ritonavir/	dasabuvir	E4	40	0% sistesis	Ne	e/0E			0.083 (0.024.0.133)
Ferenci 2014 PIV	Multinational	51	37	0% cirrhosis	Yes	28/205	1		0.137 (0.093, 0.191)
Ferenci 2014 PIII	Multinational	49	59	0% cirrhosis	Yes	9/209	1		0.043 (0.020, 0.080)
Kowaley 20146 Kumada 2015	Japan	48 61	63	0% cirrhosis	Mixed	2/79 9/215	1		0.042 (0.003, 0.088)
awitz 2015	Multinational	55	51	0% cirrhosis	Mixed	8/82	1		0.098 (0.043. 0.183)
Subtotal (1 ² = 69.8%, p = 0.003)	Multinational	47	52	S20% cirriosis	Yes	//83	1	ਤ	0.084 (0.035, 0.166) 0.065 (0.043, 0.097)
	dasahuwis + ribewisia							1 [•] !	
Feld 2014	Multinational	49	43	0% cirrhosis	Yes	112/473	1	<u>-</u>	0.237 (0.199. 0.278)
erenci 2014 PIV	Multinational	52	30	0% cirrhosis	Yes	21/100	1	_ ! — 	0.210 (0.135, 0.303)
-erenci 2014 PIII Andreone 2014	Multinational Multinational	48 54	49 50	0% cirrhosis 0% cirrhosis	Yes	11/210 19/91	1	*** ;	0.052 (0.026, 0.092) 0.209 (0.131, 0.307)
Kowdley 2014b	Multinational	50	44	0% cirrhosis	Yes	1/79	1		0.013 (0.000, 0.069)
Lalezari 2015 Dore 2016 M1	U.S. Multinational	48 48	34 39	0% cirrhosis ≤20% cirrhosis	Mixed	19/38 21/153	1		0.500 (0.334, 0.666) 0.137 (0.087, 0.202)
Dore 2016 M2	Multinational	47	46	≤20% cirrhosis	No	10/101	1		0.099 (0.049, 0.175)
Grebelv 2018b	Multinational	48	23	≤20% cirrhosis	Yes	20/87	1		0.230 (0.148. 0.332)
Vaked 2016	Egypt	49	30	≤20% cirrhosis	Yes	17/100	4	÷	0.143 (0.078, 0.252) 0.170 (0.102, 0.258)
Subtotal (I ² = 90.0%, p = 0.000)								\sim	0.152 (0.096, 0.232)
Heterogeneity between groups: (p = 0.016							1 1	
Dverall (I ² = 82.548%, p = 0.000	1)							•	0.111 (0.091, 0.135)
								0 2 4	
					Pr	oportion		v .2 .4	.0
						oportion			



					Across diff	erent genotype	S	1	
Author year	Country	Age	% Female	Fibrosis stage	Tx naïve	Events/Total	Genotype		Proportion (95% CI)
edipasvir/sofosbuvir Vidahl 2014 Kowdley 2014a Jane 2015 Abergel 2016a Ahmed 2018 Abergel 2016b Subtotal (I ² = 71.5%, p	Multinational U.S. New Zealand France Egypt France = 0.004)	52 53 47 52 51 61	41 41 42 50 35 48	≤20% cirrhosis 0% cirrhosis ≤20% cirrhosis ≤20% cirrhosis Unclear/NR ≤20% cirrhosis	Yes Yes Mixed Yes Yes Yes	24/214 24/431 6/50 4/44 1/100 3/41	1 3 or 6 4 5		$\begin{array}{c} 0.112 \ (0.073, \ 0.162) \\ 0.056 \ (0.036, \ 0.082) \\ 0.120 \ (0.045, \ 0.243) \\ 0.091 \ (0.025, \ 0.217) \\ 0.010 \ (0.000, \ 0.054) \\ 0.073 \ (0.015, \ 0.199) \\ 0.068 \ (0.042, \ 0.109) \end{array}$
Sofosbuvir/velpatasvir Everson 2015 Grebely 2018a Subtotal (l² = 49.7%, p	U.S. Multinational = 0.159)	49 48	39 28	0% cirrhosis ≤20% cirrhosis	Yes NR	7/77 4/103	1 Mixed	↓ ↓ ↓	0.091 (0.037, 0.178) 0.039 (0.011, 0.096) 0.061 (0.034, 0.108)
Elbasvir/grazoprevir Sulkowski 2015	Multinational	52	48	0% cirrhosis	Yes	5/43	1		0.116 (0.049, 0.250)
Sofosbuvir/daclatasvir Sulkowski 2014 Velson 2015 Subtotal (l² = 0.0%, p :	U.S. U.S. = 0.520)	55 53	51 43	≤20% cirrhosis ≤20% cirrhosis	Yes Yes	2/41 13/152	1 3		0.049 (0.006, 0.165) 0.086 (0.046, 0.142) 0.078 (0.047, 0.125)
Ombitasvir/paritaprevir, Ferenci 2014 PIV Ferenci 2014 PIII Andreone 2014 Kowdley 2014b Lawitz 2015 Subtotal (I ² = 71.5%, p	ritonavir/dasabuvir Multinational Multinational Multinational Multinational al Multinational = 0.007)	51 49 54 48 55	37 59 40 43 51	0% cirrhosis 0% cirrhosis 0% cirrhosis 0% cirrhosis 0% cirrhosis	Yes Yes No Yes Mixed	33/205 13/209 12/95 13/79 6/82	1 1 1 1		0.161 (0.113, 0.219) 0.062 (0.034, 0.104) 0.126 (0.067, 0.210) 0.165 (0.091, 0.265) 0.073 (0.027, 0.152) 0.111 (0.077, 0.159)
Dmbitasvir/paritaprevir/ Ferenci 2014 PIII Ferenci 2014 PIV Andreone 2014 Feld 2014 Kowdley 2014b Hezode 2015 Subtotal (I ² = 72.7%, p	ritonavir/dasabuvir + Multinational Multinational Multinational Multinational Multinational = 0.003)	ribaviri 48 52 54 49 50 47	n 49 30 50 43 44 30	0% cirrhosis 0% cirrhosis 0% cirrhosis 0% cirrhosis 0% cirrhosis 0% cirrhosis	Yes No Yes Yes Mixed	9/210 14/100 12/91 65/473 10/79 9/81	1 1 1 1 4		0.043 (0.020, 0.080) 0.140 (0.079, 0.224) 0.132 (0.070, 0.219) 0.137 (0.108, 0.172) 0.127 (0.062, 0.220) 0.111 (0.052, 0.200) 0.109 (0.078, 0.149)
leterogeneity between Overall (I ² = 70.020%,	groups: p = 0.313 p = 0.000)								0.087 (0.069, 0.110)



					Across diffe	erent genotypes		I	
Author year	Country	Age	% Female	Fibrosis stage	Tx naïve	Events/Total	Genotype		Proportion (95% CI)
_edipasvir/sofosbu	vir								
3ane 2015	New Zealand	47	42	≤20% cirrhosis	Mixed	3/50	3 or 6		0.060 (0.019, 0.170
Simeprevir/sofosbu	uvir								
Pott-Junior 2019	Brazil	53	48	0% cirrhosis	Mixed	3/60	1		0.050 (0.016, 0.144)
Sofosbuvir/velpata:	svir								
Grebely 2018a	Multinational	48	28	≤20% cirrhosis	NR	4/103	Mixed		0.039 (0.015, 0.099)
Sofosbuvir/daclata	svir								
Sulkowski 2014	U.S.	55	51	≤20% cirrhosis	Yes	1/41	1	╞╋┼──	0.024 (0.001, 0.129)
Pott-Junior 2019 Subtotal (I ² = 0.0%	Brazil 6, p = 0.700)	56	53	0% cirrhosis	Mixed	1/65	1	<u>-</u>	0.015 (0.000, 0.083) 0.019 (0.005, 0.072)
Ombitasvir/paritapr	evir/ritonavir/da	sabuv	vir + ribavirir	ı					
_alezari 2015	U.S.	48	34	0% cirrhosis	Mixed	4/38	1	 ∎	0.105 (0.029, 0.248)
3rebely 2018b Subtotal (l ² = 0.0%	Multinational 6, p = 0.806)	48	23	≤20% cirrhosis	Yes	11/87	1	$\langle \rangle$	0.126 (0.065, 0.215) 0.120 (0.074, 0.189)
Heterogeneity betw Dverall (I ² = 42.84	veen groups: p 3%, p = 0.105)	= 0.03	36					\$-	0.058 (0.034, 0.097)
leterogeneity betw Dverall (I ² = 42.84	veen groups: p 3%, p = 0.105)	= 0.03	36						0.058 (0.034

Abbreviations: CI = confidence interval; NR = not reported; Tx = treatment; U.S. = United States.

				Acro	ss different	genotypes			
Author year	Country	Age	% Female	Fibrosis stage	Tx naïve	Events/Total	Genotype		Proportion (95% CI)
edipasvir/sofosbuvir Afdahl 2014 Lawitz 2014b Kowdley 2014a Subtotal (I² = 80.3%, p = 0.00	Multinational U.S. U.S. 6)	52 47 53	41 36 40	≤20% cirrhosis 0% cirrhosis 0% cirrhosis	Yes Yes Yes	16/214 1/39 3/215	1 1 1		0.075 (0.043, 0.119) 0.026 (0.001, 0.135) 0.014 (0.003, 0.040) 0.033 (0.018, 0.088)
Simeprevir/sofosbuvir .awitz 2014a Kwo 2016 Pott-Junior 2019 Subtotal (I² = 0.0%, p = 0.631	U.S. Canada & U.S. Brazil)	56 56 53	42 47 48	0% cirrhosis 0% cirrhosis 0% cirrhosis	No Mixed Mixed	1/14 10/155 6/60	1 1 1		0.071 (0.002, 0.339) 0.065 (0.031, 0.115) 0.100 (0.038, 0.205) 0.074 (0.047, 0.116)
Sofosbuvir/velpatasvir Pianko 2015 Everson 2015 (Part A) Subtotal (l² = 44.9%, p = 0.17	Australia, New Zealand, U U.S. '8)	.S56 49	35 39	≤20% cirrhosis 0% cirrhosis	No Yes	9/80 4/77	3 Mixed		0.112 (0.053, 0.203) 0.052 (0.014, 0.128) 0.083 (0.049, 0.137)
Elbasvir/grazoprevir Sulkowski 2015	Multinational	49	30	0% cirrhosis	Yes	2/43	1		0.047 (0.012, 0.168)
Sofosbuvir/daclatasvir Pott-Junior 2019	Brazil	56	53	0% cirrhosis	Mixed	1/65	1	.	0.015 (0.002, 0.101)
Glecaprevir/pibrentasvir Chayama 2018	Japan	64	64	0% cirrhosis	No	3/129	1		0.023 (0.005, 0.066)
Dmbitasvir/paritaprevir/ritonav Ferenci 2014 PIV Ferenci 2014 PIII Andreone 2014 Dore 2016 M1 Subtotal (I ² = 65.7%, p = 0.03	ir/dasabuvir Multinational Multinational Multinational Multinational 3)	51 49 54 47	37 59 40 52	0% cirrhosis 0% cirrhosis 0% cirrhosis ≤20% cirrhosis	Yes Yes No Yes	10/205 8/209 1/95 0/83	1 1 1 1	011+	$\begin{array}{c} 0.049 \ (0.024, \ 0.088) \\ 0.038 \ (0.017, \ 0.074) \\ 0.011 \ (0.000, \ 0.057) \\ 0.000 \ (0.000, \ 0.043) \\ 0.026 \ (0.010, \ 0.067) \end{array}$
Ombitasvir/paritaprevir/ritonav erenci 2014 PIV erenci 2014 PIII Andreone 2014 elezari 2015 Jore 2016 M1 Jore 2016 M2 Subtotal (I ^e = 57.0%, p = 0.03	ir/dasabuvir + ribavirin Multinational Multinational Multinational U.S. Multinational Multinational Multinational	52 48 54 49 48 46 47	30 49 50 43 34 39 46	0% cirrhosis 0% cirrhosis 0% cirrhosis 0% cirrhosis 0% cirrhosis ≤20% cirrhosis ≤20% cirrhosis	Yes Yes No Yes Mixed Yes No	5/100 12/210 8/91 51/473 6/38 12/153 3/101	1 1 1 1 1		0.050 (0.016, 0.113) 0.057 (0.030, 0.098) 0.088 (0.039, 0.166) 0.108 (0.081, 0.139) 0.158 (0.060, 0.313) 0.078 (0.041, 0.133) 0.078 (0.045, 0.084) 0.076 (0.055, 0.103)
Heterogeneity between group Overall (l² = 69.599%, p = 0.0	s: p = 0.017 00)							⇔	0.054 (0.041, 0.071)

Abbreviations: CI = confidence interval; Tx = treatment; U.S. = United States.



Abbreviations: CI = confidence interval; DAA = direct acting antiviral; HR = hazard ratio; IFN = interferon; NR = not reported; SVR = sustained virologic response; U.S. = United States.



Abbreviations: CI = confidence interval; GT1 = genotype 1; HR = hazard ratio; IFN = interferon; NR = not reported; SVR = sustained virologic response.



Abbreviations: CI = confidence interval; HR = hazard ratio; IFN = interferon; NR = not reported; SVR = sustained virologic response; U.S. = United States.

Author year	Duration (Years)	Country	F0-F1 Only	Cirrhosis %	Treatment	Adjustment	GT1 %	N (SVR/No SVR)		HR (95% CI)
mai 1999	4.0	Asia	No	0-10%	IFN	Partial	NR	151/268	<	0.06 (0.01, 0.48)
Kasahara 1998	3.1	Asia	No	0-10%	IFN	Full	58%	313/304	<u>+</u>	0.19 (0.06, 0.58)
keda 1999	5.4	Asia	No	>10%	IFN	Full	67%	145/585		0.33 (0.12, 0.96)
Yoshida 1999	4.3	Asia	No	0-10%	IFN	Partial	70%	789/1568	_ i _	0.32 (0.14, 0.70)
Tanaka 2000	4.8	Asia	No	0-10%	IFN	Full	75%	175/419	-+	0.29 (0.07, 1.28)
Okanoue 2002	5.6	Asia	No	0-10%	IFN	Partial	NR	426/358		0.13 (0.06, 0.27)
zumi 2005	Unclear	Asia	No	0-10%	IFN	Unclear	50%	155/340		0.36 (0.04, 0.83)
ru 2006	5.2	Asia	No	>10%	IFN	Full	46%	715/342	_ _	0.24 (0.11, 0.52)
Arase 2007	7.4	Asia	No	>10%	IFN	Full	60%	140/360	_ _	0.19 (0.08, 0.45)
Kurokawa 2009	3.0	Asia	No	0-10%	IFN	Partial	89%	139/264	_ 	0.28 (0.08, 0.96)
Asahina 2010	7.5	Asia	No	0-10%	IFN	Full	71%	686/1356	- -	0.38 (0.18, 0.83)
Tateyama 2011	8.2	Asia	No	>10%	IFN	Full	72%	139/234		0.14 (0.04, 0.52)
Maruoka 2012	9.9	Asia	No	0-10%	IFN	Full	73%	221/356		0.12 (0.03, 0.41)
Osaki 2012	4.1	Asia	No	0-10%	IFN	Partial	60%	185/197		0.12 (0.01, 0.94)
Dohmen 2013	4.8	Asia	No	Unclear	IFN	Partial	67%	285/189		0.39 (0.32, 0.48)
Dieperink 2014	7.5	U.S.	No	>10%	IFN	Full	70%	222/314	÷∎-√	0.41 (0.18, 0.96)
El-Serag 2014	5.2	U.S.	Unclear	Unclear	NR	Full	55%	7577/8767	▲	0.30 (0.23, 0.38)
_ee 2017	2.6	Asia	No	>10%	IFN	Full	51%	306/183	 +	0.09 (0.02, 0.40)
oannou 2018	6.1	U.S.	No	>10%	Mixed	Full	77%	28655/23231	i	0.32 (0.28, 0.37)
Carrat 2019	2.8	Europe	No	0-10%	DAA	Full	66%	3286/146	_	0.22 (0.03, 1.76)
Mixed effects m	odel estima	ate Overal	l (I-square	d = 18.7%, p=	0.222)				\$	0.29 (0.23, 0.38)

Abbreviations: CI = confidence interval; DAA = direct acting antiviral; HCC = hepatocellular carcinoma; HR = hazard ratio; IFN = interferon; NR = not reported; SVR = sustained virologic response; U.S. = United States.

Table 1. Sustained Virologic Response Rates in Older Antiviral Regimens

Treatment	Sustained virologic response rate
Placebo	<2
Interferon monotherapy	6 to 16
Interferon plus ribavirin	33 to 41
Pegylated interferon alone	25 to 39
Pegylated interferon plus ribavirin	39 to 43 (genotypes 1 and 4)
	76 to 83 (genotypes 2 and 3)
Boceprevir or telaprevir plus pegylated interferon and ribavirin	68 to 72 (genotype 1)
Source: ^{91,277}	

Screening for Hepatitis C Virus Infection

 Table 2. Currently Recommended Direct Acting Antivirals and Alternative Regimens for Treatment

 Naïve Adults With HCV Infection Without Cirrhosis

Recommended		Duration of Treatment	
or Alternative	Regimen	(weeks)	Genotype
	Glecaprevir 300 mg + pibrentasvir 120 mg	8	1a, 1b, 2, 3, 4, 5, 6
	Ledipasvir 90 mg + sofosbuvir 400 mg	8	1a, 1b
Recommended		1.0	
Regimens	Ledipasvir 90 mg + sofosbuvir 400 mg	12	1a, 1b, 4, 5, 6
	Elbasvir 50 mg + grazoprevir 100 mg	12	1a, 1b, 4
	Sofosbuvir 400 mg + velpatasvir 100 mg	12	1a, 1b, 2, 3, 4, 5, 6
	Daclatasvir 60 mg + sofosbuvir 400 mg	12	1a, 1b, 2, 3
	Paritaprevir 150 mg + ritonavir 100 mg + ombitasvir 25 mg + weight-based ribavirin	12	4
	Simeprevir 150 mg + sofosbuvir 400 mg	12	1a, 1b
Alternative Regimens	Elbasvir 50 mg + grazoprevir 100 mg + weight-based ribavirin	16	1a
	Paritaprevir 150 mg + ritonavir 100 mg + ombitasvir 25 mg + dasabuvir ER 600 mg or dasabuvir 250 mg 2x/day + weight- based ribavirin	12	1a
	Paritaprevir 150 mg + ritonavir 100 mg + ombitasvir 25 mg + dasabuvir ER 600 mg or dasabuvir 250 mg 2x/day	12	1b
Source: AASLD/IE	USA , available at: <u>https://www.hcvguidelines.org/treatment-naive</u>		

Note: Recommended regimens are those that are favored for most patients in a given group, based on optimal efficacy, favorable tolerability and toxicity profiles, and treatment duration. Alternative regimens are those that are effective but, relative to recommended regimens, have potential disadvantages, limitations for use in certain patient populations, or less supporting data than recommended regimens. In certain situations, an alternative regimen may be an optimal regimen for an individual patient.

Abbreviations: AASLD = American Association for the Study of Liver Diseases; ER = extended release; IDSA = Infectious Disease Society of America.

Table 3. Currently Recommended Antiviral Regimens for Treatment-Experienced Adults With HCV Infection Without Cirrhosis

Recommended		Duration of	
or Alternative	Regimen	treatment (weeks)	Genotype
	Glecaprevir 300 mg + pibrentasvir 120 mg	8	1a, 1b, 2, 4, 5, 6
	Same as above	12	1
Pacammandad	Elbasvir 50 mg + grazoprevir 100 mg	12	1a, 1b, 4
Pegimens	Ledipasvir 90 mg + sofosbuvir 400 mg	12	1a, 1b, 4, 5, 6
Regimens	Sofosbuvir 400 mg + velpatasvir 100 mg	12	1a, 1b, 2, 3, 4, 5, 6
	Sofosbuvir 400 mg + velpatasvir 100mg + voxilaprevir	12	1a
	100mg		
	Daclatasvir 60 mg + sofosbuvir 400 mg	12	1a, 1b, 2, 3
	Elbasvir 50 mg + grazoprevir 100 mg + ribavirin	12	1b
	Same as above	12 to 16	1a
	Same as above	16	1a, 4
	Ledipasvir 90 mg + sofosbuvir 400 mg + ribavirin	12	1a, 1b
	Simeprevir 150 mg + sofosbuvir 400 mg	12	1a, 1b
	Ombitasvir 25 mg + paritaprevir 150 mg + ritonavir 100	12	1b
Alternative	mg + dasabuvir ER 600 mg or dasabuvir 250 mg 2x/day		
Regimens	Ombitasvir 25 mg + paritaprevir 150 mg + ritonavir 100	12	1a
	mg + dasabuvir ER 600 mg or dasabuvir 250 mg 2x/day +		
	weight-based ribavirin		
	Ombitasvir 25 mg + paritaprevir 150 mg + ritonavir 100	12	4
	mg + weight-based ribavirin		
	Sofosbuvir 400 mg + velpatasvir 100mg + voxilaprevir	12	3
	100mg		
	Glecaprevir 300 mg + pibrentasvir 120 mg	16	3

Source: AASLD/IDSA, available at: https://www.hcvguidelines.org/treatment-experienced, up to date as of June 1, 2019.

Note 1: Recommended regimens are those that are favored for most patients in a given group, based on optimal efficacy, favorable tolerability and toxicity profiles, and treatment duration. Alternative regimens are those that are effective but, relative to recommended regimens, have potential disadvantages, limitations for use in certain patient populations, or less supporting data than recommended regimens. In certain situations, an alternative regimen may be an optimal regimen for an individual patient. **Note 2:** Table does not list regimens for those with prior DAA treatment experience.

Abbreviations: AASLD = American Association for the Study of Liver Diseases; ER = extended release; IDSA = Infectious Disease Society of America.

Table 4. Currently Recommended Antiviral Regimens for Adolescents ≥12 Years Old or Weighing at Least 35 kg, Without Cirrhosis or With Compensated Cirrhosis

Regimen*	Duration of treatment (weeks)	Genotype
Ledipasvir 90 mg + sofosbuvir 400 mg for patients who are treatment-naive without cirrhosis or with compensated cirrhosis, or treatment-experienced without cirrhosis	12	1
Sofosbuvir 400 mg + weight-based ribavirin for patients who are treatment- naive or treatment-experienced without cirrhosis or with compensated cirrhosis	12	2
Sofosbuvir 400 mg + weight-based ribavirin for patients who are treatment- naive or treatment-experienced [†] without cirrhosis or with compensated cirrhosis	24	3
Ledipasvir 90 mg + sofosbuvir 400 mg for patients who are treatment-naive or treatment-experienced without cirrhosis or with compensated cirrhosis	12	4, 5, 6

Source: AASLD/IDSA https://www.hcvguidelines.org/unique-populations/children

Note: Recommended regimens are those that are favored for most patients in a given group, based on optimal efficacy, favorable tolerability and toxicity profiles, and treatment duration.

* Glecaprevir + pibrentasvir approved by the U.S. Food and Drug Administration in April 2019 for children 12 to 17 years of age for genotypes 1 through 6, but has not been incorporated in the AASLD recommendations as of June 1, 2019.

Abbreviations: AASLD = American Association for the Study of Liver Diseases; IDSA = Infectious Disease Society of America.

Table 5. U.S. Screening Guidelines

Group	Recommendation						
AASLD-IDSA65	One-time HCV testing is recommended for persons born between 1945 and 1965 (regardless of						
	country of birth) without prior ascertainment of risk.						
	Other persons should be screened for risk factors for HCV infection, and one-time testing should						
	be performed for all persons with behaviors, exposures, and conditions associated with an						
	increased risk of HCV infection						
	All persons with active HCV infection should be linked to a clinician who is prepared to provide						
	comprehensive management						
CDC ⁸⁷	Persons for whom HCV testing Is recommended:						
	Adults born from 1945 through 1965 should be tested once (without prior ascertainment of HCV						
	risk factors)						
	HCV testing is recommended for those who:						
	Currently inject drugs						
	• Ever injected drugs, including those who injected once or a few times many years ago						
	Have certain medical conditions, including persons:						
	 who received clotting factor concentrates produced before 1987 						
	 who were ever on long-term hemodialysis 						
	 with persistently abnormal ALT levels 						
	 who have HIV infection 						
	Were prior recipients of transfusions or organ transplants, including persons who:						
	 were notified that they received blood from a donor who later tested positive for HCV infection 						
	 received a transfusion of blood, blood components, or an organ transplant before July 1992 						
	 HCV- testing based on a recognized exposure is recommended for: 						
	 Healthcare, emergency medical, and public safety workers after needle sticks. 						
	sharps, or mucosal exposures to HCV-positive blood						
	 Children born to HCV-positive women 						
	Note: For persons who might have been exposed to HCV within the past 6 months, testing for						
	HCV RNA or follow-up testing for HCV antibody is recommended.						
Abbreviations: AA	SID-IDSA - American Association for the Study of Liver Diseases-Infectious Diseases Society of America:						

Abbreviations: AASLD-IDSA = American Association for the Study of Liver Diseases-Infectious Diseases Society of America; ALT = alanine aminotransferase; CDC = Centers for Disease Control and Prevention; HCV = hepatitis C virus; RNA = ribonucleic acid.

Author year <i>Quality</i>	N	Elective Cesarean or Cesarean not specified	E	Vaginal/ Emergent Cesarean	Comments/Results (95% CI)
Ceci 2001 ¹⁰⁸ <i>Fair</i>	78*	No association (data NR)	No (dat	association ta NR)	No significant association in multivariate analysis (data NR)
Gibb 2000 ¹⁰⁵ Fair	424†	0/31 (0%)	29/3	393 (7.4%)	OR 0 (0 to 0.87), p=0.04, adjusted for HIV status and breastfeeding
Mast 2005 ¹⁰⁴ Good	188*	0/12 (0%)	7/16	69 (4.1%)	RR 0.87 (0.05 to 14) Excluded from multivariate analyses due to lack of significance in univariate analysis
Resti 2002 ¹⁰⁷ Good	1,301 [‡]	22/337 (5.8%)	73/9	924 (7.9%)	OR for vaginal delivery 1.17 (0.92 to 1.41), unadjusted [§] OR for vaginal delivery 1.20 (0.93 to1.55), adjusted for maternal HCV RNA status, maternal HIV status, injection drug use, type of feeding [§]
Tovo 2005 ¹⁰⁶ EPHN <i>Good</i>	1,034*	NR	NR		OR 1.57 (0.88 to 2.83), p=0.13, unadjusted OR 1.59 (0.88 to 2.86), p=0.13 adjusted for sex, mode of delivery, prematurity, and breastfeeding
Total	3,025				

*0% HIV coinfected.

[†] 5% HIV coinfected.

[‡]14% HIV coinfected.

[§] Study appears to have reversed reference standard; Calculation to adjust reference standard gives unadjusted OR for vaginal delivery (ref): 0.85 (0.71 to 1.09); Adjusted OR for vaginal delivery (ref): 0.83 (0.65 to 1.08).

Abbreviations: CI = confidence interval; EPHN = European Paediatric Hepatitis C Virus Network; HCV = hepatitis C virus; NR = not reported; OR = odds ratio; RR = relative risk.

Table 7. Duration of Membrane Rupture and Mother-to-Infant Transmission of HCV Infection

Author year Quality	N	Duration of Membrane Rupture (hours)	Comments/Results (95% CI)
Mast 2005 ¹⁰⁴ Good	189*	<1 vs. 1 to 5 vs. 6 to 12 vs. ≥13: 0/53 vs. 1/59 (1.7%) vs. 4/40 (10%) vs. 2/30 (6.7%), p=0.02	Membrane rupture >6 hours OR, 9.3 (1.5 to 179.7), adjusted for maternal demographic characteristics, HCV RNA level, fetal monitoring, history of IVDU, and cigarette smoking during pregnancy.
Total	189		

*0% HIV coinfected.

Abbreviations: CI = confidence interval; HCV = hepatitis c virus; IVDU = intravenous drug use; OR = odds ratio; RNA = ribonucleic acid.

Table 8. Fetal Monitoring and Risk of Mother-to-Infant Transmission of HCV Infection

Author year			Comments/ Results
Quality	N	Fetal Monitoring During Delivery	(95% CI)
Mast 2005 ¹⁰⁴	188 [*]	Internal vs. external:	RR 7.7 (1.9 to 31.6), p=0.02,
Good		3/16 (18.8%) vs. 4/165 (2.4%),	unadjusted
			Internal fetal monitoring, OR 6.7
			(1.1 to 35.9), adjusted for maternal
			demographic characteristics, HCV
			RNA level, history of IVDU, and
			cigarette smoking during
			pregnancy.
Total	188		

*0% HIV coinfected. **Abbreviations:** CI = confidence interval; IVDU = intravenous drug use; HCV = hepatitis C virus; OR = odds ratio; RNA = ribonucleic acid; RR = relative risk.

Table 9. Breastfeeding and Risk of Mother-to-Infant Transmission of HCV Infection

Author year				Comments/Results
Quality	N	Breast Fed	Formula Fed	(95% CI)
Gibb 2000 ¹⁰⁵ <i>Fair</i>	414*	7.7% (2.2 to 17.8)	6.7% (3.7 to 10.6)	OR 1.52 (0.35 to 5.12), adjusted for HIV status and mode of delivery
Resti 2002 ¹⁰⁷ Good	1,281†	22/360 (6.1%)	73/921 (7.9%)	OR 0.86 (0.61 to 1.10) OR 0.95 (0.58 to 1.40), adjusted for maternal HCV RNA status, maternal HIV-1 status, maternal IVDU, type of feeding, mode of delivery
Tovo 2005 ¹⁰⁶ EPHN <i>Good</i>	1,034‡	NR	NR	OR 0.88 (0.48 to 1.61), unadjusted OR 0.92 (0.50 to 1.70), adjusted for sex, prematurity, and mode of delivery
Total	3,645			

* 5% HIV coinfected.

†14% HIV coinfected.

[‡]0% HIV coinfected.

Abbreviations: CI = confidence interval; EPHN = European Paediatric Hepatitis C Virus Network; HCV = hepatitis C virus; IVDU = intravenous drug use; NR = not reported; OR = odds ratio; RNA = ribonucleic acid.

Table 10. Trials of Sustained Virologic Response With Direct Acting Antiviral Regimens in Adults

Author year		Primary	Mean age	Proportion female	Proportion with	Proportion	
Study name	Treatment Regimen	genotype(s)	(years)	gender	cirrhosis	treatment-naïve	SVR
Chayama 2018 ¹⁹⁷ CERTAIN-1	Glecapravir + pibrentasvir	1	64	64%	0%	73%	99% (128/129)
Poordad 2017 ¹⁹⁴	Glecapravir + pibrentasvir	1	58	18%	0%	0%	92% (46/50)
Zeuzem 2018 ¹⁶⁷	Glecapravir + pibrentasvir	1	53	52%	0%	62%	99% (663/667)
Kumada 2017 ¹⁵²	Grazoprevir + elbasvir	1	61	62%	0%*	66%	97% (219/227)
Sulkowski 2015 ¹⁶⁰ C-WORTHY	Grazoprevir + elbasvir	1	51	51%	15%	100%	95% (122/129)
Zeuzem 2015 ¹⁶⁶ C-EDGE	Grazoprevir + elbasvir	1	52	46%	22%	100%	95% (273/288)
Kowdley 2014a ¹⁹⁰ ION-3	Ledipasvir + sofosbuvir	1	53	41%	0%	100%	95% (408/431)
Afdahl 2014 ¹⁸⁵ ION-1	Ledipasvir + sofosbuvir	1	52	41%	0%*	100%	100% (357/357)
Chuang 2016 ¹⁴⁵	Ledipasvir + sofosbuvir	1	55	58%	12%	100%	98% (83/85)
Lawitz 2014b ¹⁹³ LONESTAR	Ledipasvir + sofosbuvir	1	48	38%	0%	100%	97% (58/60)
Lim 2016 ¹⁵⁶	Ledipasvir + sofosbuvir	1	54	43%	9%	100%	100% (46/46)
Wei 2018 ¹⁶³	Ledipasvir + sofosbuvir	1	47	50%	16%	52%	100% (206/206)
Grebely 2018 ¹⁴⁹ D3FEAT	Ombitasvir + paritaprevir + ritonavir + dasabuvir	1	48	22%	8%	100%	91% (73/80)
Lalezari 2015 ¹⁹²	Ombitasvir + paritaprevir + ritonavir + dasabuvir	1	48	34%	0%	95%	97% (37/38)
Kwo 2016 ¹⁵³ OPTIMIST-1	Simeprevir + sofosbuvir	1	56	47%	0%	74%	97% (150/155)
Lawitz 2014a ¹⁵⁴ COSMOS	Simeprevir + sofosbuvir	1	56	30%	0%	0%	95% (61/64)
Pott-Junior 2019 ¹⁵⁹	Simeprevir + sofosbuvir	1	53	48%	0%	60%	93% (56/60)
Pott-Junior 2019 ¹⁵⁹	Sofosbuvir + daclatasvir	1	56	53%	0%	60%	100% (65/65)
Sulkowski 2014 ¹⁶¹ A1444040 Study	Sofosbuvir + daclatasvir	1	55	51%	13%	100%	98% (80/82)
Everson 2015 ¹⁴⁶	Sofosbuvir + velpatasvir	1	49	39%	0%	100%	100% (28/28)
Feld 2015 ¹³⁹ ASTRAL-1	Sofosbuvir + velpatasvir	1	54	60%	0%*	68%	98% (251/255)
Ferenci 2014 ¹⁸⁸ PEARL IV	Ombitasvir + paritaprevir + ritonavir + dasabuvir	1a	51	35%	0%	100%	92% (282/305)

Table 10. Trials of Sustained Virologic Response With Direct Acting Antiviral Regimens in Adults

Author year		Primary	Mean age	Proportion female	Proportion with	Proportion	
Study name	Treatment Regimen	genotype(s)	(years)	gender	cirrhosis	treatment-naïve	SVR
Kowdley 2014b ¹⁹¹	Ombitasvir + paritaprevir +	1a	50	42%	0%	75%	86% (183/212)
AVIATOR	ritonavir + dasabuvir						
Feld 2014 ¹⁸⁷	Ombitasvir + paritaprevir +	1a	49	43%	Unclear	100%	95% (307/322)
SAPPHIRE-1	ritonavir + dasabuvir						
Lawitz 2015 ¹⁵⁵	Ombitasvir + paritaprevir +	1b	55	51%	0%	51%	93% (76/82)
PEARL-1	ritonavir						
Andreone 2014 ¹⁸⁶	Ombitasvir + paritaprevir +	1b	54	45%	0%	0%	98% (176/179)
PEARL-II	ritonavir + dasabuvir		40	400/	00/	1000/	000/ (440/454)
Feld, 2014 ¹⁰⁷	Ombitasvir + paritaprevir +	10	49	43%	0%	100%	98% (148/151)
SAPPHIRE-1	ritonavir + dasabuvir	41-	40	E 40/	00/	4000/	000((440/440)
	Ombitasvir + paritaprevir +	1D	48	54%	0%	100%	99% (416/419)
PEARL III Kowdlov 2014b ¹⁹¹	ntonavir + dasabuvir	16	50	400/	00/	600/	1000/ (112/112)
	ombitasvir + paritaprevir +	D	50	42%	0%	68%	100% (113/113)
Kumada 2015 ¹⁵¹		16	61	62%	0%	65%	04.0% (204/215)
		1D	01	03 /0	0 /0	05 /0	94.976 (204/215)
Toyoda 2018 ¹⁹⁹		2	57	53%	0%	83%	98% (88/90)
CERTAIN-2		2	57	5570	070	0070	3078 (00/30)
Feld 2015 ¹³⁹	Sofosbuvir + velpatasvir	2	54	60%	0%*	68%	100% (93/93)
ASTRAL-1		-	•	00,0	0,0	00,0	
Foster 2015 ¹⁴⁷	Sofosbuvir + velpatasvir	2	57	36%	14%	86%	99% (133/134)
ASTRAL-2							, , , , , , , , , , , , , , , , , , ,
Zeuzem 2018 ¹⁶⁷	Glecapravir + pibrentasvir	3	47	41%	0%	100%	95% (149/157)
ENDURANCE-3							
Nelson 2015 ¹⁵⁷	Sofosbuvir + daclatasvir	3	55	41%	0%	59%	96% (105/109)
ALLY-3							
Zeuzem 2018 ¹⁶⁷	Sofosbuvir + daclatasvir	3	49	55%	0%	100%	97% (111/115)
ENDURANCE-3							
Everson 2015 ¹⁴⁶	Sofosbuvir + velpatasvir	3	50	37%	0%	100%	93% (25/27)
Foster 2015 ¹⁴⁷	Sofosbuvir + velpatasvir	3	49	39%	0%*	74%	97% (191/197)
ASTRAL-3	·						
Pianko 2015 ¹⁵⁸	Sofosbuvir + velpatasvir	3	55	34%	0%	0%	100% (53/53)
Brown 2018 ¹⁴⁴	Grazoprevir + elbasvir	4	52	58%	0%	100%	90% (9/10)
C-SCAPE					- / -		
Zeuzem 2015 ¹⁶⁶	Grazoprevir + elbasvir	4	52	46%	20%	100%	100% (18/18)
C-EDGE			-				
Abergel 2016a ¹⁴²	Ledipasvir + sofosbuvir	4	52	50%	5%	100%	96% (21/22)
Ahmed 2018 ¹⁹⁵	Ledipasvir + sofosbuvir	4	51	35%	Unclear	100%	99% (99/100)
							` '
Hezode 2015 ¹⁸⁹	Ombitasvir + paritaprevir +	4	48	29%	0%	46%	100% (91/91)
PEARLI	ritonavir + dasabuvir		1				

Table 10. Trials of Sustained Virologic Response With Direct Acting Antiviral Regimens in Adults

Author year		Primary	Mean age	Proportion female	Proportion with	Proportion	
Study name	Treatment Regimen	genotype(s)	(years)	gender	cirrhosis	treatment-naïve	SVR
Waked 2016 ¹⁶²	Ombitasvir + paritaprevir +	4	49	30%	2%	100%	94% (94/100)
AGATE-II	ritonavir + dasabuvir						
Feld 2015 ¹³⁹	Sofosbuvir + velpatasvir	4	54	60%	0%*	68%	100% (89/89)
ASTRAL-1							
Asselah 2019 ¹⁴³	Glecapravir + pibrentasvir	5	68	57%	13%	83%	96% (22/23)
ENDURANCE-5							
Abergel 2016b ¹⁴¹	Ledipasvir + sofosbuvir	5	61	48%	14%	100%	95% (20/21)
Feld 2015 ¹³⁹	Sofosbuvir + velpatasvir	5	54	60%	0%*	68%	97% (28/29)
ASTRAL-1							
Asselah 2019 ¹⁴³	Glecapravir + pibrentasvir	6	54	52%	10%	93%	98% (60/61)
ENDURANCE-6							
Gane 2015 ¹⁴⁸	Ledipasvir + sofosbuvir	6	51	36%	8%	92%	96% (24/25)
Feld 2015 ¹³⁹	Sofosbuvir + velpatasvir	6	54	60%	0%*	68%	100% (35/35)
ASTRAL-1							
Grebely 2018 ¹⁵⁰	Sofosbuvir + velpatasvir	1, 3	48	28%	9%	NR	94% (97/103)
SIMPLIFY							
Wei 2019b ¹⁶⁵	Sofosbuvir + velpatasvir	1, 3, 6	45	47%	18%	82%	97% (362/375)
Wei 2019a ¹⁶⁴	Grazoprevir + elbasvir	1, 4	48	56%	19%	100%	94% (459/486)
C-CORAL							
Sperl 2016 ¹⁹⁸	Grazoprevir + elbasvir	1, 4, 6	48	57%	17%	78%	99% (128/129)
C-EDGE							
Asselah 2018 ¹⁹⁶	Glecapravir + pibrentasvir	2, 4-6	52	52%	0%	87%	97% (196/203)
SURVEYOR II							
Part 4							
Everson 2015 ¹⁴⁶	Sofosbuvir + velpatasvir	2; 4-6	54	32%	0%	100%	95% (21/22)

*Results for subgroup with no cirrhosis.

Abbreviations: NR = not reported; SVR = sustained virologic response. Study names are not acronyms.

Table 11. Sustained Virologic Response in Comparative Trials of Direct Acting Antiviral Regimens in Adults

Comparison	Author year Study name	Treatment Regimen	Primary genotype(s)	Mean age (years)	Proportion female gender	Proportion with cirrhosis	Proportion treatment- naïve	SVR
DAA vs. Placebo	Feld 2015 ¹³⁹ ASTRAL-1	Sofosbuvir + velpatasvir	Mixed	54	60%	19%	72%	99% (618/624) vs. 0% (0/116); RR 232 (95% Cl, 14.6 to 3680)
	Dore 2016 ¹³⁷ MALACHITE-1	Placebo Ombitasvir + paritaprevir + ritonavir + dasabuvir Telaprevir +	1	46	55%	0%	100%	98% (81/83) vs. 80% (60/75); RR 1.22 (95% Cl, 1.08 to 1.37)
DAA vs. Telaprevir- containing Regimens	Dore 2016 ¹³⁷ MALACHITE-1	ribavirin Ombitasvir + paritaprevir + ritonavir + dasabuvir + ribavirin	1	46	47%	0%	100%	98% (150/153) vs. 80% (60/75); RR 1.23 (95% CI, 1.09 to 1.38)
		Telaprevir + pegylated interferon + ribavirin						
	Dore 2016 ¹³⁷ MALACHITE-2	Ombitasvir + paritaprevir + ritonavir + dasabuvir + ribavirin	1	47	46%	0%	0%	99% (100/101) vs. 66% (31/47); RR 1.50 (95% CI, 1.22 to 1.85)
		Telaprevir + pegylated interferon + ribavirin						
DAA vs. Non-	Foster 2015 ¹⁴⁷ ASTRAL-2	Sofosbuvir + velpatasvir Sofosbuvir + ribavirin	2	57	41%	14%	85%	99% (133/134) vs. 94% (124/132); RR 1.06 (95% CI, 1.01 to 1.11)
recommended DAA	Foster 2015 ¹⁴⁷ ASTRAL-3	Sofosbuvir + velpatasvir Sofosbuvir + ribavirin	3	49	38%	0%*	74%	97% (191/197) vs. 87% (163/187); RR 1.11 (95% CI, 1.05 to 1.18)

Abbreviations: CI = confidence interval; DAA = direct acting antiviral; SVR = sustained virologic response; RR = relative risk. Study names are not acronyms.

		Pooled sustained		
		virologic response rate		
Analysis	Number of trials	(95% CI)	²	p for interaction
Genotype 1 infection	32 (in 30 publications)*137,139,145,146,149,151- 156,159-161,163-167,185-188,190-194,197,198	97.7% (96.6% to 98.4%)	82%	
Ledipasvir / sofosbuvir	6 ^{145,156,163,185,190,193}	99.4% (95.2% to 99.9%)	89%	0.005 (regimens)
Simeprevir / sofosbuvir	3 ^{153,154,159}	95.7% (92.6% to 97.5%)	0%	
Sofosbuvir / velpatasvir	3 ^{139,146,165}	99.0% (95.4% to 99.8%)	27%	
Sofosbuvir / daclatasvir	2 ^{159,161}	98.6% (94.7% to 99.7%)	45%	
Glecaprevir / pibrentasvir	3 ^{167,194,197}	98.6% (94.1% to 99.7%)	78%	
Elbasvir / grazoprevir	5 ^{152,160,164,166,198}	96.7% (95.0% to 97.8%)	55%	
Ombitasvir / paritaprevir / ritonavir / dasabuvir (genotype 1, not sub-typed)	2 ^{149,192}	93.2% (87.0% to 96.6%)	27%	
Ombitasvir / paritaprevir / ritonavir / dasabuvir [†] (genotype 1a)	4 ^{137,187,188,191}	93.7% (89.0% to 96.5%)	77%	
 Ombitasvir / paritaprevir / ritonavir / dasabuvir[‡] (genotype 1b) 	7 ^{137,151,155,186-188,191}	98.2% (96.4% to 99.1%)	68%	
Good quality	12 (in 10 publications) ^{137,139,146,152,159,164,166,187,188,191}	97.2% (95.2% to 98.4%)	82%	0.42 (quality)
Fair quality	20*145,149,151,153- 156,160,161,163,165,167,185,186,190,192-194,197,198	97.9% (96.7% to 98.7%)	76%	
Cirrhosis excluded	22 (in 20 publications) §137,139,146,149,151- 155,159,160,167,185,186,188,190-194	97.1% (95.7% to 98.1%)	82%	0.22 (cirrhosis)
Some cirrhosis (<20% of population)	8 ^{145,156,161,163-166,198}	98.7% (97.1% to 99.4%)	38%	
U.S. or Canada	8 ^{146,153,154,161,190,192-194}	96.7% (93.1% to 98.4%)	82%	0.48 (geographic setting)
Multinational	12 [†]	97.7% (96.4% to 98.6%)	89%	
Other geographic setting	4	98.3% (96.1% to 99.2%)	28%	
Use of ribavirin and/or dasabuvir as recommended	26 (in 25 publications)*137,139,145,146,152- 154,156,159-161,163-166,185,187,188,190-193,197,198	98.3% (97.4% to 98.9%)	60%	
Treatment-naïve	24 (in 23 publications)*137,139,145,146,151- 156,159-161,163,164,166,185,187,188,190-193	97.4% (96.1% to 98.3%)	80%	
Genotype 2 infection	5 ^{139,147,165,196,199}	98.9% (97.5% to 99.5%)	4%	
Sofosbuvir / velpatasvir	3 ^{139,147,165}	99.7% (97.6% to 99.95%)	0%	0.06 (regimens)
Glecaprevir / pibrentasvir	2 ^{196,199}	97.9% (95.0% to 99.1%)	0%	
Good quality	1 ¹³⁹	100% (96.1% to 100%)		0.99 (quality)
Fair quality	4 ^{147,164,196,199}	98.6% (97.0% to 99.4%)	0%	
Cirrhosis excluded	3 ^{139,196,199}	98.5% (96.4% to 99.4%)	36%	0.37 (cirrhosis)
Some cirrhosis (<20% of population)	2 ^{147,164}	99.5% (96.5% to 99.9%)	0%	
U.S. or Canada	1 ¹⁴⁷	99.2% (94.9% to 99.9%)		0.62 (geographic setting)
Multinational	3 ^{139,164,196}	99.0% (97.0% to 99.7%)	33%	
Other geographic setting	1 ¹⁹⁹	97.8% (91.6% to 99.4%)	4%	
Treatment-naïve	1 ¹³⁹	100% (95.4% to 100%)		

Table 12. Sustained Virologic Response Rates With Direct Acting Antiviral Regimens in Adults by Genotype

		Pooled sustained		
		virologic response rate		
Analysis	Number of trials	(95% CI)	²	p for interaction
Genotype 3 infection	6 ^{146,147,157,158,165,167}	95.5% (91.6% to 97.7%)	66%	
Sofosbuvir / velpatasvir	4 ^{146,147,158,165}	95.6% (87.1% to 98.6%)	82%	0.78 (regimens)
Sofosbuvir / daclatasvir	2 ^{157,167}	96.4% (93.0% to 98.2%)	0%	
Glecaprevir / pibrentasvir	1 ¹⁶⁷	94.9% (90.2% to 97.8%)		
Good quality	1 ¹⁴⁶	93.2% (66.8% to 99.0%)		0.66 (quality)
Fair quality	5 ^{147,157,158,164,167}	95.7% (91.6% to 97.8%)	70%	
Cirrhosis excluded	5 ^{146,147,157,158,167}	96.4% (94.6\$ to 97.5%)	14%	0.01 (cirrhosis)
Some cirrhosis (<20% of population)	1 ¹⁶⁵	85.7% (76.5% to 91.7%)		
U.S. or Canada	3 ^{146,147,157}	96.3% (91.4% to 98.4%)	0%	0.55 (geographic setting)
Multinational	3 ^{158,164,167}	94.5% (88.2% to 97.6%)	80%	
Use of ribavirin as recommended	5 ^{146,147,157,158,164,167}	95.2% (91.4% to 97.3%)	0%	
Treatment-naïve	4 ^{146,147,157,167}	96.1% (94.5% to 97.3%)	14%	
Genotype 4 infection	10 ^{139,142,144,162,164,166,189,195,196,198}	98.2% (94.7% to 99.4%)	50%	
Ledipasvir / sofosbuvir	2 ^{142,195}	98.4% (93.7% to 99.6%)	25%	0.14 (regimens)
Sofosbuvir / velpatasvir	1 ¹³⁹	100% (95.9% to 100%)		
Elbasvir / grazoprevir	4 ^{144,164,166,198}	97.3% (83.2% to 99.6%)	0%	
Glecaprevir / pibrentasvir	1 ¹⁹⁶	93.5% (82.1% to 98.6%)		
Ombitasvir / paritaprevir / ritonavir / dasabuvir	2 ^{162,189}	98.7% (72.7% to 99.95%)	88%	
Good quality	5 ^{139,162,164,166,189}	99.1% (94.0% to 99.9%)	72%	0.31 (quality)
Fair quality	5 ^{142,144,195,196,198}	97.0% (89.1% to 99.2%)		
Cirrhosis excluded	4 ^{139,144,189,196}	98.3% (94.4% to 99.5%)	0%	0.52 (cirrhosis)
Some cirrhosis (<20% of population)	6 ^{139,142,162,164,166,198}	99.1% (91.2% to 99.9%)	0%	
U.S. or Canada	0			
Europe / Australia / New Zealand	1 ¹⁴²	96.3% (61.1% to 99.8%)		0.67 (geographic setting)
Multinational	7 ^{139,144,164,166,189,196,198}	98.8% (94.6% to 99.7%)	45%	
Other	2 ^{162,195}	97.3% (88.0% to 99.4%)	73%	
Treatment-naïve	8139,142,144,162,164,166,189,195	98.3% (94.5% to 99.5%)	52%	
Genotype 5 infection	4 ^{139,141,143,196}	96.0% (88.3% to 98.7%)	0%	
Ledipasvir / sofosbuvir	1 ¹⁴¹	95.2% (76.2% to 99.9%)		0.99 (regimens)
Sofosbuvir / velpatasvir	1 ¹³⁹	96.6% (82.2% to 99.9%)		
Glecaprevir / pibrentasvir	2 ^{143,196}	96.0% (76.4% to 99.4%)	0%	
Good quality	2 ^{139,141}	96.0% (85.4% to 99.0%)	0%	1.00 (quality)
Fair guality	2 ^{143,196}	96.0% (76.4% to 99.4%)	0%	
Cirrhosis excluded	2 ^{139,196}	96.8% (80.4% to 99.6%)	0%	0.79 (cirrhosis)
Some cirrhosis (<20% of population)	2 ^{141,143}	95.4% (83.6% to 98.9%)	0%	
U.S. or Canada	0			
Europe / Australia / New Zealand	1 ¹⁴¹	95.2% (72.9% to 99.3%)		0.85 (geographic setting)
Multinational	3 ^{139,143,196}	96.3% (86.4% to 99.1%)	0%	

Table 12. Sustained Virologic Response Rates With Direct Acting Antiviral Regimens in Adults by Genotype

		Pooled sustained		
Analysis	Number of trials	(95% CI)	1 ²	p for interaction
Treatment-naïve	2 ^{139,141}	95.6% (83.9% to 98.9%)	0%	
Genotype 6 infection	5 ^{139,143,148,165,196}	98.2% (95.4% to 99.3%)	0%	
Ledipasvir / sofosbuvir	1 ¹⁴⁸	96.0% (79.6% to 99.9%)	%	0.37 (regimens)
Sofosbuvir / velpatasvir	2 ^{139,165}	99.2% (94.9% to 99.9%)	0%	
Glecaprevir / pibrentasvir	2 ^{143,196}	97.2% (89.4% to 99.3%)	42%	
Good quality	1 ¹³⁹	100% (90% to 100%)		<0.001 (quality)
Fair quality	4 ^{143,148,164,196}	97.9% (94.6% to 99.2%)	4%	
Cirrhosis excluded	2 ^{139,196}	97.8% (85.8% to 99.7%)	63%	0.66 (cirrhosis)
Some cirrhosis (<20% of population)	2 ^{143,164}	98.7% (95.1% to 99.7%)	0%	
Cirrhosis status unclear/not reported	1 ¹⁴⁸	96.0% (76.4% to 99.4%)		
U.S. or Canada	0			
Europe / Australia / New Zealand	1 ¹⁴⁸	96.0% (76.4% to 99.4%)		0.43 (geographic setting)
Multinational	4 ^{139,143,165,196}	98.5% (95.5% to 99.5%)	0%	
Treatment-naïve	2 ^{139,148}	98.4% (89.6% to 99.8%)	35%	
Mixed genotype [¶]	2 ^{146,150}	95.4% (89.4% to 98.1%)	0%	
Sofosbuvir / velpatasvir	2 ^{146,150}	95.4% (89.4% to 98.1%)	0%	

*Two trials reported results for genotype 1a and 1b separately (Feld 2014¹⁸⁷, Kowdley 2014b¹⁹¹).

†One trial omitted dasabuvir (Kowdley 2014b¹⁹¹).
 [‡]Two trials omitted dasabuvir (Kowdley 2014b¹⁹¹, Lawitz 2015¹⁵⁵).
 [§]One trial reported results for genotype 1a and 1b separately (Kowdley 2014b¹⁹¹).

Regimens administered with or without ribavirin.

[¶]All patients were treatment-naïve.

Abbreviations: CI = confidence interval; U.S. = United States.

Author year					
Study name	Intervention(s)	Age	Sex/Gender	Race/Ethnicity	Other characteristics
Afdahl 2014 ¹⁸⁵ ION-1	A. Ledipasvir + sofosbuvir B. Ledipasvir + sofosbuvir + ribavirin	<65 years: 99% (196/197) vs. 100% (189/189) ≥65 years: 100% (15/15) vs. 100% (22/22)	Male gender: 99% (125/126) vs. 100% (124/124) Female gender: 100% (86/86) vs. 100% (87/87)	Black: 100% (24/24) vs. 100% (26/26) Non-Black: 99.5% (187.188) vs. 100% (184/184) Hispanic: 100% (26/26) vs. 100% (19/19)	NR
Andreone 2014 ¹⁸⁶ PEARL-2	A. Ombitasvir + paritaprevir + ritonavir + dasabuvir B. Ombitasvir + paritaprevir + ritonavir + dasabuvir + ribavirin	NR	Male gender: 100% (54/54) vs. 95% (41/43) Female gender: 100% (37/37) vs. 98% (44/45)	Black: 100% (5/5) vs. 100% (3/3) Other: 100% (86/86) vs. 97% (82/85)	NR
Chuang 2016 ¹⁴⁵	Ledipasvir + sofosbuvir	<65: 100% (35/35) ≥65: 100% (7/7)	Male gender: 100% (13/13) Female gender: 100% (29/29)	NR	<u>BMI</u> <25: 100% (26/26) ≥25: 100% (16/16)
Feld 2014 ¹⁸⁷ SAPPHIRE-1	A. Ombitasvir + paritaprevir + ritonavir + dasabuvir + ribavirin B. Placebo followed by open-label ombitasvir + paritaprevir + ritonavir + dasabuvir + ribavirin	<55 years: 97% (280/290) ≥55 years: 96% (175/183)	Male gender: 95% (258/271) Female gender: 98% (197/202)	Black: 96% (27/28) Non-Black: 96% (428/445)	<u>BMI</u> <30: 97% (390/402) ≥30: 92% (65/71) <u>Diabetes</u> Yes: 100% (19/19) No: 96% (436/454)
Feld 2015 ASTRAL-1 ¹³⁹	Sofosbuvir + velpatasvir	<65 years: 99% (609/615) ≥65 years: 100% (113/113)	Male gender: 99% (426/431) Female gender: 99.7% (296/297)	Black: 98% (64/65) White: 99% (570/575) Other: 100% (84/84)	<u>BMI</u> <30: 99% (568/573) ≥30: 99% (154/155)
Foster 2015 ¹⁴⁷ ASTRAL-3	A. Sofosbuvir + velpatasvir B. Sofosbuvir + ribavirin	<65 years: 95% (257/270) vs. 81% (210/261) ≥65 years: 100% (7/7) vs. 79% (11/14)	Male gender: 94% (159/170) vs. 76% (132/175) Female gender: 98% (105/107) vs. 88% (89/101)	Black: 100% (3/3) vs. 100% (1/1) White: 95% (238/250) vs. 78% (187/239) Other: 96% (23/24) vs. 94% (32/34)	NR
Grebely 2018 ¹⁵⁰ SIMPLIFY	Sofosbuvir + velpatasvir	≤41 years: 93% (26/28) >41 years: 95% (71/75)	<u>Male gender: 92%</u> (68/74) <u>Female gender:</u> 100% (29/29)	<u>NR</u>	No current opioid substitution therapy: 93% (54/58) Current opioid substitution therapy: 96% (43/45)
Grebely 2018c ¹⁴⁹ D3FEAT	Ombitasvir + paritaprevir + ritonavir + dasabuvir + ribavirin	≤54 years: 89% (59/66) >54 years: 95% (20/21)	Male gender: 91% (61/67) Female gender: 90% (18/20)	<u>NR</u>	No current opioid substitution therapy: 96% (25/26) Current opioid substitution therapy: 89% (54/61)

Table 13. Sustained Virologic Response Rates With Direct Acting Antiviral Regimens in Adults in Subgroups

Author year					
Study name	Intervention(s)	Age	Sex/Gender	Race/Ethnicity	Other characteristics
Kowdley	Ledipasvir +	8-week	8-week	8-week intervention	NR
2014a ¹⁹⁰	sofosbuvir	intervention group	intervention group	group	
ION-3		<65 years: 94%	Male: 92%	Black: 91% (41/45)	
		(185/196)	(119/130)	Non-black: 95%	
		≥65 years: 90%	Female: 98%	(161/170)	
		(17/19)	(83/85)	Hispanic: 100%	
				(13/13)	
		<u>12-week</u>	<u>12-week</u>	Non-Hispanic: 94%	
		intervention group	intervention group	(187/200)	
		<65 years: 95%	Male gender: 95%		
		(189/199)	(122/128)	<u>12-week intervention</u>	
		≥65 years: 100%	Female gender:	group	
		(17/17)	96% (84/85)	Black: 95% (40/42)	
				NON-DIACK: 95%	
				(100/173) Hispanic: 02% (12/14)	
				Non-Hispanic: 96%	
				(103/202)	
Kowdley	A Ombitasvir +	NR	NR	Black: 100% (13/13)	NR
2014h ¹⁹¹	naritanrevir +			vs 100% (13/13)	
AVIATOR	ritonavir +			Non-black: 86%	
	dasabuvir			(57/66) vs. 96%	
	B. Ombitasvir +			(63/66)	
	paritaprevir +			()	
	ritonavir +				
	dasabuvir +				
	ribavirin				
Kumada 2017 ¹⁵²	Elbasvir +	<65 years: 99%	Male gender: 98%	NR	NR
	grazoprevir	(122/123)	(85/87)		
		65-74 years: 93%	Female gender:		
		(70/75)	96% (134/140)		
		≥75 years: 93%			
Lim 2016 ¹⁵⁶	Lodiposviru	(27/29)	ND	ND	ND
	sofosbuvir	(33/33)			
	3010300011	(00/00) >65 years: 10%			
		(13/13)			
Nelson 2015 ¹⁵⁷	Daclatasvir +	<65 vears: 90%	Male gender: 86%	NR	NR
ALLY-3	sofosbuvir	(128/142)	(77/90)		
		≥65 years: 70%	Female gender:		
		(7/10)	94% (58/62)		
Sperl 2016 ¹⁹⁸	Elbasvir +	≤40 years: 100%	Male gender:	NR	NR
C-EDGE H2H	grazoprevir	(37/37)	100% (55/55)		
		41-50 years:	Female gender:		
		100% (31/31)	99% (73/74)		
		51-60 years: 98%			
		(40/41)			
		61-70 years:			
Mai 0040-164		100% (20/20)	Mala arrista - 0001	Llienenic/Letter 1000/	
	Elbasvir +	<65 years: 95%	Male gender: 96%	Hispanic/Latino: 100%	NR
C-CORAL	grazoprevir	(420/444)	(201/210) Eomolo goodor:	Non Hispania/Lating:	
		(39/42)	93% (252/270)	94% (454/481)	
Wei 2019h ¹⁶⁵	Sofosbuvir +	<65 veare: 06%	Male gender: 0/0/	NR	NR
110120130	velpatasvir	(340/353)	(186/197)		
	. Sipataovii	≥65 years: 100%	Female gender:		
		(22/22)	99% (176/178)		

Table 13. Sustained Virologic Response Rates With Direct Acting Antiviral Regimens in Adults in Subgroups

Author yoar					
Study name	Intervention(s)	Age	Sex/Gender	Race/Ethnicity	Other characteristics
Zeuzem 2015 C-EDGE ¹⁶⁶	Grazoprevir + elbasvir	<65: 94% (270/287) ≥65: 100% (29/29)	Male gender: 93% (159/171) Female gender: 97% (140/145)	Asian: 94% (51/54) Black: 97% (57/59) White: 94% (180/191) Other: 92% (11/12)	NR
Zeuzem 2018 ¹⁶⁷ ENDURANCE-1	Glecaprevir + pibrentasvir	8-week intervention group <65 years: 99% (306/309) ≥65 years: 100% (42/42) <u>12-week</u> intervention group <65 years: 99.7% (316/317) ≥65 years: 100% (35/35)	<u>8-week</u> intervention group Male gender: 99% (165/167) Female gender: 99% (183/184) <u>12-week</u> intervention group Male gender: 100% (176/176) Female gender: 99% (175/176)	8-week intervention group Black race: 100% (14/14) Other race: 99% (334/337) <u>12-week intervention</u> group Black race: 92% (12/13) Other race: 100% (339/339)	8-week intervention group No current opioid substitution therapy: 99% (336/339) Current opioid substitution therapy: 100% (12/12) 12-week intervention group No current opioid substitution therapy: 100% (326/336) Current opioid substitution therapy: 100% (336/336) Current opioid substitution therapy: 94% (15/16)
Zeuzem 2018 ¹⁶⁷ ENDURANCE-3	A. Glecaprevir 300 mg + pibrentasvir 120 mg, 8 weeks B. Glecaprevir 300 mg + pibrentasvir 120 mg, 12 weeks 3. Sofosbuvir 400 mg + daclatasvir 60 mg, 12 weeks	Age <65 years: 95% (144/152) vs. 95% (213/224) vs. 96% (107/111) Age ≥65 years: 100% (5/5) vs. 100% (9/9) vs. 100% (4/4)	Male gender: 93% (86/92) vs. 93% (112/121) vs. 92% (48/52) Female gender: 97% (63/65) vs. 98% (110/112) vs. 100% (63/63)	Black race: 100% (3/3) vs. 100% (4/4) vs. 75% (3/4) Not Black race: 95% (146/154) vs. (218/229) vs. 97% (108/111)	No current opioid substitution therapy: 94% (119/126) vs. 96% (188/195) vs. 96% (94/98) Current opioid substitution therapy: 97% (30/31) vs. 90% (34/38) vs. 100% (17/17)

Table 13. Sustained Virologic Response Rates With Direct Acting Antiviral Regimens in Adults in Subgroups

Abbreviations: BMI = body mass index; CI = confidence interval; NR = not reported. Study names are not acronyms.

Table 14. Sustained Virologic Response Rates With Direct Acting Antiviral Regimens inAdolescents With HCV Infection

Author year Country		Antiviral treatment	SVR, total	
Quality	Population characteristics	regimen	population	SVR, subgroups
Abdel Ghaffar 2019 ²⁰¹ Egypt <i>Fair</i>	n=40 Mean age 12 years 38% female Race NR Fibrosis stage F0: 35%; F1: 38%; F2 and F3: 15% Genotype 4: 100% (mixed 4 and 1a: 13%; mixed 4 and 1b: 15%) Treatment naïve: 100%	Sofosbuvir 200 to 400 mg + daclatasvir 30 to 60 mg	98% (39/40)	NR
Balistreri 2017 ¹⁷⁵ Multinational <i>Fair</i>	n=100 Mean age 15 years 63% female 90% white; 7% black; 2% Asian; 1% NR Fibrosis stage F0-F3: 42%; F4:1%; NR/unknown: 57% Genotype 1a: 81%; 1b: 19% Treatment naïve: 80% Treatment experienced 20% (prior treatment unclear; presumably IFN or pegylated IFN + ribavirin)	Ledipasvir 90 mg + sofosbuvir 400 mg*	98% (98/100)	Treatment-naïve: 98% (78/80) Treatment- experienced: 100% (20/20)
EI-Karaksy ²⁰² 2018 Egypt <i>Fair</i>	n=40 Mean age 14 years 35% female Race NR Fibrosis stage F0: 55%; F0 and F1: 13%; F1: 13%; F1 and F2: 5%; F3: 10%; F4: 5% Genotype 4: 100% Treatment-naïve: 75%	Ledipasvir 90 mg + sofosbuvir 400 mg*	100% (40/40)	NK
Jonas 2019 ¹⁷¹ DORA Multinational <i>Fair</i>	n=48 Median age 14 years 55% female 75% white; 9% black; 13% Asian; 4% mixed race Fibrosis stage F0-F1: 96%; F2: 2%; F3: 2% Genotype 1a: 51%; 1b: 28%; 2: 6%; 3: 9%; 4: 6%; no genotype 5 or 6 enrolled HIV coinfection: 4% Treatment-naïve: 77% Treatment-experienced: 23% (pegylated IFN + ribavirin)	Glecaprevir 300 mg + pibrentasvir 120 mg	100% (47/47)	NR
Leung 2018 ²⁰³ Multinational <i>Fair</i>	n=38 Median age 15 years 66% female 76% white; 13% black; 8% Asian; 3% mixed race Fibrosis stage (30/38 patients): F0 and F1: 90%; F2: 3%; F3: 3%; F4: 3% Genotype 1a: 42%; 1b: 40%; 4: 18% Treatment naïve: 66%	Ombitasvir 25 mg + paritaprevir 150 mg + ritonavir 100 mg + dasabuvir 250 mg 2x/day + weight- based ribavirin	100% (38/38)	Genotype 1a: 100% (16/16) Genotype 1b: 100% (15/15) Genotype 4: 100% (7/7) Treatment naïve: 100% (25/25) Treatment experienced: 100% (13/13)

Table 14. Sustained Virologic Response Rates With Direct Acting Antiviral Regimens inAdolescents With HCV Infection

Author year Country		Antiviral treatment	SVR, total	
Quality	Population characteristics	regimen	population	SVR, subgroups
Wirth 2017 ¹⁷³ Multinational <i>Fair</i>	n=52 Median age 15 years 40% female 90% white; 4% black; 2% Asian; 2% Hawaiian/Pacific Islander; 2% other Fibrosis stage NR; 40% no cirrhosis; 60% cirrhosis presence unknown Genotype 2: 25% Genotype 3: 75% Treatment-naive: 83%	Sofosbuvir 400 mg + weight-based ribavirin*	98% (51/52)	Genotype 2: 100% (13/13) Genotype 3: 97% (38/39)
Yakoot 2018176	n=30	Sofosbuvir +	97% (29/30)	NR
Egypt	Mean age 13 years	daclatasvir	. ,	
Good	43% female			
	Race NR			
	Fibrosis stage F0: 17%; F1: 53%; F2:			
	27%; F3: 3%			
	Genotype 4: 100%			
	Treatment naïve: 73%			

Abbreviations: IFN = interferon; NR = not reported; SVR = sustained virologic response.

Author year	Treatment Regimen(s)	Comparison	Any adverse event	Serious adverse events*	Withdrawal due to adverse events	Headache	Fatigue	Gastrointestinal	Anemia	Insomnia	Rash
Feld 2014 ¹³⁹ SAPPHIRE-1	Ombitasvir + paritaprevir + ritonavir + dasabuvir + ribavirin	DAA vs. Placebo	86% (414/473) vs. 73% (116/158); RR 1.19 (95% CI, 1.08 to 1.32)	2% (10/473) vs. 0% (0/158); RR 7.04 (95% CI, 0.42 to 120)	0.6% (3/473) vs. 0.6% (1/158); RR 1.00 (95% Cl, 0.10 to 9.56)	33% (156/473) vs. 27% (42/158); RR 1.24 (95% Cl, 0.93 to 1.66)	35% (164/473) vs. 28% (45/158); RR 1.22 (95% CI, 0.92 to 1.60)	Nausea: 24% (112/473) vs. 13% (21/158); RR 1.78 (95% Cl, 1.16 to 2.74) Diarrhea: 14% (65/473) vs. 7% (11/158); RR 1.97 (95% Cl, 1.07 to 3.64)	NR	14% (66/473) vs. 8% (12/158); RR 1.84 (95% CI, 1.02 to 3.31)	11% (51/473) vs. (9/158); RR 1.89 (95% CI, 0.95 to 3.76)
Feld 2015 ¹³⁹ ASTRAL-1	Sofosbuvir + velpatasvir	DAA vs. Placebo	78% (485/624) vs. 77% (89/116); RR 1.01 (95% CI, 0.91 to 1.13)	2% (15/624) vs. 0% (0/116); RR 5.80 (95% CI, 0.35 to 96)	0.2% (1/624) vs. 2% (2/116); RR 0.09 (95% Cl, 0.01 to 1.02)	29% (182/624) vs. 28% (33/116); RR 1.03 (95% CI, 0.75 to 1.40)	20% (126/624) vs. 20% (23/116); RR 1.02 (95% CI, 0.68 to 1.52)	Nausea: 12% (75/624) vs. 11% (13/116); RR 1.07 (95% CI, 0.62 to 1.87) Diarrhea: 8% (48/624) vs. 7% (8/116); RR 1.12 (95% CI, 0.54 to 2.30)	NR	8% (50/624) vs. 9% (11/116); RR 0.84 (95% CI, 0.45 to 1.57	NR
Kumada 2015 ¹⁵¹ GIFT-1 (Substudy 1)	Ombitasvir + paritaprevir + ritonavir	DAA vs. Placebo	69% (148/215) vs. 57% (60/106); RR 1.22 (95% CI, 1.01 to 1.47)	3% (7/215) vs. 2% (2/106); RR 1.73 (95% CI, 0.36 to 8.16)	0.9% (2/215) vs. 0% (0/106); RR 2.48 (95% Cl, 0.12 to 51)	9% (19/215) vs. 9% (10/106); RR 0.94 (95% Cl, 0.45 to 1.94)	NR	Nausea: 4% (9/215) vs. 4% (4/106); RR 1.11 (95% CI, 0.35 to 3.52)	NR	NR	NR
Wei 2019a ¹⁶⁴ C-CORAL	Elbasvir + grazoprevir	DAA vs. Placebo	47% (230/486) vs. 50% (62/123)	2% (8/486) vs. 2% (2/123)	0.6% (3/486) vs. 2% (2/123)	6% (27/486) vs. 5% (6/123)	5% (22/486) vs. 7% (9/123)	NR	NR	NR	NR

Author year	Treatment Regimen(s)	Comparison	Any adverse event	Serious adverse events*	Withdrawal due to adverse events	Headache	Fatigue	Gastrointestinal	Anemia	Insomnia	Rash
Dore 2016 ¹³⁷ MALACHITE-1	Ombitasvir + paritaprevir mg + ritonavir + dasabuvir	DAA vs. telaprevir / pegylated interferon / ribavirin	49% (41/83) vs. 100% (37/37); RR 0.50 (95% Cl, 0.40 to 0.62)	0% (0/83) vs. 11% (4/37); RR 0.05 (95% CI, 0.003 to 0.91)	0% (0/83) vs. (3/37); RR 0.07 (95% CI, 0.003 to 1.25)	19% (16/83) 30% (11/37); RR 0.65 (95% CI, 0.33 to 1.26)	5% (4/83) vs. 30% (11/37); RR 0.16 (95% CI, 0.06 to 0.48)	Nausea: 8% (7/83) vs. 41% (15/37); RR 0.21 (95% Cl, 0.09 to 0.47)	1% (1/83) vs. 46% (17/37); RR 0.03 (95% Cl, 0.004 to 0.19)	NR	0% (0/83) vs. (8/37); RR 0.03 (95% CI, 0.002 to 0.45)
Dore 2016 ¹³⁷ MALACHITE-1	Ombitasvir + paritaprevir + ritonavir + dasabuvir + ribavirin	DAA vs. telaprevir / pegylated interferon / ribavirin	75% (115/153) 97% vs. (37/38); RR 0.77 (95% Cl, 0.69 to 0.86)	0.7% (1/153) vs. (5/38); RR 0.05 (95% Cl, 0.01 to 0.41)	1% (1/153) vs. (3/38); RR 0.08 (95% CI, 0.01 to 0.75)	27% (41/153) vs. 32% (12/38); RR 0.85 (95% Cl, 0.50 to 1.45)	14% (21/153) vs. 32% (12/38); RR 0.43 (95% CI, 0.24 to 0.80)	Nausea: 21% (32/153) vs. 39% (15/38); RR 0.53 (95% Cl, 0.32 to 0.87)	7% (10/153) vs. 45% (17/38); RR 0.15 (95% CI, 0.07 to 0.29)	NR	8% (12/153) vs. (9/38); RR 0.33 (95% Cl, 0.15 to 0.73)
Dore 2016 ¹³⁷ MALACHITE-2	Ombitasvir + paritaprevir + ritonavir + dasabuvir + ribavirin	DAA vs. telaprevir / pegylated interferon / ribavirin	62% (63/101) vs. (43/47); RR 0.68 (95% Cl, 0.57 to 0.81)	1% (1/101) vs. (2/47); RR 0.23 (95% Cl, 0.02 to 2.50)	0% (0/101) vs. 11% (5/47); RR 0.04 (95% Cl, 0.002 to 0.76)	29% (29/101) vs. 45% (21/47); RR 0.64 (95% CI, 0.41 to 1.00)	12% (12/101) vs. 25% (12/47); RR 0.47 (95% CI, 0.23 to 0.96)	Nausea: 10% (10/101) vs. 43% (20/47); RR 0.23 (95% Cl, 0.12 to 0.46)	3% (3/101) vs. 34% (16/47); RR 0.09 (95% CI, 0.03 to 0.38)	6% (6/101) vs. 21% (10/47); RR 0.28 (95% CI, 0.11 to 0.72)	3% (3/101) vs. (8/47); RR 0.17 (95% CI, 0.05 to 0.63)
Abergel 2016a ¹⁴²	Ledipasvir + sofosbuvir	NA	71% (31/44)	0%	0%	25% (11/44)	20% (9/44)	Nausea: 9% (4/44) Diarrhea: 9% (4/44)	NR	NR	NR
Abergel 2016b ¹⁴¹	Ledipasvir + sofosbuvir	NA	80% (33/41)	2% (1/41)	0%	27% (11/41)	10% (4/41)	Diarrhea: 7% (3/41)	NR	NR	NR
Afdahl 2014 ¹⁸⁵ ION-1	Ledipasvir + sofosbuvir	NA	79% (169/214)	0.5% (1/214)	0%	25% (53/214)	21% (44/214)	Nausea: 11% (24/214) Diarrhea: 11% (24/214)	0%	8% (17/214)	7% (16/214)

Author year	Treatment Regimen(s)	Comparison	Any adverse event	Serious adverse events*	Withdrawal due to adverse events	Headache	Fatigue	Gastrointestinal	Anemia	Insomnia	Rash
Ahmed 2018 ¹⁹⁵ Egypt	Ledipasvir + sofosbuvir	NA	26% (26/100)	NR	NR	2% (2/100)	18% (18/100)	Diarrhea: 1% (1/100)	NR	2% (2/100)	NR
Andreone 2014 ¹⁸⁶ PEARL II	Ombitasvir + paritaprevir + ritonavir + dasabuvir	NA	77.9% (74/95)	3% (3/95)	0%	23% (22/95)	16% (15/95)	Nausea: 6% (6/95) Diarrhea: 13% (12/95)	0% (0/95)	NR	1% (1/95)
Asselah 2018 ¹⁹⁶ SURVEYOR	Glecaprevir + pibrentasvir	NA	63% (128/203)	1% (2/203)	0%	18% (37/203)	14% (28/203)	Nausea: 11% (23/203)	NR	NR	NR
Asselah 2019 ¹⁴³ ENDURANCE 5 and 6	Glecaprevir + pibrentasvir	NA	55% (46/84)	6% (5/84)	0% (0/84)	13% (11/84)	13% (11/84)	NR	NR	NR	NR
Brown 2018 ¹⁴⁴ C-SCAPE	Elbasvir + grazoprevir	NA	79% (15/19)	0%	5% (1/19)	26% (5/19)	16% (3/19)	Nausea: 5% (1/19)	NR	NR	NR
Chayama 2018 ¹⁹⁷	Glecaprevir + pibrentasvir	NA	57% (74/129)	0%	0%	5% (6/129)	NR	NR	NR	NR	2% (3/129)
Chuang 2016 ¹⁴⁵	Ledipasvir + sofosbuvir	NA	60% (51/60)	NR	1% (1/85)	14% (12/85)	9% (8/85)	Nausea: 6% (5/85)	NR	NR	NR
Everson 2015 ¹⁴⁶ (Part A)	Sofosbuvir + velpatasvir	NA	70% (54/77)	1% (1/77)	0%	18% (14/77)	18% (14/77)	Nausea: 10% (8/77) Diarrhea: 9% (7/77)	NR	6% (5/77)	5% (4/77)
Ferenci 2014 ¹⁸⁸ PEARL III	Ombitasvir + paritaprevir + ritonavir + dasabuvir	NA	67% (140/209)	2% (4/209)	0%	23% (49/209)	23% (48/209)	Nausea: 4% (9/209)) Diarrhea: 6% (13/209)	NR	NR	3% (8/209)
Ferenci 2014 ¹⁸⁸ PEARL IV	Ombitasvir + paritaprevir + ritonavir + dasabuvir + ribavirin	NA	92% (92/100)	3.0% (3/100)	0%	25% (25/100)	46% (46/100)	Nausea: 21% (21/100) Diarrhea: 14% (14/100)	NR	NR	5% (5/100)

Author year	Treatment Regimen(s)	Comparison	Any adverse event	Serious adverse events*	Withdrawal due to adverse events	Headache	Fatigue	Gastrointestinal	Anemia	Insomnia	Rash
Foster 2015 ¹⁴⁷ ASTRAL-2 and ASTRAL-3	Sofosbuvir + velpatasvir	NA	82% (337/411)	2% (7/411)	0.2% (1/411)	28% (114/411)	22% (91/411)	Nausea: 15% (60/411)	NR	9% (37/411)	NR
Gane 2015 ¹⁴⁸	Ledipasvir + sofosbuvir	NA	92% (46/50)	10% (5/50)	2% (1/50)	24% (12/50)	22% (11/50)	Nausea: 18% (9/50) Diarrhea: 12% (6/50) Vomiting: 6% (3/50)	NR	NR	NR
Grebely 2018 ¹⁵⁰ SIMPLIFY	Sofosbuvir + velpatasvir	NA	83% (85/103)	7% (7/103)	1% (1/103)	18% (19/103)	22% (23/103)	Nausea: 14% (14/103) Vomiting: 4% (4/103) Diarrhea: 4% (4/103)	NR	9% (9/103)	NR
Grebely 2018c ¹⁴⁹ D3FEAT	Ombitasvir + paritaprevir + ritonavir + dasabuvir + ribavirin	NA	61% (53/87)	6% (5/87)	0%	5% (12/87)	10% (25/87)	Nausea: 8% (20/87) Vomiting: 4% (11/87)	5% (12/87)	4% (11/87)	NR
Hezode 2015 ¹⁸⁹ PEARL I	Ombitasvir + paritaprevir + ritonavir + ribavirin	NA	88% (80/91)	0%	0%	31% (28/91)	15% (14/91)	Nausea: 14% (13/91) Diarrhea: 14% (6/42)	NR	13% (12/91)	NR
Kowdley 2014b ¹⁹¹ AVIATOR	Ombitasvir + paritaprevir + ritonavir + dasabuvir	NA	NR	3% (2/79)	0%	19% (15/79)	20% (16/79)	Nausea: 3% (2/79) Diarrhea: 16% (13/79)	1% (1/79)	NR	NR
Author year	Treatment Regimen(s)	Comparison	Any adverse event	Serious adverse events*	Withdrawal due to adverse events	Headache	Fatique	Gastrointestinal	Anemia	Insomnia	Rash
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Kowdley 2014b ¹⁹¹ AVIATOR	Ombitasvir + paritaprevir + ritonavir + dasabuvir + ribavirin	NA	NR	1% (1/79)	3% (2/79)	27% (21/79)	28% (22/79)	Nausea: % 1% (1/79) Diarrhea: 13% (10/79)	9% (7/79)	NR	NR
Kowdley 2014a ¹⁹⁰ ION-3	Ledipasvir + sofosbuvir	NA	67% (355/431)	2% (9/431)	0.5% (2/431)	15% (63/431)	22% (94/431)	Nausea: 9% (39/431) Diarrhea: 6% (24/431)	0.7% (3/431)	6% (26/431)	1% (3/215)
Kumada 2015 ¹⁵¹	Ombitasvir + paritaprevir + ritonavir	NA	69% (148/215)	3% (7/215)	0.9% (2/215)	9% (19/215)	NR	Nausea: 4% (9/215)	NR	NR	NR
Kumada 2017 ¹⁵² (Part 2 only)	Elbasvir + grazoprevir	NA	96% (219/227)	5% (11/227)	1% (3/227)	NR	NR	NR	NR	NR	NR
Kwo 2016 ¹⁵³ OPTIMIST-1	Simeprevir + sofosbuvir	NA	66% (103/155)	1% (1/155)	0%	14% (22/155)	12% (19/155)	15% (23/155)	NR	NR	6% (10/155)
Lalezari 2015 ¹⁹²	Ombitasvir + paritaprevir + ritonavir + dasabuvir + ribavirin	NA	92% (35/38)	8% (3/38)	3% (1/38)	32% (12/38)	47% (18/38)	Nausea: 50% (19/38) Vomiting: 11% (4/38)	11% (4/38)	19% (7/38)	16% (6/38)
Lawitz 2014a ¹⁵⁴ COSMOS	Simeprevir + sofosbuvir	NA	79% (11/14)	0%	0%	NR	NR	NR	0%	NR	7% (1/14)
Lawitz 2014b ¹⁹³ LONESTAR	Ledipasvir + sofosbuvir	NA	45% (17/39)	3% (1/39)	0%	5% (2/39)	NR	Nausea: 8% (3/39)	NR	NR	3% (1/39)
Lawitz 2015 ¹⁵⁵ PEARL 1	Ombitasvir + paritaprevir + ritonavir	NA	93% (76/82)	2% (2/82)	0%	29% (24/82)	7% (6/82)	Nausea: 10% (8/82) Diarrhea: 7% (6/82)	NR	NR	NR
Lim 2016 ¹⁵⁶	Ledipasvir + sofosbuvir	NA	49% (46/93)	3% (3/93)	1% (1/93)	8% (7/93)	8% (7/93)	ŇR	NR	NR	NR

Author year	Treatment Regimen(s)	Comparison	Any adverse event	Serious adverse events*	Withdrawal due to adverse events	Headache	Fatigue	Gastrointestinal	Anemia	Insomnia	Rash
Nelson 2015 ¹⁵⁷ ALLY-3	Sofosbuvir + daclatasvir	NA	NR	0.7% (1/152)	NR	20% (30/152)	19% (29/152)	Nausea: 12% (18/152)	NR	6% (9/152)	NR
								Diarrhea: 9% (13/152)			
Poordad 2017 ¹⁹⁴ MAGELLAN-1	Glecapravir + pibrentasvir	NA	82% (23/28)	4% (1/28)	0%	32% (9/28)	18% (5/28)	Nausea: 18% (5/28)	NR	0%	NR
Pott-Junior 2019 ¹⁵⁹ Group A	Sofosbuvir + daclatasvir	NA	NR	NR	NR	15% (10/65)	23% (15/65)	Nausea: 6% (4/65)	NR	6% (4/65)	2% (1/65)
								Vomiting: 2% (1/65)			
Pott-Junior 2019 ¹⁵⁹ Group B	Simeprevir + sofosbuvir	NA	NR	NR	NR	28% (17/60)	28% (17/60)	Nausea: 13% (8/60)	NR	10% (6/60)	10% (6/60)
								Vomiting: 5% (3/60)			
Sperl 2016 ¹⁹⁸ C-EDGE Head- 2-Head	Elbasvir + grazoprevir	NA	52% (67/129)	0.8% (1/129)	0%	NR	NR	NR	NR	NR	NR
Sulkowski 2014 ¹⁶¹	Sofosbuvir + daclatasvir	NA	93% (38/41)	2% (1/41)	0%	34% (14/41)	39% (16/41)	Nausea: 20% (8/41)	NR	10% (4/41)	NR
Study								Vomiting: 2% (1/41)			
								Diarrhea: 5% (2/41)			
Sulkowski 2015 ¹⁶⁰ C-WORTHY	Elbasvir + grazoprevir	NA	56% (24/43; drug-related adverse events)	0%	0%	35% (15/43)	23% (10/43)	Nausea: 16% (7/43) Diarrhea: 12% (5/43)	NR	NR	NR
Toyoda 2018 ¹⁹⁹ CERTAIN-2	Glecaprevir + pibrentasvir	NA	48% (43/90)	2% (2/90)	1% (1/90)	7% (6/90)	NR	Nausea: 3% (3/90)	0%	NR	NR

Author year	Treatment Regimen(s)	Comparison	Any adverse event	Serious adverse events*	Withdrawal due to adverse events	Headache	Fatigue	Gastrointestinal	Anemia	Insomnia	Rash
Waked 2016 ¹⁶² AGATE-II	Ombitasvir + paritaprevir + ritonavir + ribavirin	NA	80% (80/100)	2% (2/100)	0%	41% (41/100)	35% (35/100)	Dyspepsia: 17% (17/100)	NR	9% (9/100)	NR
Wei 2018 ¹⁶³	Ledipasvir + sofosbuvir	NA	58% (120/206)	1% (3/206)	0%	NR	NR	NR	NR	NR	NR
Wei 2019b ¹⁶⁵	Sofosbuvir + velpatasvir	NA	50% (189/375)	1% (3/375)	0%	5% (18/375)	NR	NR	NR	NR	NR
Zeuzem 2015 ¹⁶⁶ C-EDGE	Elbasvir + grazoprevir	NA	71% (175/246)	3% (7/246)	0.8% (2/246)	NR	NR	NR	NR	NR	NR
Zeuzem 2018 ¹⁶⁷ ENDURANCE-1	Glecaprevir + pibrentasvir	NA	64% (450/703)	1% (9/703)	0.1% (1/703)	18% (130/703)	11% (74/703)	Nausea: 7% (48/703)	NR	NR	NR
Zeuzem 2018 ¹⁶⁷ ENDURANCE-3 (Glecaprevir + pibrentasvir arm)	Glecaprevir + pibrentasvir	NA	71% (275/390)	2% (7/390)	0.8% (3/390)	23% (91/390)	16% (64/390)	Nausea: 13% (51/390)	NR	NR	NR
Zeuzem 2018 ¹⁶⁷ ENDURANCE-3 (Sofosbuvir + daclatasvir arm)	Sofosbuvir + daclatasvir	NA	70% (80/115)	2% (2/115)	0.9% (1/115)	20% (23/115)	14% (16/115)	Nausea: 13% (15/115)	NR	NR	NR

*Serious adverse events listed in Appendix B Table 12

Abbreviations: CI = confidence interval; DAA = direct acting antiviral; NA = not applicable; NR = not reported; RR = relative risk. Study names are not acronyms.

Table 16. Pooled Rates With Direct Acting Antiviral Regimens in Adults for Any Adverse Event, Serious Adverse Events, and Withdrawals Due to Adverse Events

			Withdrawal due to adverse event:
	Any adverse event: Pooled rate	Serious adverse events: Pooled rate	Pooled rate (95% CI): I ² ; number of
Analysis	(95% CI); I ² ; number of studies (k)	(95% CI); I ² ; number of studies (k)	studies (k)
All studies	73.3% (68.0% to 78.1%); I ² =95%; k=44 ^{137,139,141-156,159-167,185-190,192-199}	1.9% (1.5% to 2.4%); $I^2=31\%$; k=42 ^{137,139,141-144,146-157,160-167,185-194,196-199}	0.4% (0.3% to 0.6%); $I^2=0\%$; k=41 ^{137,139,141-156,160,161,163-167,185-194,196-199}
Ledipasvir / sofosbuvir	$\begin{array}{l} 69.4\% \ (54.8\% \ to \ 80.9\%); \ l^2 = 95\%; \\ k = 10^{141,142,145,148,156,163,185,190,193,195} \end{array}$	2.0% (1.0% to 3.9%); l ² =47%; k=8 ^{141,142,148,156,163,185,190,193}	0.4% (0.2% to 1.0%); l ² =0%; k=9 ^{141,142,145,148,156,163,185,190,193}
Simeprevir / sofosbuvir	67.5% (60.0% to 74.1%); l ² =0%; k=2 ^{153,156}	0.6% (0.1% to 4.1%); l ² =0%; k=2 ^{153,156}	0% (0% to 21.6%); k=2* ^{153,156}
Sofosbuvir / daclatasvir	82.7% (58.5% to 94.2%); l ² =90%; k=2 ^{161,167}	1.3% (0.5% to 3.4%); l ² =0%; k=3 ^{157,161,167}	0.6% (0.1% to 4.4%); l ² =0%; k=2 ^{161,167}
Sofosbuvir / velpatasvir	74.6% (63.5% to 83.2%); l ² =96%; k=6 ^{139,146,147,150,165}	1.9% (0.1% to 4.1%); l ² =57%; k=6 ^{139,146,147,150,165}	0.2% (0.1% to 0.6%); l ² =0%; k=6 ^{139,146,147,150,165}
Glecaprevir / pibrentasvir	62.3% (56.1% to 68.1%); l ² =78%; k=7 ^{143,167,194,196,197,199}	1.7% (1.1% to 2.7%); l ² =44%; k=7 ^{143,167,194,196,197,199}	0.3% (0.1% to 0.9%); l ² =0%; k=7 ^{143,167,194,196,197,199}
Elbasvir / grazoprevir	79.1% (50.0% to 86.8%); l ² =98%; k=6 ^{144,152,160,164,166,198}	2.1% (1.1% to 3.9%); l ² =42%; k=6 ^{144,152,160,164,166,198}	0.9% (0.5% to 1.6%); $l^2=0\%$; k=6 ^{144,152,160,164,166,198}
Ombitasvir / paritaprevir / ritonavir / dasabuvir	75.1% (62.3% to 84.6%); l ² =92%; k=6 ^{137,151,155,186,188}	1.9% (1.2% to 3.2%); l ² =31%; k=7 ^{137,151,155,186,188,191}	0.1% (0% to 4.0%); l ² =0%; k=7 ^{137,151,155,186,188,191}
Ombitasvir / paritaprevir / ritonavir / dasabuvir/ ribavirin	81.1% (74.2% to 86.5%); l ² =87%; k=10 ^{137,149,162,186-189,192}	2.1% (1.5% to 3.0%); l ² =26%; k=11 ^{137,149,162,186-189,191,192}	0.6% (0.3% to 1.1%); l ² =11%; k=11 ^{137,149,162,186-189,191,192}
Patients with cirrhosis excluded	75.5% (69.0% to 81.1%); I ² =94%; k=24 ^{137,144,146,151-155,160,167,186-190,192-} 194,196,197,199	1.8% (1.3% to 2.5%); I ² =19%; k=24 ^{144,146,151-154,160,167,186-194,196,197,199}	0.5% (0.3% to 0.7%); I ² =14%; k=23 ^{144,146,151-155,160,167,186-194,196,197,199}
Some patients (<20% of sample) with cirrhosis	72.4% (64.6% to 79.0%); I ² =95%; k=19 ^{139,141-143,145,147-150,156,161-166,185,198}	2.0% (1.4% to 2.7%); I ² =51%; k=21 ^{137,139,141-143,147-150,156,157,161-166,185,198}	0.3% (0.2% to 0.6%); I ² =0%; k=21 ^{137,139,141,143,145,147-150,156,161,162,164-} 166,185,198
Treatment-naïve	74.0% (66.6% to 80.2%); I ² =95%; k=23 ^{137,141,142,144-146,148,149,156,160- 162,164,166,167,185,187-190,193,195}	$\begin{array}{l} 1.8\% \ (1.4\% \ to \ 2.4\%); \ l^2 = 16\%; \\ k = 24^{137,141,142,144,146,148,149,156,157,160-} \\ {}_{162,164,166,167,185,187-191,193} \end{array}$	0.5% (0.3% to 0.8%); I ² =0%; k=23 ^{137,141,142,144-146,148,149,156,160-} 162,164,166,167,185,187-191,193
Treatment-experienced	76.6% (61.5% to 87.0%); l ² =72%; k=5 ^{137,154,186,189,194}	1.7% (0.7% to 4.0%); l ² =0%; k=5 ^{137,154,186,189,194}	0.5% (0.1% to 2.1%); l ² =0%; k=5 ^{*137,154,186,189,194}
Mixed treatment-naïve and experienced	71.0% (62.0% to 78.6%); $l^2=93.4\%$; k=16 ^{139,143,147,148,151-153,155,163,165,192,196-199}	1.9% (1.4% to 2.6%); $l^2=51\%$; k=17 ^{139,143,147,148,151-153,155,163,165,192,196-199}	0.3% (0.2% to 0.5%); l ² =9%; k=17 ^{139,143,147,148,151-} 153,155,163,165,167,192,196-199

*No events reported

Abbreviation: CI = confidence interval.

	Anemia: Pooled rate (95%			
Analysis	CI); I ² ; number of studies (k)	Fatigue: Pooled rate (95% CI); I ² ; number of studies (k)	Headache: Pooled rate (95% CI): I ² ; number of studies (k)	Insomnia: Pooled rate (95% CI); I ² ; number of studies (k)
All studies	2.4% (0.9% to 6.3%); l ² =85%:	18.4% (15.6% to 21.7%); $l^2=90\%$; k=37 ^{137,139,141-150,153,155-157,159-}	18.7% (15.6% to 22.2%); I ² =90%; k=42 ^{137,139,141-151,153,155-157,159-}	8.3% (6.8% to 10.1%); l ² =58%; k=18 ^{139,146,147,149,150,157,159-}
	k=13 ^{137,149,154,185,186,190-} 192,199	162,164,167,185-192,194-196	162,164,165,167,185-197,199	162,185,187,189,190,192,194,195
Ledipasvir / sofosbuvir	0.5% (0.2% to 1.4%); l ² =44%; k=2 ^{185,190}	16.2% (12.2% to 21.0%); l ² =67%; k=8 ^{141,142,145,148,156,185,190,195}	13.7% (8.4% to 21.5%); l ² =85%; k=9 ^{141,142,145,148,156,185,190,193,195}	6.0% (4.5% to 8.0%); l ² =58%; k=3 ^{185,190,195}
Simeprevir / sofosbuvir	0% (0% to 23.2%); k=1* ¹⁵⁴	18.4% (9.8% to 31.8%; l ² =86%); k=2 ^{153,159}	19.5% (11.7% to 30.8%; l ² =81%); k=2 ^{153,159}	10.0% (3.8% to 20.5%); k=1 ¹⁵⁹
Sofosbuvir / velpatasvir		20.8% (17.9% to 24.0%); l ² =44%; k=5 ^{139,146,147,150}	18.0% (10.8% to 28.5%); l ² =96%; k=6 ^{139,146,147,150,165}	8.3% (6.7% to 10.2%); l ² =32%; k=5 ^{139,146,147,150}
Sofosbuvir / daclatasvir		21.7% (14.9% to 30.1%); $l^2=72\%$; k=4 ^{157,159,160,167}	20.6% (16.8% to 25.1%); I ² =41%; k=4 ^{157,159,161,167}	6.4% (4.0% to 10.1%); l ² =0%; k=4 ^{157,159,161,194}
Glecaprevir / pibrentasvir	0% (0% to 4.0%); k=1 ¹⁹⁹	13.3% (10.8% to 16.3%; $l^2=54\%$); k=5 ^{143,167,194,196}	14.7% (9.4% to 22.2%) l ² =87%; k=7 ^{143,167,194,196,197,199}	0% (0% to 15.4%); k=1 ¹⁹⁴
 Elbasvir / grazoprevir 		10.9% (4.3% to 25.1%; l ² =88%); k=3 ^{144,160,164}	17.1% (6.1% to 39.5%) l ² =94%; k=3 ^{144,160,164}	7.0% (2.3% to 19.5%); k=1 ¹⁶⁰
Ombitasvir / paritaprevir / ritonavir / dasabuvir	0.8% (0.2% to 3.1%); l ² =0%%; k=3 ^{137,186,191}	15.8% (9.1% to 26.1%); 91%; k=6 ^{137,155,186,188,191}	20.7% (15.6% to 26.9%); l ² =83%; k=7 ^{137,151,155,186,188,191}	
Ombitasvir / paritaprevir / ritonavir / dasabuvir / ribavirin	8.3% (5.8% to 11.8%); I ² =49%; k=6 ^{137,149,186,191,192}	26.9% (20.5% to 34.4%); l ² =88%; k=11 ^{137,149,162,186-189,191,192}	27.6% (24.0% to 31.5%); l ² =61%; k=11 ^{137,149,162,187-189,191,192}	13.3% (11.1% to 15.9%); l ² =0%; k=5 ^{149,162,187,189,192}
Patients with cirrhosis excluded	2.1% (0.6% to 7.3%); l ² =81%; k=6 ^{154,186,190-192,199}	20.2% (16.0% to 25.3%); I ² =92%; k=18 ^{144,146,153,155,159,160,167,186- 192,194,196}	19.6% (15.5% to 24.3%); $l^2=87\%$; k=22 ^{144,146,151,153,155,159,160,167,186-194,196,197,199}	9.0% (7.0% to 11.5%); l ² =68%; k=10 ^{146,157,159,161,162,187,189,190,192,194}
 Some cirrhosis (≤20%) 	2.9% (0.7% to 11.0%); l ² =92%; k=4 ^{137,149,185}	16.7% (13.1% to 21.2%); l ² =90%; k=18 ^{137,139,141-143,145,147-} 150,156,157,161,162,164,185	19.1% (14.9% to 24.1%); l ² =94%; k=19 ^{137,139,141-143,145,147} 150,156,157,161,162,164,165,185	8.4% (6.4% to 10.9%); l ² =12%; k=8 ^{139,147,149,150,160,185,194}
Treatment-naïve	2.2% (0.7% to 6.7%); I ² =90%; k=6 ^{137,149,185,190,191}	18.1% (14.5% to 22.2%); $l^2=92\%$; k=24 ^{141,142,144-146,148,149,156,157,160-162,164,167,185,187-191,195}	21.1% (16.8% to 26.1%); $l^2=91\%$; k=24 ^{137,141,142,144-} 146,148,149,156,157,160,162,164,167,185,187- 191,193,195	7.9% (5.9% to 10.7%); I ² =68%; k=10 ^{146,149,160,162,185,187,189,190,194,195}
Treatment- experienced	3.6% (0.8% to 14.5%); l ² =0%; k=3 ^{137,154,186}	23.2% (14.7% to 34.6%); l ² =51%; k=4 ^{137,186,189,194}	23.5% (14.4% to 35.8%); l ² =0%; k=4 ^{137,186,189,194}	10.0% (5.2% to 18.5%); I ² =68%; k=3 ^{161,189,194}
 Mixed treatment- naïve and experienced 	2.1% (0.2% to 18.1%); l ² =89%; k=2 ^{192,199}	17.6% (12.8% to 23.7%) $l^2=87\%$; k=11 ^{139,143,147,148,153,155,159,164,192,196}	14.5% (10.6% to 19.5%); l ² =93%; k=15 ^{139,143,147,148,151,153,155,159,167,192,} 196,197,199	8.3% (5.9% to 11.4%); l ² =53%; k=6 ^{139,147,157,159,192}

Abbreviation: CI = confidence interval.

				Vomiting: Pooled rate	
_		Nausea: Pooled rate (95% CI); I ² ;	Diarrhea: Pooled rate (95% CI);	(95% CI); I ² ; number of	Rash: Pooled rate (95% CI):
Ana	alysis	number of studies (k)	I ² ; number of studies (k)	studies (k)	I ² ; number of studies (k)
All s	studies	11.1% (9.1% to 13.5%); l ² =82%;	8.7% (6.9% to 11.0%); l ² =70%;	5.8% (3.4% to 9.7%);	5.4% (4.1% to 7.1%); l ² =70%,
		k=36 ^{137,139,142,144-151,153,157,159-}	k=18 ^{141,142,146,148,150,155,157,160,161,185-}	$l^2=43\%$; k=6 ^{148-150,159,161,192}	k=17 ^{137,146,153,154,158-160,185-}
		162,167,185,186,188-196,199	191,195		188,190,192,193,197
•	Ledipasvir / sofosbuvir	8.4% (5.7% to 12.1%); l ² =60%; k=7 ^{142,145,148,185,190,193,195}	6.8% (4.2% to 10.9%); I ² =72%; k=6 ^{141,142,148,185,190,195}	6.0% (1.9% to 17.0%); k=1 ¹⁴⁸	3.3% (1.8% to 8.8%); l ² =80%; k=3 ^{185,190,193}
•	Simeprevir / sofosbuvir	14.4% (10.3% to 19.8%); I ² =0%; k=2 ^{153,159}		5.0% (1.6% to 14.4%); k=1 ¹⁵⁹	7.4% (4.7% to 11.6%); l ² =0%; k=3 ^{153,154,159}
•	Sofosbuvir / daclatasvir	12.1% (9.1% to 15.8%); l ² =32%; k=4 ^{157,159,161,167}	7.8% (4.7% to 12.5%); l ² =0%; k=2 ^{157,161}	1.9% (0.5% to 7.2%); I ² =0%; k=2 ^{159,161}	1.5% (0.2% to 10.1%); k=1 ¹⁵⁹
•	Sofosbuvir / velpatasvir	12.9% (11.0% to 15.0%); l ² =13%; k=5 ^{139,146,147,150}	6.1% (3.4% to 10.8%); l ² =50%; k=2 ^{146,150}	3.9% (1.5% to 9.9%); k=1 ¹⁵⁰	8.3% (4.9% to 13.7%); l ² =45%; k=2 ^{146,158}
•	Glecaprevir / pibrentasvir	9.3% (6.4% to 13.4%); I ² =79%; k=5 ^{167,194,196,199}			2.3% (0.5% to 6.6%); k=1 ¹⁹⁷
•	Elbasvir / grazoprevir	12.9% (6.6% to 23.7%); l ² =19%; k=2 ^{144,160}	11.6% (4.9% to 25.0%); k=1 ¹⁶⁰		4.7% (1.2% to 16.8%); k=1 ¹⁶⁰
•	Ombitasvir / paritaprevir / ritonavir / dasabuvir	6.5% (4.3% to 9.7%); l ² =70%; k=7 ^{137,151,155,186,188,191}	11.1% (7.7% to 15.9%); l ² =72%; k=5 ^{155,186,188,191}		2.6% (1.0% to 6.7%); l ² =66%; k=4 ^{137,186,188}
•	Ombitasvir / paritaprevir / ritonavir / dasabuvir/ ribavirin	15.2% (9.6% to 23.2%); l ² =90%; k=11 ^{137,149,162,186-189,191,192}	10.9% (7.8% to 14.9%); l ² =73%; k=6 ^{186-189,191}	12.0% (7.4% to 18.9%); l ² =0%; k=2 ^{149,192}	7.6% (5.5% to 10.3%); l ² =57%; k=7 ^{137,186-188,192}
•	Patients with cirrhosis excluded	10.6% (8.2% to 13.5%); $l^2=89\%$; k=21 ^{144,146,151,153,155,160,167,186-194,196,199}	10.1% (7.9% to 12.8%); l ² =80%; k=10 ^{146,155,160,186-191}	5.2% (2.1% to 12.4%); l ² =65%; k=2 ^{159,192}	5.2% (3.8% to 7.1%); l ² =69%; k=13 ^{146,153,154,159,160,186-} 188,190,192,193,197
•	Some patients (<20% of sample) with cirrhosis	12.9% (9.6% to 17.1%); I ² =43%; k=14 ^{137,139,142,145,147-150,157,161,162,185}	8.0% (5.5% to 11.6%); l ² =8%; k=7 ^{141,142,148,150,157,161,185}	6.1% (3.2% to 11.4%); l ² =51%; k=4 ^{148-150,161}	6.2% (3.7% to 10.1%); l ² =49%; k=4 ^{137,158,185}
•	Treatment-naïve	11.8% (9.0% to 15.2%); $l^2=86\%$; k=22 ^{137,142,144-146,148,149,157,160-162,167,185,187-191,193,195}	8.9% (6.9% to 11.4%); l ² =77%; k=15 ^{141,142,146,148,157,160,161,185,187-} 191,195	9.6% (5.3% to 16.9%); l ² =51%; k=3 ^{148,149,161}	5.2% (3.6% to 7.3%); $I^2=74\%$; k=9 ^{137,146,154,160,185,187,188,190}
•	Treatment-experienced	12.2% (7.2% to 20.1%); l ² =0%; k=4 ^{137,186,189,194}	10.0% (5.0% to 18.9%); l ² =0%; k=2 ^{186,189}		4.8% (2.8% to 8.2%); l ² =50%; k=5 ^{137,154,158,186,197}
•	Mixed treatment-naïve	9.6% (6.6% to 13.6%); I ² =87%;	10.1% (4.5% to 21.1%); I ² =39%;	4.3% (2.1% to 8.6%);	7.6% (4.2% to 13.6%);
	and experienced	k=12 ^{139,147,148,151,153,155,159,167,192,196,199}	k=2 ^{148,155}	I ² =54%; k=3 ^{148,159,192}	I ² =47%; k=3 ^{153,159,192}

Abbreviation: CI = confidence interval.

Table 19. Adverse Events With Direct Acting Antiviral Regimens in Adolescents

Author, year				Withdrawal due				
Country	Antiviral treatment	Any adverse	Serious	to adverse				
Quality	regimen	event	adverse events	events	Headache	Fatigue	Gastrointestinal	Insomnia
Abdel Ghaffar 2019 ²⁰¹ Eqypt	Sofosbuvir 200-400 mg + daclatasvir 30- 60 mg	NR	NR	NR	3% (1/40)	5% (2/40)	Vomiting: 3% (1/40)	NR
Fair	oo mg							
Balistreri 2017 ¹⁷⁵ Multinational <i>Fair</i>	Ledipasvir 90 mg + sofosbuvir 400 mg*	71% (71/100)	0% (0/100)	0% (0/100)	27% (27/100)	13% (13/100)	Nausea: 11% (11/100) Vomiting: 11% (11/100)	NR
El-Karaksy 2018 ²⁰² Egypt <i>Fair</i>	Ledipasvir 90 mg + sofosbuvir 400 mg*	NR	NR	NR	48% (19/40)	53% (21/40)	Nausea: 28% (11/40) Diarrhea: 23% (9/40)	23% (9/40)
Leung 2018 ²⁰³ Multinational <i>Fair</i>	Ombitasvir 25 mg + paritaprevir 150 mg + ritonavir 100 mg + dasabuvir 250 mg 2x/day + weight- based ribavirin	84% (32/38)	0% (0/38)	0% (0/38)	21% (8/38)	18% (7/38)	NR	NR
Wirth 2017 ¹⁷³ Multinational <i>Fair</i>	Sofosbuvir 400 mg + weight-based ribavirin*	81% (41/52)	2% (1/52)	0% (0/52)	23% (12/52)	12% (6/52)	Nausea: 27% (14/52) Diarrhea: 6% (3/52)	NR
Yakoot 2018 ¹⁷⁶ Egypt <i>Good</i>	Sofosbuvir + daclatasvir	27% (8/30)	0% (0/30)	0% (0/30)	10% (3/30)	13% (4/30)	Nausea: 10% (3/30)	NR

*Currently recommended regimen.

Abbreviation: NR = not reported.

Table 20. Studies on the Association Between Sustained Virologic Response After AntiviralTherapy Versus No Sustained Virologic Response and Clinical Outcomes

Author year	Duration of	N, by treatment	Percent with Cirrhosis	Percent with Genotype 1	Statistical adjustments for
Arase 2007 ²⁰⁴	7 4 years	SVR· 140	14%	60%	Yes
Japan*	(mean)	No SVR: 360	11/0	0070	
Asahina 2010 ²¹⁷	7.5 years	SVR: 686	5%	70%	Yes
Japan [†]	(mean)	No SVR: 1.356	0,0	1070	
Backus 2011 ⁶⁹	3.8 years	SVR: 7.461	13%	72%	Yes
U.S.‡	(median)	No SVR: 9.403		/ •	
Butt 2017 ²⁰⁵	1.5 years	SVR: 6.371	15%	85%	Yes
U.S.‡		No SVR: 599			
Carrat 2019 ¹⁶⁸	2.8 years	SVR: 3,286	0% (subgroup)	67%	Yes
	(median)	No SVR: 146			
		Unknown SVR:			
		1,089			
Cozen 2013 ²⁰⁶	10 years	SVR: 112	5%	67%	Yes in San Francisco VA cohort
U.S.‡	(mean)	No SVR: 91			Partial in UCSF cohort
		Relapse: 43			
		Early treatment			
		discontinuation or			
Diaparink 2014207	7.5.10000		210/	709/	Vaa
	(median)	SVR. 222 No SVR: 314	2170	70%	res
Dobmen 2013 ²¹⁸		SV/R· 285	NR	67%	Partial
Japan	(median)	No SVR: 189		07 /0	
El-Serag 2014 ²¹⁵	5 2 years	SVR: 7 577	NR	55%	Yes
U.S.‡	(mean)	No SVR: 8.767		0070	
Ikeda 1999 ²¹⁹	5.4 years	SVR: 145	0%	67%	Yes
Japan*	(median)	No SVR: 585	• / •	01 /0	
Imai 1999 ²²⁰	4 years	SVR: 151	8%	NR	Partial
Japan	(median)	Relapse: 120			
		No SVR: 148			
Imazeki 2003 ²⁰⁸	8.2 years	SVR: 116	13%	74%	Partial
Japan [§]	(mean)	No SVR: 239			
Innes 2011 ²⁰⁹	5.3 years	SVR: 560	14%	35%	Yes
U.K.	(mean)	No SVR: 655			
loannou 2018 ²²¹	6.1 years	SVR: 28,655	17%	77%	Yes
U.S.	(mean)	NO SVR: 23,231	4.0/	500/	
IZUMI 2005	Unclear	SVR: 155	1%	50%	Unclear
Japan Kasabara 1009223	2 1 voore	NU SVR. 340	0%	590/	Voc
lanan [¶]	(mean)	Delanse: 304	0%	30%	res
Japan	(mean)	No SV/R: 405			
Kasahara 2004 ²¹⁰	5 7 vears	SV/R· 738	4%	NR	Partial
Japan [¶]	(mean)	No SVR: 1.930	170		
Kurokawa	3 vears	SVR: 139	2%	89%	Partial
2009 ²²⁴	(median)	No SVR: 264			
Japan [¶]	· /				
Lee 2017 ²²⁵	2.6 years	SVR: 306	13%	51%	Yes
South Korea	(median)	No SVR: 183			
Maruoka 2012 ²¹¹	9.9 years	SVR: 221	10%	73%	Yes
Japan [§]	(mean)	No SVR: 356			
Okanoue 2002 ²²⁶	5.6 years	SVR: 426	4%	NR	Partial
Japan	(mean)	Relapse: 358			
0		No SVR: 426			
Osaki 2012 ²²⁷	4.1 years	SVR: 185	0%	60%	Partial
Japan	(median)	NO SVR: 197	040/	000/	¥
Singai 2013-12	5 years	5VK: 83	21%	68%	Yes
U.J. Sinn 2009231	(median)	NU SVK: 159	Ungloor	469/	No
South Korea	4.0 years	3VR. 290	Unclear	40%	INU
South Noted	(mealail)	130 0 911. 134			

 Table 20. Studies on the Association Between Sustained Virologic Response After Antiviral

 Therapy Versus No Sustained Virologic Response and Clinical Outcomes

Author year Country	Duration of followup	N, by treatment response	Percent with Cirrhosis	Percent with Genotype 1	Statistical adjustments for age, sex, fibrosis, genotype
Tanaka 2000 ²²⁸	4.8 years	SVR: 175	3%	75%	Yes
Japan	(mean)	Relapse: 165 No SVR: 254			
Tateyama 2011 ²²⁹	8.2 years (mean)	SVR: 139 No SVR: 234	17%	72%	Yes
Japan					
Tseng 2016 ²¹⁶ Taiwan	5.5 years (mean)	SVR: 95 No SVR: 50	NR	61%	Partial
Yoshida 1999 ²³⁰ Japan [#]	4.3 years (mean)	SVR: 789 No SVR: 1,568	10%	70%	Partial
Yoshida 2002 ²¹³ Japan [#]	5.4 years (mean)	SVR: 817 No SVR: 1,613	10%	NR	Partial
Yu 2006 ²¹⁴	5.2 years	SVR: 715	16%	46%	Yes
Taiwan	(mean)	No SVR: 342			

* Study populations overlap.

† Study populations overlap.

‡ Study population appears to overlap with Ioannou 2018.

§ Study populations overlap.

Study population appears to overlap with Backus 2011, Butt 2017, Cozen 2013, Dieperink 2014, and El-Serag 2014.

¶ Study populations likely overlap.

Study populations appear to overlap.

Abbreviations: NR = not reported; SVR = sustained virologic response; UCSF = University of California, San Francisco; U.K. = United Kingdom; U.S. = United States; VA = Veterans Affairs.

Table 21. Pooled Estimates on the Association Between Sustained Virologic Response After Antiviral Therapy Versus No Sustained Virologic Response and Clinical Outcomes

		Adjusted HR (95%			p for
Outc	ome	CI)	²	Number of studies	interaction
All-c	ause mortality	0.40 (0.28 to 0.56)	52%	13 ^{69,168,204-214}	
•	Exclude overlapping	0.37 (0.25 to 0.56)	62%	10 ^{69,168,204,205,209-214}	
	studies				
•	Fully adjusted*	0.42 (0.29 to 0.62)	55%	10 ^{69,168,204-207,209,211,212,214}	0.34
•	Partially adjusted	0.29 (0.15 to 0.55)	0%	3 ^{208,210,213}	
٠	Duration >5 years	0.33 (0.24 to 0.46)	0%	9 ^{204,206-211,213,214}	0.003
•	Duration <5 years	0.64 (0.56 to 0.74)	58%	4 ^{69,168,205,212}	
٠	U.S./Europe	0.48 (0.30 to 0.79)	54%	7 ^{69,168,205-207,209,212}	0.10
•	Asia	0.29 (0.19 to 0.45)	0%	6 ^{204,208,210,211,213,214}	
٠	Cirrhosis 0-10%	0.33 (0.18 to 0.60)	0%	4 ^{168,206,211,213}	0.58
•	Cirrhosis >10%	0.41 (0.28 to 0.62)	56%	9 ^{69,204,205,207-210,212,214}	
Live	r mortality [†]	0.11 (0.04 to 0.27)	0%	4 ^{204,208,210,213}	
٠	Fully adjusted*	0.13 (0.03 to 0.59)		1 ²⁰⁴	0.79
•	Partially adjusted	0.10 (0.03 to 0.30)	0%	3 ^{208,210,213}	
٠	Cirrhosis 0-10%	0.13 (0.03 to 0.61)		1 ²¹³	0.82
٠	Cirrhosis >10%	0.10 (0.03 to 0.30)	0%	3 ^{204,208,210}	
Cirrh	iosis‡	0.36 (0.33 to 0.40)	0%	4 ^{206,215,216}	
•	Exclude overlapping	0.36 (0.33 to 0.40)	0%	3 ^{206,215,216}	
	studies				
•	Fully adjusted*	0.36 (0.33 to 0.40)	0%	2 ^{206,215}	0.76
•	Partially adjusted	0.31 (0.12 to 0.78)	0%	2 ^{206,216}	
•	U.S./Europe	0.36 (0.33 to 0.40)	0%	3 ^{206,215}	0.71
•	Asia	0.29 (0.10 to 0.76)		1 ²¹⁶	
٠	Cirrhosis 0 to 10%	0.36 (0.13 to 1.03)	0%	2 ²⁰⁶	0.99
٠	Cirrhosis unclear	0.36 (0.33 to 0.50)	0%	2 ^{215,216}	
Нера	atocellular carcinoma	0.29 (0.23 to 0.38)	19%	20168,204,207,211,214,215,217-230	
٠	Exclude overlapping	0.25 (0.19 to 0.35)	34%	16 ^{168,204,211,214,217,218,220,221,223-230}	
	studies				
•	Fully adjusted*	0.30 (0.27 to 0.34)	0%	13 ^{168,204,207,211,214,215,217,219,221,223,225,228,229}	0.26
•	Partially adjusted	0.26 (0.16 to 0.42)	51%	7 ^{218,220,222,224,226,227,230}	
٠	Duration >5 years	0.30 (0.27 to 0.34)	23%	10 ^{204,207,211,214,215,217,221,226,229}	0.18
•	Duration <5 years	0.29 (0.16 to 0.52)	17%	9 ^{168,218,220,223-225,227,228,230}	
٠	U.S./Europe	0.32 (0.28 to 0.36)	0%	4 ^{168,207,215,221}	0.37
٠	Asia	0.24 (0.18 to 0.33)	34%	16 ^{204,211,2} 14,217-220,222-230	
•	Cirrhosis 0 to 10%	0.22 (0.16 to 0.31)	0%	11 ^{168,211,2} 17,220,222-224,226-228,230	0.08
٠	Cirrhosis >10%	0.31 (0.27 to 0.35)	7%	7 ^{204,20} 7,214,219,221,225,229	

*Study accounted for age, sex, fibrosis stage, and HCV genotype in analysis.

[†]All studies conducted in Asia and had duration >5 years.

[‡]All studies had duration >5 years.

Abbreviations: CI = confidence interval; HR = hazard ratio; U.S. = United States.

			HCV		Antiviral		Rates of	Incremental cost-	
Screening	Author	Screening	prevalence	Background	therapy costs	HCV infection	linkage to	effectiveness	
General	Barocas 2018 ²⁴⁷	A: ≥18 years B: ≥30 years C: ≥40 years D: Birth cohort	(range) NR (incidence in PWID 12 cases/100 person-years)	Per 100 person-years PWID: 33.1 Non-PWID: 2.6 to 2.7	(range) \$69,078 (\$0 to \$114,000)	F0 to F3: 0.94 (0.0 to 1.0) F4: 0.75 (0.6 to 0.9) Decompensated: 0.60 (0.48 to 0.75)	 care <30 years: 17.9% ≥30 years: 28.9% 	A: \$28,000/QALY B: Dominated C: Dominated D: Reference	HCV Cost-Effectiveness model. All screening strategies included risk- based screening; model included reinfection
adult population	Eckman 2018 ²⁴⁸	A: ≥18 years B: Birth cohort C: No screening	Birth cohort: 2.6% Non-birth cohort: 0.29%	Not included in model	\$24,270 (\$24,270 to \$74,760)	F0 to F3: 0.79 (NR) F4: 0.79 (NR) Decompensated: 0.72 (NR) Post-transplant: 0.75 (NR) HCC: 0.72 (NR)	100%	A: \$11,378/QALY B: Reference C: Dominated	Screening strategies did not include risk-based screening; model did not include reinfection
15 to 30 years old	Assoumou 2018 ²⁴⁹	9 1-time HCV screening strategies in 15 to 30 year olds vs. risk- based testing	NR (incidence 15.6/100 person-years)	PWID: 5% Non-PWID: 3%	\$71,950 to \$137,820 (\$26,480 to \$206,730)	F0 to F3: NR F4: 0.62 (0.55 to 0.75) Decompensated: 0.48 (0.40-0.60)	53%	Counselor-initiated, routine rapid testing: \$71,000/QALY Physician-ordered, counselor- performed targeted rapid testing: \$40,000/QALY Counselor-initiated, targeted rapid testing: \$44,000/QALY Other screening strategies: Dominated Risk-based testing: Reference	Hepatitis C Cost- Effectiveness model. Screening strategies varied with respect to routine vs. expanded targeted vs. current risk- based screening; counselor/tester vs. physician-initiated; rapid vs. standard test. Counselor-initiated, routine rapid testing associated with greater average QALY gain (0.007 to 0.11) compared with the other two non-dominated strategies and below \$100,000/QALY willingness-to-pay threshold

Table 22. Hepatitis C Cost Effectiveness Analyses

			HCV		Antiviral		Rates of	Incremental cost-	
Screening	Author	Screening	prevalence	Background	therapy costs	HCV infection	linkage to	effectiveness	
population	year	strategies	(range)	testing rates	(range)	utilities (range)	care	ratios	Comments
Prenatal screening	Chaillon 2019 ²⁵⁰	A: Prenatal screening B: Risk-based screening	0.73%	5% per year	\$25,000 (no range reported)	F0: 0.93 (0.83 to 1.0) F1, F2: 0.86 (0.78 to 0.94) F3: 0.83 (0.78 to 0.89) F4: 0.81 (0.68 to 0.89) Decompensated cirrhosis: 0.70 (0.56 to 0.79) HCC: 0.67 (0.56 to 0.78) Post-transplant: 0.71 (0.69 to 0.79)	Appears to be 100%	A: \$2,826/QALY B: Reference	Costs and effects on neonate not modelled; antiviral therapy administered postpartum; model did not appear to include reinfection
	Tasillo 2019 ²⁵¹	A: Prenatal screening B: Current practice	0.38%	During pregnancy: 14% No risk behaviors: 4 per 100 person-years With risk behavior: 40 per 100 person-years	No cirrhosis: \$39,600 (\$19,800 to \$59,400) Cirrhosis: \$68,773 (\$47,833 to \$89,712)	F0 to F3: 0.94 (0.94 to 1.0) F4: 0.75 (NR) Decompensated cirrhosis: 0.60 (NR)	Linked to care: 25% Initiated treatment if linked: 92%	A: \$41,000/QALY B: Reference	HCV Cost-Effectiveness model. Costs and effects on neonate not modelled; antiviral therapy offered 6 months postpartum

Abbreviations: HCC = hepatocellular carcinoma; HCV = hepatitis C virus; NR = not reported; PWID = people who inject drugs; QALY = quality-adjusted life year.

KQ	Number of Studies (k) Number of participants (n) Study Designs	Summary of Findings	Consistency and Precision	Overall quality	Body of Evidence Limitations	EPC Assessment of Strength of Evidence for KQ	Applicability
KQ 1a. Benefits of screening	No studies						
KQ 1b. Prenatal screening and vertical transmission	No studies						
KQ 2.Screening strategies	No studies						
KQ 3. Screening strategies and yield	Prior review: k=5 studies (n=8,044) New evidence: k=1 study (n=5,917)	The prior review included 5 studies that found risk-based screening associated with sensitivities of >90% and numbers needed to screen to identify 1 case of HCV infection of <20. One new study found that perfect application of risk-based guidelines would identify 82% of HCV cases with a number needed to screen to identify one case of HCV infection of 14.6, while applying a birth cohort strategy would result in 76% of cases identified a number needed to screen of 28.7.	Reasonable consistent and precise.	Fair	Studies were retrospective and in some studies significant proportions of patients were not tested. No studies of the yield of one-time versus repeat screening, alternative screening strategies in different risk groups, or the yield of currently recommended screening versus expanded screening strategies.	Low	Most studies conducted in high- prevalence settings. One study assumed perfect application of risk-based screening, which has not been attainable.
KQ 4. Harms of screening	Prior review: k=5 studies (n=288) New evidence: No new studies	Poor-quality evidence from the prior review suggested potential negative psychological and social effects of screening. No new studies on harms of screening were identified.	Low consistency and precision	Poor	Small sample sizes, no unscreened comparison group, reliance on retrospective recall, poorly defined outcomes.	Low	Studies were conducted in the era of interferon- based treatments

	Number of Studies (k)					EPC Assessment	
	Number of narticipants (n)		consistency	Overall	Body of Evidence	of Strength	
KQ	Study Designs	Summary of Findings	Precision	quality	Limitations	for KQ	Applicability
KQ 5. Effectiveness of interventions to prevent vertical transmission	Prior review: k=4 studies (n=1,724) New evidence: k=1 study (n=1,301)	Mode of delivery and risk of mother- to-infant transmission (5 studies, 1 new): No clear association Prolonged rupture of membrane (1 study from prior review): Adjusted OR 9.3, 95% Cl, 1.5 to 180 Internal fetal monitoring (1 study from prior review): Adjusted OR 6.7, 95% Cl, 1.1 to 35.9 Breastfeeding (3 studies, 1 new): No clear association	Mode of delivery: Inconsistent; some imprecision Rupture of membranes and fetal monitoring: Unable to assess consistency, imprecise Fetal monitoring: NA; imprecise Breastfeeding: Inconsistent; some imprecision	Fair	All studies were observational. Most studies from prior review were poor-quality and didn't perform statistical adjustment for potential confounders and were excluded. Prolonged rupture of membranes and internal monitoring only evaluated in 1 study each.	Low	Studies were conducted in the U.S. or Europe One study excluded women who were HIV positive; in the remaining 4 studies, HIV infection rates ranged from 5% to 15%

KQ	Number of Studies (k) Number of participants (n) Study Designs	Summary of Findings	Consistency and Precision	Overall quality	Body of Evidence Limitations	EPC Assessment of Strength of Evidence for KQ	Applicability
KQ 6. Effect of treatment on health outcomes - Adults	Prior review: NA (outdated regimens) New evidence: k=37 (34 trials [n=4,434], 2 pooled analyses [n=2,706], and 3 observational studies [n=58,892])	Two pooled analyses of 3 and 4 trials each and data from 3 other trials not included in pooled analyses found small, short-term improvements in quality of life scale scores after compared with before DAA therapy. In 31 DAA trials reporting short-term (<1 year) mortality, there were no deaths in 21 trials; mortality was low in the remaining 10 trials (0.4% [17/3,848] overall.) Two large observational studies found use of both DAA associated with lower rates of cardiovascular events and hepatocellular cancer. These associations were not found in a third, smaller observational study with shorter duration of followup.	Consistent, imprecise	Fair	Trials reporting quality of life and function were not randomized, used an open-label design, and did not have a non-DAA comparison group. Trials provided short- term followup, and were not designed to assess health outcomes. Event rates for mortality were low across studies, and other health outcomes were not widely reported. Evidence on long-term clinical outcomes was limited to 3 observational studies.	Low	Trials did not enroll a high proportion of patients with cirrhosis at baseline and evaluated current DAA regimens. Evidence on effects on hepatocellular cancer and cardiovascular events was primarily derived from a VA database that included few female subjects (3- 4%).
KQ 6. Effect of treatment on health outcomes - Adolescents	k=5 (3 trials [n=230] and 2 post-hoc observational studies [n=152])	There were no deaths in 3 trials of DAA regimens reporting short-term mortality. Sofosbuvir with ledipasvir or ribavirin and glecaprevir with pibrentasvir were associated with small improvements in Pediatric Quality of Life Inventory scores compared to baseline.	Cannot determine (for quality of life); imprecise	Fair	Trials were not designed to assess long-term health outcomes. The only evidence on quality of life outcomes is based on a post-hoc analysis of trial data.	Low	One trial evaluated a DAA regimen not FDA-approved for use in adolescents.

	Number of					EDC	
ĸq	Studies (k) Number of participants (n) Study Designs	Summary of Findings	Consistency and Precision	Overall guality	Body of Evidence Limitations	Assessment of Strength of Evidence for KQ	Applicability
KQ 7. Effect of treatment on SVR - Adults	Prior review: NA (outdated regimens) New evidence: k=49 trials (n=9,917; 27 multi- arm trials and 22 single arm trials)	DAA vs. placebo (1 RCT): SVR 99% vs. 0%, RR 231.6, 95% Cl, 14.6 to 3680 DAA vs. telaprevir (2 RCTs): SVR 98% vs. 80%, RR 1.22 (95% Cl, 1.09 to 1.37) and 99% vs. 66%, RR 1.50 (95% Cl, 1.22 to 1.85) In 49 trials, SVR rates with DAA therapies ranged from 95% to 100% across genotypes. Estimates were consistent in subgroup analyses based on study quality, geographic setting, fibrosis status, prior treatment experience, and other factors. Results were also similar in trials that stratified patients according to age, sex, race or ethnicity, or treatment-experience.	Consistent; precise	Good	All studies were industry-funded. Most DAA trials did not include a non-DAA comparison group. Evidence was most robust for genotype 1 and more limited for genotypes 2 through 6.	High	SVR rates based on currently recommended DAA regimens. Trials did not enroll a high proportion of patients with cirrhosis at baseline. Most trials enrolled predominantly white participants. Persons with current or recent drug use excluded from most trials. Most trials were conducted in the U.S. or Europe or were multinational.
KQ 7. Effect of treatment on SVR - Adolescents	Prior review: NA k=7 single arm trials (n=348)	In seven trials, the SVR rate ranged from 97% to 100%. Rates were similar when stratified according to DAA treatment regimen, genotype and treatment history.	Consistent; imprecise	Fair	Evidence in adolescents with genotype 2 and 4 infection was very limited (n=20) Four trials were industry funded.	Fair	Three trials evaluated DAA regimens not FDA- approved for use in adolescents. Four trials were multinational (primarily U.S. and Europe) and three were conducted in Egypt.

	Number of Studies (k) Number of participants (n)		Consistency and	Overall	Body of Evidence	EPC Assessment of Strength of Evidence	
KQ	Study Designs	Summary of Findings	Precision	quality	Limitations	for KQ	Applicability
KQ 8. Harms – Adults: DAA vs. placebo	k=4 trials (n=2,113)	 Pooled adverse event rates, DAA versus placebo: Any adverse event (4 trials): RR 1.12, 95% Cl, 1.02 to 1.24, l²=46 Serious adverse events (4 trials): RR 1.90, 95% Cl, 0.73 to 4.95, l²=0% Withdrawal due to adverse events (4 trials): RR 1.90, 95% Cl, 0.73 to 4.95, l²=0% Withdrawal due to adverse events (4 trials): RR 0.47, 95% Cl, 0.14 to 1.58, l²=14% Headache (4 trials): RR 1.12, 95% Cl, 0.91 to 1.37, l²=0% Nausea (3 trials): RR 1.42, 95% Cl, 1.00 to 2.03, l²=10% Diarrhea (2 trials): RR 1.53, 95% Cl, 0.78 to 2.68, l²=29% Fatigue (3 trials): RR 1.05, 95% Cl, 0.78 to 1.40; l²=32% Anemia (1 trial): RR 2.21, 95% Cl, 0.11 to 46 	Consistent; precise	Fair	Most trials did not have a non-DAA comparison group. Reporting of methods used to assess and define was suboptimal. Trials did not report long-term follow-up.	Moderate	See KQ 7

ко	Number of Studies (k) Number of participants (n) Study Designs	Summary of Findings	Consistency and Precision	Overall	Body of Evidence Limitations	EPC Assessment of Strength of Evidence for KQ	Applicability
KQ 8. Harms – Adults: DAA vs. other treatment	k=2 trials (n=459)	 Pooled adverse event rates, DAA versus other treatment: Any adverse event (2 trials): RR 0.65, 95% Cl, 0.50 to 0.84, l²=87% Serious adverse events (2 trials): RR 0.08, 95% Cl, 0.02 to 0.34, l²=0% Headache (2 trials): RR 0.78, 95% Cl, 0.58 to 1.04; l²=0% Headache (2 trials): RR 0.78, 95% Cl, 0.58 to 1.04; l²=0% Withdrawal due to adverse events (2 trials): RR 0.06, 95% Cl, 0.01 to 0.29, l²=0% Fatigue (2 trials): RR 0.37, 95% Cl, 0.21 to 0.63, l²=32% Headache (2 trials): RR 0.70, 95% Cl, 0.52 to 0.95; l²=0% Nausea (2 trials): RR 0.31, 95% Cl, 0.16 to 0.59, l2=65% Anemia (2 trials): RR 0.09, 95% Cl, 0.04 to 0.23, l²=41% Rash (2 trials): RR 0.19, 95% Cl, 0.06 to 0.58, l²=48% 	Consistent; precise	Fair	Most trials did not have a non-DAA comparison group. Reporting of methods used to assess and define was suboptimal. Trials did not report long-term follow-up.	Moderate	See KQ 7

ĸQ	Number of Studies (k) Number of participants (n) Study Designs	Summary of Findings	Consistency and Precision	Overall quality	Body of Evidence Limitations	EPC Assessment of Strength of Evidence for KQ	Applicability
KQ 8. Harms of treatment – Adults: Overall	Prior review: NA (outdated regimens) New evidence: k=49 trials (n=9,917)	 Pooled adverse events rates for currently recommended DAA regimens were: Any adverse event (44 trials): 73.3%, 95% Cl, 68.0% to 78.1%; l²=95% Serious adverse events (44 trials): 1.9%, 95% Cl, 1.5% to 2.4%; l²=31% Withdrawal due to adverse events (44 trials): 1.9%, 95% Cl, 1.5% to 2.4%; l²=31% Withdrawal due to adverse events (44 trials): 0.4%, 95% Cl, 0.3% to 0.6%; l²=0% Anemia (13 trials): 2.4%, 95% Cl, 0.9% to 6.3%; l²=85% Fatigue (37 trials): 18.4%, 95% Cl, 0.9% to 6.3%; l²=85% Fatigue (37 trials): 18.4%, 95% Cl, 15.6% to 21.7%; l²=90% Headache (42 trials): 18.7%, 95% Cl, 15.6% to 22.2%; l²=90% Insomnia (18 trials): 8.3%, 95% Cl, 6.8% to 10.1%; l²=58% Nausea (36 trials): 11.1%; 95% Cl, 9.1% to 13.5%, l²=82% Diarrhea (18 trials): 8.7%, 95% Cl, 6.9% to 11.0%; l²=70% Vomiting (6 trials): 5.8%, 95% Cl, 3.4% to 9.7%; l²=43% Rash (17 trials): 5.4%, 95% Cl, 4.1% to 7.1%; l²=70% 	Consistent; precise	Fair	Most trials did not have a non-DAA comparison group. Reporting of methods used to assess and define was suboptimal. Trials did not report long-term follow-up.	Moderate	See KQ 7

ĸQ	Number of Studies (k) Number of participants (n) Study Designs	Summary of Findings	Consistency and Precision	Overall quality	Body of Evidence Limitations	EPC Assessment of Strength of Evidence for KQ	Applicability
KQ 8. Harms of treatment – Adolescents	Prior review: NA New evidence: k=7 trials (n=348)	Five trials reported no withdrawals due to adverse events. There was one serious adverse event (grade 3 joint injury) in 1 trial. The rate of any adverse event was 27% in one trial and 71 to 87% in four trials. Specific adverse event rates were: • Headache (7 trials): 3 to 48% • Fatigue (7 trials): 5 to 53% • Gastrointestinal adverse events (5 trials): 3 to 28% • Insomnia (1 trial): 23%	Inconsistent; imprecise	Fair	Trials did not have a non-DAA comparison group. There was high variability in adverse event rates, with no clear trends when results were stratified according to regimen. Reporting of methods used to assess harms was suboptimal and long-term followup (>48 weeks) was not reported	Fair	See KQ 6 - Adolescents

	Number of Studies (k) Number of		Consistency	Overall	Body of Evidence	EPC Assessment of Strength	
KQ	Study Designs	Summary of Findings	Precision	quality	Limitations	for KQ	Applicability
KQ 9. Association between SVR and health outcomes	Prior review: 19 studies (n=30,692) New evidence: k=30 (n=116,821 [n=27,367 from studies included in the prior report + n=89,454 from new studies)	 Pooled estimates for health outcomes for SVR versus no SVR, in studies in which <25% of the population had cirrhosis at baseline: All-cause mortality (13 studies, 5 new): HR 0.40, 95% Cl, 0.28 to 0.56; l²=52% Liver mortality (4 studies, 0 new): HR 0.11, 95% Cl, 0.04 to 0.27; l²=0% Cirrhosis (4 cohorts reported in 3 studies, all new): HR 0.36, 95% Cl, 0.33 to 0.40; l²=0%) Hepatocellular carcinoma (20 studies, 16 new): HR 0.29, 95% Cl, 0.23 to 0.38; l²=19% Estimates were consistent in analyses stratified according to duration of follow-up, geographic setting, and level of statistical adjustment for potential confounders. 	Consistent, precise	Fair	Studies are observational and susceptible to confounding. Some studies appeared to evaluate overlapping patient populations. About half (k=13) of the studies did not address four pre-specified potential confounders in analyses (age, sex, fibrosis stage, and genotype).	Fair	Most studies evaluated SVR after interferon- based therapy; evidence on SVR after DAA therapy was limited to two studies, one of which reported imprecise estimates. Studies did not enroll a high proportion of patients with cirrhosis at baseline. Patients primarily received interferon- containing therapy. Six of seven U.S. studies conducted in VA populations. Over half of studies conducted in Asia, though results similar in U.S./Europe studies.

Abbreviations: ARD = adjusted risk difference; CI = confidence interval; DAA = direct acting antiviral; EPC = Evidence-based Practice Center; FDA = US Food and Drug Administration; HCV = hepatitis C virus; HR = hazard ratio; KQ = Key Question; NA = not applicable; OR = odds ratio; RCT = randomized controlled trial; RR = relative risk; SVR = sustained virologic response; U.S. = United States; VA = Veterans Affairs.

Key Questions 1-4

Database: Ovid MEDLINE(R) 1996 to February Week 1 2019

- 1. Hepatitis C/ or Hepatitis C, Chronic/ or Hepacivirus/
- 2. ("Hepatitis C" or hepacivirus* or HCV).ti,ab.
- 3. 1 or 2
- 4. Mass Screening/ or Population Surveillance/ or Sentinel Surveillance/ or Seroepidemiologic Studies/ or Prenatal Diagnosis/ or Neonatal Screening/ or Infectious Disease Transmission, Vertical/ or Disease Transmission, Infectious/ or tm.fs. or transmi*.ti,ab. or ((public* or communit* or universal* or widespread or open* or unrestricted or group* or adult* or adolescen* or pregnan* or antibod*) adj3 (screen* or test* or surveillance).ti,ab.
- 5. 3 and 4
- 6. limit 5 to yr="2012 -Current"
- 7. 6 and (random* or control* or trial or cohort or group*).ti,ab.
- 8. limit 6 to (clinical trial, all or comparative study or randomized controlled trial)
- 9. 7 or 8
- 10. limit 9 to (english language and humans)

Database: EBM Reviews - Cochrane Central Register of Controlled Trials February 2019

- 1. Hepatitis C/ or Hepatitis C, Chronic/ or Hepacivirus/
- 2. ("Hepatitis C" or hepacivirus* or HCV).ti,ab.
- 3. 1 or 2
- 4. Mass Screening/ or Population Surveillance/ or Sentinel Surveillance/ or Seroepidemiologic Studies/ or Prenatal Diagnosis/ or Neonatal Screening/ or Infectious Disease Transmission, Vertical/ or Disease Transmission, Infectious/ or tm.fs. or transmi*.ti,ab. or ((public* or communit* or universal* or widespread or open* or unrestricted or group* or adult* or adolescen* or pregnan* or antibod*) adj3 (screen* or test* or surveillance).ti,ab.
- 5. 3 and 4
- 6. limit 5 to yr="2012 -Current"

Key Question 5

Database: Ovid MEDLINE(R) 1996 to February Week 1 2019

- 1. Hepatitis C/ or Hepatitis C, Chronic/ or Hepacivirus/
- 2. ("Hepatitis C" or hepacivirus* or HCV).ti,ab.
- 3. 1 or 2
- 4. Infectious Disease Transmission, Vertical/ or Pregnancy Complications, Infectious/
- 5. Maternal-Fetal Exchange/
- 6. exp Breast Feeding/ or (breastfeed or breast feed* or breastfed or breast fed or breast milk).ti,ab.
- 7. (pregnan* or mother or maternal or child* or infan* or neonat* or prenatal or perinatal).ti,ab.
- 8. and tm.fs.
- 9. 3 and (4 or 5 or 6 or 8)
- 10. (random\$ or control\$ or trial or cohort or group* or compar*).ti,ab.
- 11. limit 9 to (clinical trial, all or comparative study or randomized controlled trial)
- 12. 9 and 10
- 13. 11 or 12
- 14. limit 13 to yr="2012 -Current"
- 15. limit 14 to (english language and humans)

Database: EBM Reviews - Cochrane Central Register of Controlled Trials Feburary 2019

- 1. Hepatitis C/ or Hepatitis C, Chronic/ or Hepacivirus/
- 2. ("Hepatitis C" or hepacivirus* or HCV).ti,ab.
- 3. 1 or 2
- 4. Infectious Disease Transmission, Vertical/ or Pregnancy Complications, Infectious/
- 5. Maternal-Fetal Exchange/
- 6. exp Breast Feeding/ or (breastfeed or breast feed* or breastfed or breast fed or breast milk).ti,ab.

Appendix A1. Search Strategies

- 7. (pregnan* or mother or maternal or child* or infan* or neonat* or prenatal or perinatal).ti,ab.
- 8. 7 and tm.fs.
- 9. 3 and (4 or 5 or 6 or 8)
- 10. limit 9 to yr="2012 -Current"

Key Questions 6-7

Database: Ovid MEDLINE(R) 1946 to February Week 1 2019

1 (Hepatitis C/ or Hepatitis C, Chronic/ or Hepacivirus/) and dt.fs.

2 ("Hepatitis C" or hepacivirus* or HCV).ti,ab.

3 1 or 2

4 Antiviral Agents/ad, tu

5 (daclatasvir or dasabuvir or elbasvir or glecaprevir or grazoprevir or ledipasvir or ombitasvir or paritaprevir or pibrentasvir or ribavirin or simeprevir or sofosbuvir or velpatasvir or voxilaprevir).ti,ab,kw

64 or 5

7 3 and 6

8 7 not (transplant* or HIV or "hepatitis B").ti.

9 limit 8 to yr="2012 -Current"

10 9 and exp Clinical Studies as Topic/

11 limit 9 to (clinical trial, all or meta analysis or randomized controlled trial or systematic reviews)

12 9 and (random* or control* or trial or "systematic review" or "meta-analysis" or metaanalysis).ti,ab.

13 10 or 11 or 12

14 limit 13 to (english language and humans)

Database: EBM Reviews - Cochrane Central Register of Controlled Trials February 2019

- 1. (Hepatitis C/ or Hepatitis C, Chronic/ or Hepacivirus/) and dt.fs.
- 2. ("Hepatitis C" or hepacivirus* or HCV).ti,ab.
- 3. 1 or 2
- 4. Antiviral Agents/ad, tu [Administration & Dosage, Therapeutic Use]
- 5. (daclatasvir or dasabuvir or elbasvir or glecaprevir or grazoprevir or ledipasvir or ombitasvir or paritaprevir or pibrentasvir or ribavirin or simeprevir or sofosbuvir or velpatasvir or voxilaprevir).mp.
- 6. 4 or 5
- 7. 3 and 6
- 8. 7 not (transplant* or HIV or "hepatitis B").ti.
- 9. limit 8 to yr="2012 -Current"

Key Question 8

Database: Ovid MEDLINE(R) 1996 to February Week 1 2019

- 1. (Hepatitis C/ or Hepatitis C, Chronic/ or Hepacivirus/) and dt.fs.
- 2. ("Hepatitis C" or hepacivirus* or HCV).ti,ab.
- 3. 1 or 2
- 4. Antiviral Agents/ad, tu [Administration & Dosage, Therapeutic Use]
- 5. (daclatasvir or dasabuvir or elbasvir or glecaprevir or grazoprevir or ledipasvir or ombitasvir or paritaprevir or pibrentasvir or ribavirin or simeprevir or sofosbuvir or velpatasvir or voxilaprevir).mp.
- 6. 4 or 5
- 7. 3 and 6
- 8. 7 not (transplant* or HIV or "hepatitis B").ti.
- 9. limit 8 to yr="2012 -Current"
- 10. 9 and exp Clinical Studies as Topic/
- 11. limit 9 to (clinical trial, all or meta analysis or randomized controlled trial or systematic reviews)
- 12. 9 and (random* or control* or trial or "systematic review" or "meta-analysis" or metaanalysis).ti,ab.
- 13. 10 or 11 or 12
- 14. limit 13 to (english language and humans)
- 15. 9 not 14
- 16. 15 and (ae or co or mo or po or to or ct).fs.

Appendix A1. Search Strategies

- 17. 15 and (adverse or safety or harm* or complication* or "side-effect*" or "treatment emerg*").ti,ab.
- 18. 16 or 17
- 19. limit 18 to (english language and humans)

Database: EBM Reviews - Cochrane Central Register of Controlled Trials February 2019

- 10. (Hepatitis C/ or Hepatitis C, Chronic/ or Hepacivirus/) and dt.fs.
- 11. ("Hepatitis C" or hepacivirus* or HCV).ti,ab.
- 12. 1 or 2
- 13. Antiviral Agents/ad, tu [Administration & Dosage, Therapeutic Use]
- 14. (daclatasvir or dasabuvir or elbasvir or glecaprevir or grazoprevir or ledipasvir or ombitasvir or paritaprevir or pibrentasvir or ribavirin or simeprevir or sofosbuvir or velpatasvir or voxilaprevir).mp.
- 15. 4 or 5
- 16. 3 and 6
- 17. 7 not (transplant* or HIV or "hepatitis B").ti.
- 18. limit 8 to yr="2012 -Current"

Key Question 9

Database: Ovid MEDLINE(R) 1996 to February Week 1 2019

- 1. (Hepatitis C/ or Hepatitis C, Chronic/ or Hepacivirus/) and dt.fs.
- 2. ("Hepatitis C" or hepacivirus* or HCV).ti,ab.
- 3. 1 or 2
- 4. Antiviral Agents/ad, tu [Administration & Dosage, Therapeutic Use]
- 5. (daclatasvir or dasabuvir or elbasvir or glecaprevir or grazoprevir or ledipasvir or ombitasvir or paritaprevir or pibrentasvir or ribavirin or simeprevir or sofosbuvir or velpatasvir or voxilaprevir).mp.
- 6. 4 or 5
- 7. 3 and 6
- 8. sustained virologic response/
- 9. ("sustained virologic response" or svr).ti,ab.
- 10. 8 or 9
- 11. 7 and 10
- 12. Liver Cirrhosis/
- 13. Liver Transplantation/
- 14. (cirrho* or transplant* or decompensat* or morbidity or mortality or death*).ti,ab.
- 15. 11 and (12 or 13 or 14)
- 16. limit 15 to yr="2012 -Current"
- 17. limit 16 to (english language and humans)

Database: EBM Reviews - Cochrane Central Register of Controlled Trials February 2019

- 1. (Hepatitis C/ or Hepatitis C, Chronic/ or Hepacivirus/) and dt.fs.
- 2. ("Hepatitis C" or hepacivirus* or HCV).ti,ab.
- 3. 1 or 2
- 4. Antiviral Agents/ad, tu [Administration & Dosage, Therapeutic Use]
- 5. (daclatasvir or dasabuvir or elbasvir or glecaprevir or grazoprevir or ledipasvir or ombitasvir or paritaprevir or pibrentasvir or ribavirin or simeprevir or sofosbuvir or velpatasvir or voxilaprevir).mp.
- 6. 4 or 5
- 7. 3 and 6
- 8. sustained virologic response/
- 9. ("sustained virologic response" or svr).ti,ab.
- 10. 8 or 9
- 11. 7 and 10
- 12. Liver Cirrhosis/
- 13. Liver Transplantation/
- 14. (cirrho* or transplant* or decompensat* or morbidity or mortality or death*).ti,ab.
- 15. 11 and (12 or 13 or 14)
- 16. limit 15 to yr="2012 -Current"

All Key Questions

Database: EBM Reviews - Cochrane Database of Systematic Reviews 2005 to February 6, 2019

- 1. ("Hepatitis C" or hepacivirus* or HCV).ti.
- 2. (daclatasvir or dasabuvir or elbasvir or glecaprevir or grazoprevir or ledipasvir or ombitasvir or paritaprevir or pibrentasvir or ribavirin or simeprevir or sofosbuvir or velpatasvir or voxilaprevir).ti,ab.
- 3. 1 and 2
- 4. screen*.mp. [mp=title, short title, abstract, full text, keywords, caption text]
- 5. 1 and 4
- 6. 3 or 5
- 7. limit 6 to full systematic reviews

PICOTS	Inclusion Criteria	Exclusion Criteria
Populations	Screening in nonpregnant adolescents and adults	Screening in nonpregnant adolescents and
	(KQs 1a, 2–4) Screening in pregnant adolescents	adults (KQs 1a, 2-4) Persons with known
	and adults (KQs 1–4) Asymptomatic, pregnant and	abnormal liver function tests, hepatitis B virus
	nonpregnant adolescents (ages 12 to 17 years) and	infection, or HIV infection; children age <12 years
	adults without prior HCV infection	Screening in pregnant adolescents and adults
	Labor and delivery and perinatal interventions (KQ	(KQs 1–4) Persons with known abnormal liver
	5) Pregnant adolescents and adults with HCV infection	function tests, hepatitis B virus infection, or HIV
	Antiviral treatment (KQs 6–8) Persons with screen-	infection
	detected or asymptomatic HCV infection (patients with a	Labor and delivery and perinatal interventions
	METAVIR fibrosis stage of 0–3, if symptom status is	(KQ 5)
	NR); persons with no prior antiviral treatment; includes	Other populations
	pregnant women	Antiviral treatment (KQS 6–8) Association
	Association between improvements in sustained	between improvements in sustained virologic
	Persona with UCV infaction being tracted with antiviral	response and clinical outcomes (RQ 9) Persons
	therease	transplant notionts, persons with repairing 6 virus of HIV,
Intonyontiono	Sereening in nennregnant adelessents and adults	Sereening in perprogram adelessents and
Interventions	(KOs 1a, 2-4) Screening in program adolescents	$\frac{1}{2}$
	and adults (KOs 1-4) Screening	addlescents and adults (KOs $1-4$) I abor and
	Labor and delivery and perinatal interventions (KO	delivery and perinatal interventions (K0.5) Other
	5) Mode of delivery labor management strategies	interventions
	breastfeeding practices	Antiviral treatment (KQs 6–8) Interferon-based
	Antiviral treatment (KQs 6–8) Currently recommended	treatment and other nonrecommended regimens*
	direct acting antiviral regimens*	a called a called the called a construction of the called a construction o
	Association between improvements in sustained	
	virologic response and clinical outcomes (KQ 9)	
	Direct acting antiviral regimens or other antiviral	
	treatment	
Comparisons	Screening in nonpregnant adolescents and adults	Screening in nonpregnant adolescents and
	(KQs 1a, 2–4) Screening in pregnant adolescents	adults (KQs 1a, 2–4) Screening in pregnant
	and adults (KQs 1-4) Screening vs. no screening, one	adolescents and adults (KQs 1–4) Labor and
	screening method vs. another, screening interval	delivery and perinatal interventions (KQ 5) Other
	comparisons	comparisons
	Labor and delivery and perinatal interventions (KQ	
	5) Elective cesarean delivery vs. vaginal or emergency	
	cesarean delivery, internal retai monitoring vs. no	
	monitoring, longer vs. shorter duration of rupture of	
	Antiviral treatment (KOs 6–9) Another direct acting	
	antiviral regimen or older antiviral regimen; includes	
	clinical trials without a comparison group	
	Association between improvements in sustained	
	virologic response and clinical outcomes (KO 9)	
	Persons who experience a sustained virologic response	
	vs. those who do not	

PICOTS	Inclusion Criteria	Exclusion Criteria
Outcomes	Screening in nonpregnant adolescents and adults	Screening in nonpregnant adolescents and
	(KQs 1a, 2–4) Mortality, morbidity (e.g., cirrhosis,	adults (KQs 1a, 2-4) Other outcomes, including
	hepatic decompensation, liver transplant, extrahepatic	intermediate outcomes
	manifestations of HCV infection), quality of life, HCV	Screening in pregnant adolescents and adults
	transmission, harms (e.g., labeling, anxiety, drug-related	(KQs 1–4) Labor and delivery and perinatal
	harms), screening yield (number of new diagnoses per	interventions (KQ 5) Other outcomes
	tests performed) (KQ 3)	Antiviral treatment (KQs 6–8) Association
	Screening in pregnant adolescents and adults (KQs	between improvements in sustained virologic
	1–4) Perinatal transmission, mortality, morbidity, quality	response and clinical outcomes (KQ 9)
	of life, harms (e.g., labeling, anxiety, drug-related	Histologic outcomes, liver function tests
	harms), screening yield (number of new diagnoses per	
	tests performed) (KQ 3)	
	Labor and delivery and perinatal interventions (KQ	
	Perinatal transmission of HCV infection	
	Antiviral treatment (KQs 6–8) Sustained virologic	
	response (KQ 7); morbidity (e.g., cirrhosis, hepatic	
	decompensation, liver transplant, extrahepatic	
	manifestations of HCV infection), mortality, quality of	
	life, HCV transmission (KQ 6), harms of treatment (KQ	
	8); behavioral outcomes will be included for Contextual	
	Question 3	
	Association between improvements in sustained	
	virologic response and clinical outcomes (KQ 9)	
	Morbidity (e.g., cirrhosis, hepatic decompensation, liver	
	transplant), mortality	
Setting	Screening in nonpregnant adolescents and adults	
	(KQs 1a, 2–4) Screening in pregnant adolescents	
	and adults (KQS 1–4) U.S. primary care,	
	obstetrics/gynecology, emergency department, and	
	primary care-applicable settings, including settings that	
	backh ages (a group structure to primary care and behavioral	
	Labor and delivery and peripatel interventions (KO	
	5) U.S. Jabor and delivery settings	
	Antiviral treatment (KOs 6–8) Association between	
	improvements in sustained virologic response and	
	clinical outcomes (KQ 9) Clinical settings in which	
	HCV antiviral treatments are prescribed	
Study design	Screening in nonpregnant adolescents and adults	Screening in nonpregnant adolescents and
jg.:	(KQs 1a, 2–4) Screening in pregnant adolescents	adults (KQs 1a, 2–4) Screening in pregnant
	and adults (KQs 1–4) Labor and delivery and	adolescents and adults (KQs 1-4) Uncontrolled
	perinatal interventions (KQ 5) RCTs, controlled	studies
	observational studies	Labor and delivery and perinatal interventions
	Antiviral treatment (KQs 6–8) RCTs and uncontrolled	(KQ 5) Antiviral treatment (KQs 6–8) Case
	clinical trials; for harms and clinical outcomes (KQ 6),	reports, studies not reporting original data
	will also include large cohort and case-control studies;	Association between improvements in
	will consider good-quality systematic reviews of clinical	sustained virologic response and clinical
	trials	outcomes (KQ 9) Case-control studies, case
	Association between improvements in sustained	reports, studies not reporting original data
	virologic response and clinical outcomes (KQ 9)	
	Cohort studies	

*For clinical outcomes (KQs 6 and 9), previously recommended regimens will be used.

Abbreviations: HCV = hepatitis C virus; KQ = Key Question; NR = not reported; PICOTS = population, interventions, comparisons, outcomes, setting, study design; RCTs = randomized controlled trials; U.S. = United States.



*Other sources include prior reports, reference lists of relevant articles, systematic reviews, etc. *Some studies were included for multiple KQs.

Abbreviation: KQ = key question.

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Abergel A, Asselah T, Metivier S, et al. Ledipasvir-sofosbuvir in patients with hepatitis C virus genotype 5 infection: an openlabel, multicentre, single-arm, phase 2 study. Lancet Infect Dis. 2016b;16(4):459-64. doi: 10.1016/S1473-3099(15)00529-0. PMID: 26803446.

Abergel A, Metivier S, Samuel D, et al. Ledipasvir plus sofosbuvir for 12 weeks in patients with hepatitis C genotype 4 infection. Hepatology. 2016a;64(4):1049-56. doi: 10.1002/hep.28706. PMID: 27351341.

Afdhal N, Zeuzem S, Kwo P, et al. Ledipasvir and sofosbuvir for untreated HCV genotype 1 infection. N Engl J Med. 2014;370(20):1889-98. doi: 10.1056/NEJMoa1402454. PMID: 24725239.

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Andreone P, Colombo MG, Enejosa JV, et al. ABT-450, ritonavir, ombitasvir, and dasabuvir achieves 97% and 100% sustained virologic response with or without ribavirin in treatment-experienced patients with HCV genotype 1b infection. Gastroenterology. 2014;147(2):359-65.e1. doi: 10.1053/j.gastro.2014.04.045. PMID: 24818763.

Arase Y, Ikeda K, Suzuki F, et al. Long-term outcome after interferon therapy in elderly patients with chronic hepatitis C. Intervirology. 2007;50(1):16-23. doi: 10.1159/000096308. PMID: 17164553.

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Appendix A5. List of Excluded Studies With Reasons For Exclusion

Abad S, Vega A, Hernandez E, et al. Universal sustained viral response to the combination of ombitasvir/paritaprevir/ritonavir and dasabuvir with/without ribavirin in patients on hemodialysis infected with hepatitis C virus genotypes 1 and 4. Am J Nephrol. 2017;45(3):267-72. doi: 10.1159/000454819. PMID: 28166520. Excluded for wrong study design for Key Question.

Abad S, Vega A, Rincon D, et al. Effectiveness of direct-acting antivirals in hepatitis C virus infection in haemodialysis patients. Nefrologia. 2017;37(2):158-63. doi: 10.1016/j.nefro.2016.10.003. PMID: 27914803. Excluded for wrong population.

Abdel-Aziz AM, Ibrahim MA, El-Sheikh AA, et al. Effect of sofosbuvir plus daclatasvir in hepatitis C virus genotype-4 patients: promising effect on liver fibrosis. J Clin Exp Hepatol. 2018;8(1):15-22. doi: 10.1016/j.jceh.2017.06.006. PMID: 29743792. Excluded for wrong population.

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Abdel-Moneim A, Aboud A, Abdel-Gabaar M, et al. Efficacy and safety of sofosbuvir plus daclatasvir with or without ribavirin: large real-life results of patients with chronic hepatitis C genotype 4. Hepatol Int. 2018;12(4):348-55. doi: 10.1007/s12072-018-9868-8. PMID: 29754329. Excluded for wrong population.

Abd-Elsalam S, Badawi R, Elnawasany S, et al. Sofosbuvir, pegylated interferon and ribavirin in treatment of an Egyptian cohort with hepatitis C virus infection in real life clinical practice. Infect Disord Drug Targets. 2018;12:12. doi: 10.2174/1871526518666180912121835. PMID: 30207250. Excluded for wrong intervention.

Abergel A, Hezode C, Leroy V, et al. Peginterferon alpha-2b plus ribavirin for treatment of chronic hepatitis C with severe fibrosis: a multicentre randomized controlled trial comparing two doses of peginterferon alpha-2b. J Viral Hepat. 2006;13(12):811-20. doi: 10.1111/j.1365-2893.2006.00768.x. PMID: 17109680. Excluded for outdated medication.

Abouelkheir Abdalla D, Ali Elhadidy T, Besheer T, et al. Respiratory adverse effects of sofosbuvir-based regimens for treatment of chronic hepatitis C virus. Egypt J Chest Dis Tuberc. 2016;10. Excluded for wrong intervention.

Adler H, Lambert JS. Daclatasvir for the treatment of hepatitis C virus infection. Expert Rev Gastroenterol Hepatol. 2014;8(7):725-38. doi: 10.1586/17474124.2014.925798. PMID: 24882552. Excluded for systematic review or meta-analysis used as a source document only to identify individual studies.

Afdhal N, Reddy KR, Nelson DR, et al. Ledipasvir and sofosbuvir for previously treated HCV genotype 1 infection. N Engl J Med. 2014;370(16):1483-93. doi: 10.1056/NEJMoa1316366. PMID: 24725238. Excluded for wrong population.

Aghemo A, Prati GM, Rumi MG, et al. Sustained virological response prevents the development of insulin resistance in patients with chronic hepatitis C. Hepatology. 2012;56(5):1681-7. doi: 10.1002/hep.25867. PMID: 22619107. Excluded for wrong population.

Ahmed OA, Elsebaey MA, Fouad MHA, et al. Outcomes and predictors of treatment response with sofosbuvir plus daclatasvir with or without ribavirin in Egyptian patients with genotype 4 hepatitis C virus infection. Infect Drug Resist. 2018;11:441-5. doi: 10.2147/IDR.S160593. PMID: 29628768. Excluded for wrong population.

Ahn SH, Lim YS, Lee KS, et al. A phase 3b study of sofosbuvir plus ribavirin in treatment-naive and treatment-experienced Korean patients chronically infected with genotype 2 hepatitis C virus. J Viral Hepat. 2016;23(5):358-65. doi: 10.1111/jvh.12499. PMID: 26864153. Excluded for wrong intervention.

Airoldi J, Berghella V. Hepatitis C and pregnancy. Obstet Gynecol Surv. 2006;61(10):666-72. doi: 10.1097/01.ogx.0000238671.13495.33. PMID: 16978426. Excluded for systematic review or meta-analysis used as a source document only to identify individual studies.

Akahane T, Kurosaki M, Itakura J, et al. Real-world efficacy and safety of sofosbuvir + ribavirin for hepatitis C genotype 2: a nationwide multicenter study by the Japanese Red Cross Liver Study Group. Hepatol Res. 2018;01:01. doi: 10.1111/hepr.13246. PMID: 30171740. Excluded for wrong intervention.

Akiyama MJ, Kaba F, Rosner Z, et al. Hepatitis C screening of the "Birth Cohort" (born 1945-1965) and younger inmates of New York City jails. Am J Public Health. 2016;106(7):1276-7. doi: 10.2105/AJPH.2016.303163. PMID: 27196656. Excluded for wrong study design for Key Question.

Akiyama MJ, Kaba F, Rosner Z, et al. Correlates of hepatitis C virus infection in the targeted testing program of the New York City jail system. Public Health Rep. 2017;132(1):41-7. doi: 10.1177/0033354916679367. PMID: 28005477. Excluded for wrong study design for Key Question.

Akuta N, Kobayashi M, Suzuki F, et al. Liver fibrosis and body mass index predict hepatocarcinogenesis following eradication of hepatitis C virus RNA by direct-acting antivirals. Oncology. 2016;91(6):341-7. doi: 10.1159/000450551. PMID: 27694754. Excluded for wrong comparator.

Appendix A5. List of Excluded Studies With Reasons For Exclusion

Akuta N, Suzuki F, Seko Y, et al. Efficacy and anticarcinogenic activity of ribavirin combination therapy for hepatitis C virusrelated compensated cirrhosis. Intervirology. 2013;56(1):37-45. doi: 10.1159/000342746. PMID: 23037768. Excluded for wrong population.

Alavi M, Spelman T, Matthews GV, et al. Injecting risk behaviours following treatment for hepatitis C virus infection among people who inject drugs: the Australian trial in acute hepatitis C. Int J Drug Policy. 2015;26(10):976-83. doi: 10.1016/j.drugpo.2015.05.003. PMID: 26115881. Excluded for wrong study design for Key Question.

Alfaleh FZ, Alswat K, Helmy A, et al. The natural history and long-term outcomes in patients with chronic hepatitis C genotype 4 after interferon-based therapy. Liver Int. 2013;33(6):871-83. doi: 10.1111/liv.12127. PMID: 23490034. Excluded for poor quality.

Almario CV, Vega M, Trooskin SB, et al. Examining hepatitis C virus testing practices in primary care clinics. J Viral Hepat. 2012;19(2):e163-9. doi: 10.1111/j.1365-2893.2011.01539.x. PMID: 22239514. Excluded for wrong outcome.

Alqahtani SA, Afdhal N, Zeuzem S, et al. Safety and tolerability of ledipasvir/sofosbuvir with and without ribavirin in patients with chronic hepatitis C virus genotype 1 infection: analysis of phase III ION trials. Hepatology. 2015;62(1):25-30. doi: 10.1002/hep.27890. PMID: 25963890. Excluded for systematic review or meta-analysis used as a source document only to identify individual studies.

American College of Obstetricians and Gynecologists' Committee on Gynecological Practice. Committee opinion no. 655: hepatitis B, hepatitis C, and human immunodeficiency virus infections in obstetrician-gynecologists. Obstet Gynecol. 2016;127(2):e70-4. doi: 10.1097/AOG.0000000001315. PMID: 26942390. Excluded for not a study: letter, editorial, non-systematic review article, no original data.

Ampuero J, Reddy KR, Romero-Gomez M. Hepatitis C virus genotype 3: meta-analysis on sustained virologic response rates with currently available treatment options. World J Gastroenterol. 2016;22(22):5285-92. doi: 10.3748/wjg.v22.i22.5285. PMID: 27298572. Excluded for systematic review or meta-analysis used as a source document only to identify individual studies.

Anderson EM, Mandeville RP, Hutchinson SJ, et al. Evaluation of a general practice based hepatitis C virus screening intervention. Scott Med J. 2009;54(3):3-7. doi: 10.1258/rsmsmj.54.3.3. PMID: 19728405. Excluded for poor quality.

Anderson ES, Galbraith JW, Deering LJ, et al. Continuum of care for hepatitis C virus among patients diagnosed in the emergency department setting. Clin Infect Dis. 2017;64(11):1540-6. doi: 10.1093/cid/cix163. PMID: 28207069. Excluded for wrong outcome.

Anderson ES, Pfeil SK, Deering LJ, et al. High-impact hepatitis C virus testing for injection drug users in an urban ED. Am J Emerg Med. 2016;34(6):1108-11. doi: 10.1016/j.ajem.2016.03.004. PMID: 27037135. Excluded for wrong study design for Key Question.

Andriulli A, Cursaro C, Cozzolongo R, et al. Early discontinuation of ribavirin in HCV-2 and HCV-3 patients responding to peginterferon alpha-2a and ribavirin. J Viral Hepat. 2009;16(1):28-35. doi: 10.1111/j.1365-2893.2008.01044.x. PMID: 18761603. Excluded for outdated medication.

Anonymous. Corrections to: "Ledipasvir-sofosbuvir with or without ribavirin to treat patients with HCV genotype 1 infection and cirrhosis non-responsive to previous protease-inhibitor therapy: a randomised, double-blind, phase 2 trial (SIRIUS)" [Lancet Infect Dis (2015), 15: 397-404]. Lancet Infect Dis. 2015;15(7):761. doi: 10.1016/S1473-3099(15)00111-5. PMID: 26122440. Excluded for not a study: letter, editorial, non-systematic review article, no original data.

Anonymous. Erratum: simeprevir with pegylated interferon alfa 2a plus ribavirin in treatment-naive patients with chronic hepatitis C virus genotype 1 infection (QUEST-1): a phase 3, randomised, double-blind, placebo-controlled trial. Lancet. 2016;387(10030):1816. Excluded for not a study: letter, editorial, non-systematic review article, no original data.

Anonymous. Correction: ombitasvir, paritaprevir, and ritonavir plus ribavirin for chronic hepatitis C virus genotype 4 infection in Egyptian patients with or without compensated cirrhosis (AGATE-II): a multicentre, phase 3, partly randomised open-label trial. Lancet Gastroenterol Hepatol. 2016;1(1):e1. Excluded for not a study: letter, editorial, non-systematic review article, no original data.

Aqel B, Leise M, Vargas HE, et al. Multicenter experience using ledipasvir/sofosbuvir +/- RBV to treat HCV GT 1 relapsers after simeprevir and sofosbuvir treatment. Ann Hepatol. 2018;17(5):815-21. doi: 10.5604/01.3001.0012.3142. PMID: 30145562. Excluded for wrong population.

Aqel BA, Pungpapong S, Leise M, et al. Multicenter experience using simeprevir and sofosbuvir with or without ribavirin to treat hepatitis C genotype 1 in patients with cirrhosis. Hepatology. 2015;62(4):1004-12. doi: 10.1002/hep.27937. PMID: 26096332. Excluded for wrong population.

Appendix A5. List of Excluded Studies With Reasons For Exclusion

Arora S, O'Brien C, Zeuzem S, et al. Treatment of chronic hepatitis C patients with persistently normal alanine aminotransferase levels with the combination of peginterferon alpha-2a (40 kDa) plus ribavirin: impact on health-related quality of life. J Gastroenterol Hepatol. 2006;21(2):406-12. doi: 10.1111/j.1440-1746.2005.04059.x. PMID: 16509866. Excluded for wrong outcome.

Asahina Y, Itoh Y, Ueno Y, et al. Ledipasvir-sofosbuvir for treating Japanese patients with chronic hepatitis C virus genotype 2 infection. Liver Int. 2018;38(9):1552-61. doi: 10.1111/liv.13685. PMID: 29297980. Excluded for wrong population.

Ascione A, De Luca M, Tartaglione MT, et al. Peginterferon alfa-2a plus ribavirin is more effective than peginterferon alfa-2b plus ribavirin for treating chronic hepatitis C virus infection. Gastroenterology. 2010;138(1):116-22. doi: 10.1053/j.gastro.2009.10.005. PMID: 19852964. Excluded for outdated medication.

Asselah T. Sofosbuvir-based interferon-free therapy for patients with HCV infection. J Hepatol. 2013;59(6):1342-5. doi: 10.1016/j.jhep.2013.07.023. PMID: 23891655. Excluded for wrong study design for Key Question.

Asselah T, Hezode C, Qaqish RB, et al. Ombitasvir, paritaprevir, and ritonavir plus ribavirin in adults with hepatitis C virus genotype 4 infection and cirrhosis (AGATE-I): a multicentre, phase 3, randomised open-label trial. Lancet Gastroenterol Hepatol. 2016;1(1):25-35. doi: 10.1016/S2468-1253(16)30001-2. PMID: 28404108. Excluded for wrong population.

Asselah T, Tran T, Alves K, et al. Efficacy and safety of glecaprevir/pibrentasvir in patients with HCV genotype 5 or 6 infection: The ENDURANCE-5,6 study. J Hepatol. 2018;68:S39. doi: 10.1016/s0168-8278(18)30294-0. Excluded for not a study: letter, editorial, non-systematic review article, no original data.

Atsukawa M, Tsubota A, Toyoda H, et al. Efficacy and safety of ombitasvir/paritaprevir/ritonavir and ribavirin for chronic hepatitis patients infected with genotype 2a in Japan. Hepatol Res. 2018;28:28. doi: 10.1111/hepr.13292. PMID: 30485638. Excluded for wrong population.

Attia D, El Saeed K, Elakel W, et al. The adverse effects of interferon-free regimens in 149 816 chronic hepatitis C treated Egyptian patients. Aliment Pharmacol Ther. 2018;47(9):1296-305. doi: 10.1111/apt.14538. PMID: 29504152. Excluded for wrong population.

Aziz H, Aziz M, Gill ML. Analysis of host and viral-related factors associated to direct acting antiviral response in hepatitis C virus patients. Viral Immunol. 2018;31(3):256-63. doi: 10.1089/vim.2017.0124. PMID: 29664710. Excluded for wrong intervention.

Azzaroli F, Accogli E, Nigro G, et al. Interferon plus ribavirin and interferon alone in preventing hepatocellular carcinoma: a prospective study on patients with HCV related cirrhosis. World J Gastroenterol. 2004;10(21):3099-102. PMID: 15457551. Excluded for wrong population.

Babatin MA, AlGhamdi AS, Assiri AM, et al. Treatment efficacy of ledipasvir/sofosbuvir for 8 weeks in non-cirrhotic chronic hepatitis C genotype 4 patients. Saudi J Gastroentero. 2019;25(1):55-60. doi: 10.4103/sjg.SJG_189_18. PMID: 30117490. Excluded for wrong intervention.

Backus LI, Belperio PS, Shahoumian TA, et al. Comparative effectiveness of ledipasvir/sofosbuvir +/- ribavirin vs. ombitasvir/paritaprevir/ritonavir + dasabuvir +/- ribavirin in 6961 genotype 1 patients treated in routine medical practice. Aliment Pharmacol Ther. 2016;44(4):400-10. doi: 10.1111/apt.13696. PMID: 27291852. Excluded for wrong study design for Key Question.

Bang C, Song I. Impact of antiviral therapy on hepatocellular carcinoma and mortality in patients with chronic hepatitis C: systematic review and meta-analysis. BMC Gastroenterol. 2017;17(1):46. doi: 10.1186/s12876-017-0606-9. PMID: 28376711. Excluded for systematic review or meta-analysis used as a source document only to identify individual studies.

Barron J, Xie Y, Wu SJ, et al. Treatment of chronic hepatitis C infection with sofosbuvir-based regimens in a commercially insured patient population. Am Health Drug Benefits. 2016;9(6):327-35. PMID: 27924186. Excluded for wrong study design for Key Question.

Bassiony MM, Yousef A, Yousef U, et al. Major depressive disorder and generalized anxiety disorder and response to treatment in hepatitis C patients in Egypt. Int J Psychiatry Med. 2015;50(2):147-62. doi: 10.1177/0091217415605029. PMID: 26405268. Excluded for wrong intervention.

Basu P, Shah NJ, Aloysius MM, Brown RS Jr. . Interferon ineligible naïve chronic hepatitis C genotype I subjects treated with simeprevir and sofosbuvir in special population (psychiatric). A clinical pilot study; Inspire C study; interim results. HPB (Oxford). 2015;S2(17):46. Excluded for wrong comparator.

Bell AM, Wagner JL, Barber KE, et al. Elbasvir/Grazoprevir: a review of the latest agent in the fight against hepatitis C. Int J Hepatol. 2016;2016:3852126. doi: 10.1155/2016/3852126. PMID: 27403342. Excluded for not a study: letter, editorial, non-systematic review article, no original data.
Belperio PS, Shahoumian TA, Loomis TP, et al. Real-world effectiveness of daclatasvir plus sofosbuvir and velpatasvir/sofosbuvir in hepatitis C genotype 2 and 3. J Hepatol. 2019;70(1):15-23. doi: 10.1016/j.jhep.2018.09.018. PMID: 30266283. Excluded for wrong study design for Key Question.

Berden FA, Aaldering BR, Groenewoud H, et al. Identification of the best direct-acting antiviral regimen for patients with hepatitis C virus genotype 3 infection: a systematic review and network meta-analysis. Clin Gastroenterol Hepatol. 2017;15(3):349-59. doi: 10.1016/j.cgh.2016.10.034. PMID: 27840182. Excluded for systematic review or meta-analysis used as a source document only to identify individual studies.

Bernardinello E, Cavalletto L, Chemello L, et al. Long-term clinical outcome after beta-interferon therapy in cirrhotic patients with chronic hepatitis C. TVVH Study Group. Hepatogastroenterology. 1999;46(30):3216-22. PMID: 10626189. Excluded for wrong population.

Bernstein D, Kleinman L, Barker CM, et al. Relationship of health-related quality of life to treatment adherence and sustained response in chronic hepatitis C patients. Hepatology. 2002;35(3):704-8. doi: 10.1053/jhep.2002.31311. PMID: 11870387. Excluded for outdated medication.

Bezemer G, Van Gool AR, Verheij-Hart E, et al. Long-term effects of treatment and response in patients with chronic hepatitis C on quality of life. An international, multicenter, randomized, controlled study. BMC Gastroenterol. 2012;12:11. doi: 10.1186/1471-230X-12-11. PMID: 22292521. Excluded for wrong intervention.

Bini EJ, Mehandru S. Sustained virological response rates and health-related quality of life after interferon and ribavirin therapy in patients with chronic hepatitis C virus infection and persistently normal alanine aminotransferase levels. Aliment Pharmacol Ther. 2006;23(6):777-85. doi: 10.1111/j.1365-2036.2006.02819.x. PMID: 16556180. Excluded for wrong outcome.

Bonkovsky HL, Woolley JM. Reduction of health-related quality of life in chronic hepatitis C and improvement with interferon therapy. The Consensus Interferon Study Group. Hepatology. 1999;29(1):264-70. doi: 10.1002/hep.510290124. PMID: 9862876. Excluded for wrong outcome.

Boyd SD, Harrington P, Komatsu TE, et al. HCV genotype 4, 5 and 6: Distribution of viral subtypes and sustained virologic response rates in clinical trials of approved direct-acting antiviral regimens. J Viral Hepat. 2018;25(8):969-75. doi: 10.1111/jvh.12896. PMID: 29577495. Excluded for wrong study design for Key Question.

Braks RE, Ganne-Carrie N, Fontaine H, et al. Effect of sustained virological response on long-term clinical outcome in 113 patients with compensated hepatitis C-related cirrhosis treated by interferon alpha and ribavirin. World J Gastroenterol. 2007;13(42):5648-53. PMID: 17948941. Excluded for wrong population.

Bronowicki JP, Ouzan D, Asselah T, et al. Effect of ribavirin in genotype 1 patients with hepatitis C responding to pegylated interferon alfa-2a plus ribavirin. Gastroenterology. 2006;131(4):1040-8. doi: 10.1053/j.gastro.2006.07.022. PMID: 17030174. Excluded for wrong intervention.

Bronowicki JP, Pol S, Thuluvath PJ, et al. Randomized study of asunaprevir plus pegylated interferon-alpha and ribavirin for previously untreated genotype 1 chronic hepatitis C. Antivir Ther. 2013;18(7):885-93. doi: 10.3851/IMP2660. PMID: 23804631. Excluded for wrong intervention.

Brook RA, Kleinman NL, Su J, et al. Absenteeism and productivity among employees being treated for hepatitis C. Am J Manag Care. 2011;17(10):657-64. PMID: 22106459. Excluded for wrong outcome.

Bruix J, Poynard T, Colombo M, et al. Maintenance therapy with peginterferon alfa-2b does not prevent hepatocellular carcinoma in cirrhotic patients with chronic hepatitis C. Gastroenterology. 2011;140(7):1990-9. doi: 10.1053/j.gastro.2011.03.010. PMID: 21419770. Excluded for wrong population.

Bruno G, Saracino A, Fabrizio C, et al. Safety and effectiveness of a 12-week course of sofosbuvir and simeprevir+/-ribavirin in HCV-infected patients with or without HIV infection: a multicentre observational study. Int J Antimicrob Agents. 2017;49(3):296-301. doi: 10.1016/j.ijantimicag.2016.11.030. PMID: 28163136. Excluded for wrong study design for Key Question.

Bruno S, Stroffolini T, Colombo M, et al. Sustained virological response to interferon-alpha is associated with improved outcome in HCV-related cirrhosis: a retrospective study. Hepatology. 2007;45(3):579-87. doi: 10.1002/hep.21492. PMID: 17326216. Excluded for wrong population.

Buti M, Calleja JL, Lens S, et al. Simeprevir in combination with sofosbuvir in treatment-naive and -experienced patients with hepatitis C virus genotype 4 infection: a Phase III, open-label, single-arm study (PLUTO). Aliment Pharmacol Ther. 2017;45(3):468-75. doi: 10.1111/apt.13883. PMID: 27896822. Excluded for wrong intervention.

Buti M, Dominguez-Hernandez R, Casado MA, et al. Healthcare value of implementing hepatitis C screening in the adult general population in Spain. PLoS One. 2018;13(11):e0208036. doi: 10.1371/journal.pone.0208036. PMID: 30485377. Excluded for wrong study design for Key Question.

Buti M, Flisiak R, Kao JH, et al. Alisporivir with peginterferon/ribavirin in patients with chronic hepatitis C genotype 1 infection who failed to respond to or relapsed after prior interferon-based therapy: FUNDAMENTAL, a Phase II trial. J Viral Hepat. 2015;22(7):596-606. doi: 10.1111/jvh.12360. PMID: 25412795. Excluded for wrong intervention.

Butt AA, Ren Y, Marks K, et al. Do directly acting antiviral agents for HCV increase the risk of hepatic decompensation and decline in renal function? Results from ERCHIVES. Aliment Pharmacol Ther. 2017;45(1):150-9. doi: 10.1111/apt.13837. PMID: 27813162. Excluded for wrong study design for Key Question.

Butt AA, Yan P, Marks K, et al. Adding ribavirin to newer DAA regimens does not affect SVR rates in HCV genotype 1 infected persons: results from ERCHIVES. Aliment Pharmacol Ther. 2016;44(7):728-37. doi: 10.1111/apt.13748. PMID: 27459341. Excluded for wrong study design for Key Question.

Butt AA, Yan P, Shaikh OS, et al. Hepatitis B reactivation and outcomes in persons treated with directly acting antiviral agents against hepatitis C virus: results from ERCHIVES. Aliment Pharmacol Ther. 2018a;47(3):412-20. doi: 10.1111/apt.14426. PMID: 29181838. Excluded for wrong study design for Key Question.

Cacoub P, Desbois AC, Comarmond C, et al. Impact of sustained virological response on the extrahepatic manifestations of chronic hepatitis C: a meta-analysis. Gut. 2018:2025-34. doi: 10.1136/gutjnl-2018-316234. PMID: 29703790. Excluded for systematic review or meta-analysis used as a source document only to identify individual studies.

Cacoub P, Lidove O, Maisonobe T, et al. Interferon-alpha and ribavirin treatment in patients with hepatitis C virus-related systemic vasculitis. Arthritis Rheum. 2002;46(12):3317-26. doi: 10.1002/art.10699. PMID: 12483738. Excluded for wrong intervention.

Cardoso AC, Moucari R, Figueiredo-Mendes C, et al. Impact of peginterferon and ribavirin therapy on hepatocellular carcinoma: incidence and survival in hepatitis C patients with advanced fibrosis. J Hepatol. 2010;52(5):652-7. doi: 10.1016/j.jhep.2009.12.028. PMID: 20346533. Excluded for wrong population.

Caroleo B, Colangelo L, Perticone M, et al. Efficacy and safety of elbasvir-grazoprevir fixed dose in the management of polytreated HCV patients: evidence from real-life clinical practice. J Clin Pharmacol. 2018;58(10):1248-53. doi: 10.1002/jcph.1135. PMID: 29746724. Excluded for wrong outcome.

Cha RR, Lee SS, Lee CM, et al. Clinical features and outcomes of patients with genotype 3 hepatitis C virus infection in Korea: a retrospective observational study. Medicine. 2016;95(6):e2755. doi: 10.1097/MD.00000000002755. PMID: 26871824. Excluded for wrong population.

Chahine EB, Sucher AJ, Hemstreet BA. Sofosbuvir/velpatasvir: the first pangenotypic direct-acting antiviral combination for hepatitis C. Ann Pharmacother. 2017;51(1):44-53. doi: 10.1177/1060028016668897. PMID: 27609942. Excluded for systematic review or meta-analysis used as a source document only to identify individual studies.

Chamorro-de-Vega E, Gimenez-Manzorro A, Rodriguez-Gonzalez CG, et al. Effectiveness and safety of ombitasvirparitaprevir/ritonavir and dasabuvir with or without ribavirin for HCV genotype 1 infection for 12 weeks under routine clinical practice. Ann Pharmacother. 2016;50(11):901-8. doi: 10.1177/1060028016659306. PMID: 27422641. Excluded for wrong population.

Chan HLY, Tsang OTY, Hui YT, et al. Real-life efficacy and safety of paritaprevir/ ritonavir, ombitasvir and dasabuvir combination with or without ribavirin in difficult-to-treat genotype 1 chronic hepatitis C patients in Hong Kong. J Gastroenterol Hepatol. 2016;31(378). Excluded for wrong population.

Chang KC, Ye YH, Wu CK, et al. Risk factors for development of hepatocellular carcinoma in patients with chronic hepatitis C without sustained response to combination therapy. J Formos Med Assoc. 2018;117(11):1011-8. doi: 10.1016/j.jfma.2017.11.008. PMID: 29254684. Excluded for wrong population.

Chang SC, Yang SS, Chang CC, et al. Assessment of health-related quality of life in antiviral-treated Taiwanese chronic hepatitis C patients using SF-36 and CLDQ. Health Qual Life Outcomes. 2014;12:97. doi: 10.1186/1477-7525-12-97. PMID: 24941994. Excluded for wrong intervention.

Chayama K, Notsumata K, Kurosaki M, et al. Randomized trial of interferon- and ribavirin-free ombitasvir/paritaprevir/ritonavir in treatment-experienced hepatitis C virus-infected patients. Hepatology. 2015;61(5):1523-32. doi: 10.1002/hep.27705. PMID: 25644279. Excluded for wrong intervention.

Chen J, Shi J, Xie WF, et al. Meta-analysis: amantadine may lower the efficacy of pegylated interferon plus ribavirin in treatment-naive hepatitis C genotype 1 patients. Int J Infect Dis. 2012;16(10):e748-52. doi: 10.1016/j.ijid.2012.06.002. PMID: 22836046. Excluded for wrong intervention.

Cheng PN, Chiu YC, Chien SC, et al. Real-world effectiveness and safety of sofosbuvir plus daclatasvir with or without ribavirin for genotype 2 chronic hepatitis C in Taiwan. J Formos Med Assoc. 2018;11:11. doi: 10.1016/j.jfma.2018.09.016. PMID: 30316677. Excluded for wrong population.

Cho Y, Cho EJ, Lee JH, et al. Sofosbuvir-based therapy for patients with chronic hepatitis C: early experience of its efficacy and safety in Korea. Clin Mol Hepatol. 2015;21(4):358-64. doi: 10.3350/cmh.2015.21.4.358. PMID: 26770924. Excluded for wrong study design for Key Question.

Chopp S, Vanderwall R, Hult A, et al. Simeprevir and sofosbuvir for treatment of hepatitis C infection. Am J Health Syst Pharm. 2015;72(17):1445-55. PMID: 26294237. Excluded for wrong study design for Key Question.

Chung W, Kim KA, Jang ES, et al. Cost-effectiveness of sofosbuvir plus ribavirin therapy for hepatitis C virus genotype 2 infection in South Korea. J Gastroenterol Hepatol. 2018;21:21. doi: 10.1111/jgh.14554. PMID: 30462841. Excluded for wrong outcome.

Conte D, Fraquelli M, Prati D, et al. Prevalence and clinical course of chronic hepatitis C virus (HCV) infection and rate of HCV vertical transmission in a cohort of 15,250 pregnant women. Hepatology. 2000;31(3):751-5. doi: 10.1002/hep.510310328. PMID: 10706568. Excluded for poor quality.

Cornberg M, Petersen J, Schober A, et al. Real-world use, effectiveness and safety of anti-viral treatment in chronic hepatitis C genotype 3 infection. Aliment Pharmacol Ther. 2017;45(5):688-700. doi: 10.1111/apt.13925. PMID: 28078723. Excluded for wrong study design for Key Question.

Coughlan B, Sheehan J, Carr A, et al. Evaluation of a brief group based psychological/educational treatment programme for women with an iatrogenic chronic hepatitis C virus infection. J Clin Psychol Med Settings. 2004;11(4):303-14. Excluded for poor quality.

Coverdale SA, Khan MH, Byth K, et al. Effects of interferon treatment response on liver complications of chronic hepatitis C: 9year follow-up study. Am J Gastroenterol. 2004;99(4):ajg2004123. doi: 10.1111/j.1572-0241.2004.04085.x. PMID: 15089895. Excluded for wrong population.

Coyle C, Moorman AC, Bartholomew T, et al. The HCV care continuum: linkage to HCV care and treatment among patients at an urban health network, Philadelphia, PA. Hepatology. 2019 doi: 10.1002/hep.30501. PMID: 30633811. Excluded for wrong outcome.

Cucchetti A, D'Amico G, Trevisani F, et al. Effect of direct-acting antivirals on future occurrence of hepatocellular carcinoma in compensated cirrhotic patients. Dig Liver Dis. 2018;50(2):156-62. doi: 10.1016/j.dld.2017.10.004. PMID: 29102521. Excluded for wrong outcome.

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Randomized Controlled Trials (RCTs) and Cohort Studies

Criteria:

- Initial assembly of comparable groups:
 - For RCTs: Adequate randomization, including first concealment and whether potential confounders were distributed equally among groups
- For cohort studies: Consideration of potential confounders, with either restriction or measurement for adjustment in the analysis; consideration of inception cohorts
- Maintenance of comparable groups (includes attrition, cross-overs, adherence, contamination)
- Important differential loss to followup or overall high loss to followup
- Measurements: equal, reliable, and valid (includes masking of outcome assessment)
- Clear definition of interventions
- All important outcomes considered
- Analysis: adjustment for potential confounders for cohort studies or intention-to treat analysis for RCTs

Definition of ratings based on above criteria:

Good: Meets all criteria: Comparable groups are assembled initially and maintained throughout the study (followup \geq 80%); reliable and valid measurement instruments are used and applied equally to all groups; interventions are spelled out clearly; all important outcomes are considered; and appropriate attention to confounders in analysis. In addition, intention-to-treat analysis is used for RCTs.

Fair: Studies are graded "fair" if any or all of the following problems occur, without the fatal flaws noted in the "poor" category below: Generally comparable groups are assembled initially, but some question remains whether some (although not major) differences occurred with followup; measurement instruments are acceptable (although not the best) and generally applied equally; some but not all important outcomes are considered; and some but not all potential confounders are accounted for. Intention-to-treat analysis is used for RCTs.

Poor: Studies are graded "poor" if any of the following fatal flaws exists: Groups assembled initially are not close to being comparable or maintained throughout the study; unreliable or invalid measurement instruments are used or not applied equally among groups (including not masking outcome assessment); and key confounders are given little or no attention. Intention-to-treat analysis is lacking for RCTs.

Systematic Reviews

Criteria:

- Comprehensiveness of sources considered/search strategy used
- Standard appraisal of included studies
- Validity of conclusions
- Recency and relevance (especially important for systematic reviews)

Definition of ratings based on above criteria:

Good: Recent, relevant review with comprehensive sources and search strategies; explicit and relevant selection criteria; standard appraisal of included studies; and valid conclusions.

Fair: Recent, relevant review that is not clearly biased but lacks comprehensive sources and search strategies.

Poor: Outdated, irrelevant, or biased review without systematic search for studies, explicit selection criteria, or standard appraisal of studies.

Diagnostic Accuracy Studies

Criteria:

- Screening test relevant, available for primary care, and adequately described
- Credible reference standard, performed regardless of test results
- Reference standard interpreted independently of screening test
- Indeterminate results handled in a reasonable manner
- Spectrum of patients included in study
- Sample size
- Reliable screening test

Definition of ratings based on above criteria:

Good: Evaluates relevant available screening test; uses a credible reference standard; interprets reference standard independently of screening test; assesses reliability of test; has few or handles indeterminate results in a reasonable manner; includes large number (greater than 100) of broad-spectrum patients with and without disease.

Fair: Evaluates relevant available screening test; uses reasonable although not best standard; interprets reference standard independent of screening test; has moderate sample size (50 to 100 subjects) and a "medium" spectrum of patients.

Poor: Has a fatal flaw, such as: uses inappropriate reference standard; improperly administers screening test; biased ascertainment of reference standard; has very small sample size or very narrow selected spectrum of patients.

Source: U.S. Preventive Services Task Force Procedure Manual. Accessed at https://www.uspreventiveservicestaskforce.org/Page/Name/methods-and-processes

Appendix B Table 1. Key Question 5: Evidence Table Interventions to Prevent Mother-to-Infant Transmission of HCV Infection – Study Characteristics

Author year Country Study Name	Study Type	Definition of mother-to- infant transmission	Confounders assessed in	Duration of	Eligibility	Exclusion	Number screened/ eligible/ enrolled/	Demographic characteristics of study	HCV genotype HCV viral load HIV infection
	Droopostivo			24 months				Motornal aga (n=79)	
	cohort	anti HCV	rick factors	24 11011115	HUV pogativo	niv-positive	2447/ 79/	Modian (rango): 20 (21 to	status (n=78)
naiy	study	antibodies	(exposure to		women		78/	42)	$P_{0}(11-70)$
Fair	Study	hevond 18	blood		women		78	*Characteristics of HCV-	Negative: 18 (23%)
r an		months or HCV-	products and				70	RNA positive mothers	Negative. 10 (2070)
		nositive on two						(n-60)	*Characteristics of HCV-
		senarate tests	viral load					HCV risk factors	RNA positive mothers
			HCV					Absent: 25 (42%)	(n=60)
			genotype					Blood transfusion: 14 (23%)	denotype
			gestational					IVDU: 20 (33%)	1a: 9 (15%)
			age mode of					Blood transfusion and	1b: 25 (42%)
			delivery, birth					IVDU: 1 (2%)	2a: 20 (33%)
			weight					Mode of delivery	3: 6 (10%)
								Vaginal: 43 (72%)	Viral load
								Cesarean: 17 (28%)	<0.2X106: 9 (15%)
								Gestational age	>0.2X106: 51 (85%)
								<36 weeks: 9 (15%)	
								≥36 weeks: 51 (85%́)	
								Birth weight	
								<2500g: 14 (23%)	
								≥2500g: 46 (77%)	
								HCV risk factors	
								Absent: 25 (42%)	
								Blood transfusion: 14 (23%)	
								IVDU: 20 (33%)	
								Blood transfusion and	
								IVDU: 1 (2%)	
								Mode of delivery	
								Vaginal: 43 (72%)	
								Cesarean: 17 (28%)	
								Gestational age	
								<36 weeks: 9 (15%)	
								≥36 weeks: 51 (85%)	
								Birth weight	
								<2500g: 14 (23%)	
								≥2500g: 46 (77%)	
European	Multicenter	Children	Account for	Children	HCV infected	Second-born	1787/	Maternal age (n=1205)	Maternal HIV infection
Pediatric Hep C	prospective	considered	differences	received	mothers and	twins and	1479/	Mean (SD): 31.7 (5.17)	(n=1391)
Virus Network	cohort	infected if they	between	clinical	their singleton	second- and	1479	Median (range): 32 (17.1 to	Yes: 208 (15%)
2005 (Tovo) ¹⁰⁶	study	had ≥2 positive	centers in the	examinatio	infants or first-	third-born	/1220 (1034 HIV-)	45.1)	No: 1183 (85%)
		HCV RNA PCR	HCV RNA	n at birth, 6	born infants	triplets were		Mode of delivery (n=1455)	

Appendix B Table 1. Key Question 5: Evidence Table Interventions to Prevent Mother-to-Infant Transmission of HCV Infection – Study Characteristics

Author year Country		Definition of mother-to-	Confounders				Number screened/	Demographic	HCV genotype HCV viral load
Study Name		infant	assessed in	Duration of			eligible/ enrolled/	characteristics of study	HIV infection
Quality	Study Type	transmission	analysis	followup	Eligibility	Exclusion	analyzed	population	IVDU
Italy, Spain,		test results	PCR assays	weeks, and	from multiple	excluded.		Vaginal: 764 (52.5%)	Child HIV infection
Germany,		and/or were anti-	used to	3, 6, 9, 12,	pregnancies	Mother-infant		Emergency cesarean	(n=1435)
Ireland, U.K.,		HCV antibody	determine	18, and 24	with	pairs with		section: 160 (11%)	Yes: 10 (0.7%)
Norway, Sweden		positive after 18	infection,	months;	confirmed	infants of		Elective cesarean: 480	No: 1397 (97.4%)
Good		months.	allow for	and	HCV infection	indeterminate		(33%)	Indeterminate: 28
		Children	center-	thereafter	status.	infection		Cesarean section	(1.9%)
		considered	associated	every 6		status were		(unspecified): 51 (3.5%)	Maternal IVDU
		uninfected if	unobserved	months if		excluded.		Infant feeding type	(n=1162)
		they had <2	differences in	infected or				(n=1357)	History: 448 (38.6%)
		positive HCV	background	every year				Breast-fed: 452 (32.7%)	No history: 714 (61.4%)
		RNA PCR test	characteristic	lT uninfactad				Formula fed: $930 (67.3\%)$	
		results and ≤z	s, autions	unimected				Sex of child $(n=1470)$	
			a random					F_{0}	
		results and/or	a flact in					$G_{estational} = 200 (+3.476)$	
		were anti-HCV	multivariable					<34 weeks: 97 (7%)	
		antibody	models at the					35 to 36 weeks: 122 (8 8%)	
		negative after 18	center level					≥37 weeks [.] 1163 (84 2%)	
		months.							
Gibb 2000 ¹⁰⁵	Prospective	Positive result	Adjusted for	24 months	Mother known	U.K. children	499/	Maternal age (n=441)	Maternal HIV infection
Ireland, U.K.	cohort	for HCV	HIV status,		to be HCV	born before	441/	Mean (SD): 27 (6) Race	(n=441) Yes: 22 (5%)
Fair	study	antibody within	breastfeeding,		infected	1996	441/	(n=441)	No: 328 (74%)
		90 days of birth	and mode of		during		441	White: 413 (94%)	Unknown: 91 (21%)
			delivery		pregnancy or			Non-white: 28 (6%)	Maternal IVDU (n=441)
					if child had			Breastfeeding (n=414) Yes:	History: 343 (78%)
					positive result			59 (14%)	No history: 98 (22%)
					for HCV			No: 355 (86%)	
					antibody			Mode of delivery (n=424)	
					within 90 days			Vaginal: 339 (80%)	
					or dirth			Emergency cesarean: 54	
								(13%)	
Mast 2005 ¹⁰⁴	Prospective	Infant serum	Variables with	Infants horn	Women	Mothers with	75 909/	$\Delta q_{p} (p - 242)$	Mother HCV/ RNA+
ILS (Houston &	cohort	collected at hirth	n < 1 from the	to HCV+	nresenting for	serum testing	567/	$\sim 20.7 (2.9\%)$	(n-242) At enrollment or
Honolulu)	study	and 8 well-child	univariate	mothers	nrenatal care	as RIRA	332/	20 to 29: 103 (42 6%)	delivery: 194 (79 5%)
Good	otady	visits. Testing	analysis and	followed	(and in	indeterminate	242 women & 244	30 to 39: 120 (49.6%)	Both: 179 (77.2%)
		included	maternal	from birth to	Houston.	and HCV	infants	≥40: 12 (4.9%)	Delivery: 5 (2.2%)
		detection of	demographic	≥12	those who did	RNA negative		Race (n=242)	Enrollment: 4 (1.7%)
		antibody to	characteristic	months.	not receive	were		White: 79 (32.6%)	Maternal HIV infection
		HCV, detection	s included in	HCV-	prenatal care	excluded from		Black: 77 (31.8%)	(n=242): Yes: 11 (4.5%)
		of HCV RNA		infected	who	the analysis.		Hispanic: 49 (20.3%)	
Appendix B Table 1. Key Question 5: Evidence Table Interventions to Prevent Mother-to-Infant Transmission of HCV Infection – Study Characteristics

Author year		Definition of							HCV genotype
Country		mother-to-	Confounders				Number screened/	Demographic	HCV viral load
Study Name		infant	assessed in	Duration of			eligible/ enrolled/	characteristics of study	HIV infection
Quality	Study Type	transmission	analysis	followup	Eligibility	Exclusion	analyzed	population	IVDU
		(qualitative and	multivariate	infants	presented for				HIV and HCV RNA+
		quantitative),	analysis	followed	delivery at 2				(n=242)
		and genotyping.		annually	county				7 (2.9%)
				until age 5	hospitals)				Maternal IVDU (n=242)
					were offered				126 (52.3%) Geometric
					testing.				mean HCV RNA level at
					Women with				delivery (n=194) HIV-:
					positive anti-				2.38*106
					HCV test				Maternal HCV genotype
					results were				(n=116) 18.76 (66%)
					eproll (those				10.10(14%) 2b: 10(9%)
					with				20. 10 (376) 3a: 13 (11%)
					indeterminate				4a: 1 (01%)
					status were				
					invited to				
					enroll until				
					HCV status				
					was				
					confirmed).				
Resti 2002 ¹⁰⁷	Prospective	HCV RNA-	Maternal HCV	24 months	Anti-HCV	Twin pairs &	NR/	n=1372 mother-infant pairs	Maternal HCV viremia:
Italy	cohort	positive at any	RNA status,		positive	siblings	1493/	Maternal age: NR	Positive: 897 (65.4%)
Good		testing or	maternal HIV-		women		1493/	Type of delivery: Cesarean:	Negative: 387 (28.2%)
		persistence of	1 status,		attending 24		1372	377 (27.5%)	Missing: 88 (6.4%)
		anti-HCV	maternal		study sites			Vaginal: 924 (67.3%)	Maternal HIV-1 status:
		beyond age 2	IVDU, type of		between April			Missing: 71 (5.2%)	Positive: 194 (14.1%)
		years	reeding, mode		December			Propert: 260 (26 2%)	Negative: 1178 (85.9%)
			of delivery		1006			Eormula: $921(67.1\%)$	Maternal IV/DI I: Ves:
					1990			Missing: 91 (6.7%)	461 (33.6%)
								Birth weight a	No [•] 911 (66 4%)
								<2500: 145 (10.6%)	Missing: 0
								>2500: 1042 (83.2%)	
								Missing: 185 (6.2%)	
								Gestational age, weeks:	
								<36: 107 (7.8%)	
								>36: 1127 (82.1%)	
1								Missing 138 (10 1%)	

 Abbreviations: HCV = hepatitis C virus; IVDU = injection drug use; NR = not reported; PCR = polymerase chain reaction; RIBA = recombinant-immunoblot-assay; RNA = ribonucleic acid; SD = standard deviation; U.K. = United Kingdom; U.S. = United States.

Appendix B Table 2. Key Question 5: Evidence Table Interventions to Prevent Mother-to-Infant Transmission of HCV Infection – Study Outcomes

Author year Country Study Name <i>Quality</i>	Overall transmission	Transmission by labor management: IUPC	Transmission by labor management: Fetal monitoring	Transmission by labor management: Rupture of membranes	Transmission by route of delivery	Transmission by type of infant feeding
Ceci 2001 ¹⁰⁸ Italy <i>Fair</i>	Overall transmission (n=78) 2 consecutive positive tests: 8 (10%) 24 month followup: 2 (3%) not adjusted	NR	NR	NR	No association (data NR)	NR
European Pediatric Hep C Virus Network 2005 (Tovo) ¹⁰⁶ Italy, Spain, Germany, Ireland, U.K., Norway, Sweden <i>Good</i>	91/1479 6.2% (95% CI, 5.0% to 7.5%)	NR	NR	NR	Elective cesarean vs. emergency cesarean or vaginal delivery (n=1220) OR 1.66 (95% CI, 1.00 to 2.74) unadjusted, p=0.05 OR 1.46 (95% CI, 0.86 to 2.48) adjusted, p=0.16 HIV- mothers elective vs. emergency cesarean or vaginal delivery (n=1034) 1.57 (95% CI, 0.88 to 2.83) unadjusted, p=0.13 1.59 (95% CI, 0.88 to 2.86) adjusted, p=0.13 Adjusted for: sex, mode of delivery, prematurity, and infant feeding type	Breast vs. formula (n=1220) OR 0.74 (95% Cl, 0.42 to 1.31) unadjusted, p=0.30 OR .88 (95% Cl, 0.48 to 1.61) adjusted, p=0.68 HIV- mothers breast vs. formula (n=1034) OR 0.88 (95% Cl, 0.48 to 1.61) unadjusted, p=0.68 OR 0.92 (95% Cl, 0.50 to 1.70) adjusted, p=0.60

Appendix B Table 2. Key Question 5: Evidence Table Interventions to Prevent Mother-to-Infant Transmission of HCV Infection – Study Outcomes

Author year Country Study Name <i>Quality</i>	Overall transmission	Transmission by labor management: IUPC	Transmission by labor management: Fetal monitoring	Transmission by labor management: Rupture of membranes	Transmission by route of delivery	Transmission by type of infant feeding
Gibb 2000 ¹⁰⁵ Ireland, U.K. <i>Fair</i>	Overall (n=441) 6.7% (95% Cl, 4.1 to 10.2) unadjusted	NR	NR	NR	Elective cesarean vs. emergency cesarean vs. vaginal (n=424) 0% (95% Cl, 0 to 7.4) vs. 5.9% (95% Cl, 1.0 to 17.8) vs. 7.7% (4.5 to 11.9) OR elective cesarean 0 (95% Cl, 0 to 0.86) vs. OR emergency cesarean 0.84 (95% Cl, 0.12 to 3.63) Adjusted for HIV status and breastfeeding Elective cesarean vs. vaginal/emergency cesarean (n=424) 0% (85% Cl, 0 to 7.4) vs. 7.4% (95% Cl, 4.5 to 11.3) OR 0 (95% Cl, 0 to 0.87) Adjusted for: HIV status and breastfeeding	Breast vs. formula (n=414) 7.7% (95% Cl, 2.2 to 17.8) vs. 6.7% (95% Cl, 3.7 to 10.6) OR 1.52 (95% Cl, 0.35 to 5.12) Adjusted for: HIV status and mode of delivery
Mast 2005 ¹⁰⁴ U.S. (Houston & Honolulu) <i>Good</i>	9/244 (3.7%)	NR	Results are for HCV RNA+/HIV- mothers (n=188) Internal vs. external 3/16 (18.8%) vs. 4/165 (2.4%), RR 7.7 (1.9-31.6), p=0.02 Internal fetal monitoring AOR, 6.7 (95% CI, 1.1 to 35.9)	Results are for HCV RNA+/HIV- mothers (n=189) Rupture of membranes before onset of laboryes vs. no 4/45 (8.9%) vs. 3/137 (2.2%), RR 4.1 (95% CI, 0.9 to 17.5), p=0.06 Duration of membrane rupture <1 vs. 1-5 vs. $6-12$ vs. $\geq 130/53$ vs. 1/59 (1.7%) vs. 4/40 (10%) vs. 2/30 (6.7%), p=0.02 Membrane rupture >6 hours OR, 9.3 (95% CI, 1.5 to 179.7) Adjusted	Results are for HCV RNA+/HIV- mothers (n=188) Elective cesarean vs. emergency cesarean vs. vaginal delivery 0/12 (0%) vs. 1/18 (5.5%) vs. 6/151 (4%), elective cesarean RR undefined, emergency cesarean RR 1.4 (95% CI, 0.2 to 1.1), p=0.55 Elective cesarean vs. emergency cesarean/vaginal 0/12 vs. 7/169 (4%), RR 0.87 (95% CI, 0.05 to 14)	Results are for HCV RNA+/HIV- mothers (n=189) Breast vs. formula 2/62 (3.2%) vs. 5/120 (4.2%), RR 0.8 (95% CI, 0.2 to 3.9), p=1.0

Appendix B Table 2. Key Question 5: Evidence Table Interventions to Prevent Mother-to-Infant Transmission of HCV Infection – Study Outcomes

Author year Country		Transmission by labor	Transmission by labor	Transmission by labor management:	Transmission by route of	Transmission by time of infort
Study Name	Overall transmission		Fotol monitoring	Rupture of	delivery	fooding
						Dreast va farmula (n. 1001):
Resti 2002107	98/13/2 (7.1%, 95% CI,	NR	NR	NR	Cesarean vs. vaginai (n=1301):	Breast vs. formula (n=1281):
Italy	2.2 to 7.2%)				22/377 (5.8%) vs. 73/924 (7.9%);	22/360 (6.1%) vs. 73/921 (7.9%);
Good					Calculated OR (95% CI): OR 0.85	p=0.26; OR (95% CI): 0.86 (0.61
					(0.71 to 1.09)	to 1.10); AOR for breast (95% CI):
					Calculated AOR (95% CI): 0.83	0.95 (0.58 to 1.40)
					(0.65 to 1.08)	
					Per study for cesarean vs. vaginal	
					(ref): OR (95% CI): 1 17 (0 92 to	
					1 41; p=0.19; AOR for vaginal	
					(05%) (0.1): 1.20 (0.02 to 1.55):	
					(95% CI). 1.20 (0.93 to 1.55),	
					p=0.15	
					Note: Appears to have reversed	
					reference	

Abbreviations: AOR = adjusted odds ratio; CI = confidence interval; HCV = hepatitis C virus; IUPC = Intra-uterine pressure catheter; NR = not reported; OR = odds ratio; RNA = ribonucleic acid; RR = relative risk; U.K. = United Kingdom; U.S. = United States.

Appendix B Table 3. Key Question 5: Evidence Table Interventions to Prevent Mother-to-Infant Transmission of HCV Infection – Additional Study Outcomes

Author year Country Study Name <i>Quality</i>	Transmission by other risk factors (maternal)	Transmission by other risk factors (child)	Subgroup analyses	Adverse events	Funding source
Ceci 2001 ¹⁰⁸	Transmission from women with no known risk of	NR	NR	NR	NR
Italy	infection was significantly lower (RR=0.17%, 0.04-				
Fair	0.7370, p=0.0003)				
	By maternal blood transfusion (n=38)				
	2+ positive tests vs. 0 positive tests 3/8 (37.5%) vs. 2/30 (6.7%), p<0.05				
	By maternal viremia (n=38) 2+ positive tests vs. 0 positive tests 6.90 +/- 5.87 x 106 vs. 3.93 +/- 2.94 x 106				
	Note: Multivariate analysis found significant associations between HCV transmission and high maternal viral load, possession of HCV risk factors, and history of blood transfusion (p<0.05 for all, but no data shown); also states that no other variables				
	were found to be significantly associated with HCV transmission				

Appendix B Table 3. Key Question 5: Evidence Table Interventions to Prevent Mother-to-Infant Transmission of HCV Infection – Additional Study Outcomes

Author year Country					
Study Name	Transmission by other	Transmission by other	Subgroup	Adverse	Funding
Quality	risk factors (maternal)	risk factors (child)	analyses	events	source
European Pediatric Hep C Virus Network 2005 (Tovo) ¹⁰⁶ Italy, Spain, Germany, Ireland, U.K., Norway, Sweden <i>Good</i>	Mother HIV positive vs. negative (n=1220) OR 1.89 (95% Cl, 1.05 to 3.40) unadjusted, p=0.03 OR 1.82 (95% Cl, 0.94 to 3.52) adjusted, p=0.06	Female vs. male (n=1220) OR 2.12 (95% CI, 1.27 to 3.56) unadjusted, p=0.004 OR 2.07 (95% CI, 1.23 to 3.48) adjusted, p=0.006 Premature vs. term (n=1220) OR 0.54 (95% CI, 0.23 to 1.26) unadjusted, p=0.15 OR 0.45 (95% CI, 0.19 to 1.08) adjusted, p=0.07 HIV- mothers female vs. male (n=1034) OR 1.79 (95% CI, 1.00 to 3.22) unadjusted, p=0.05 OR 1.80 (95% CI, 1.00 to 3.24) adjusted, p=0.07 HIV- mothers premature vs. term (n=1034) OR 0.83 (95% CI, 0.32 to 2.13) unadjusted, p=0.69 OR 0.83 (95% CI, 0.32 to 2.15) adjusted, p=0.07	NR	NR	European Commission Regione Piemonte, Italy; U.K. Medical Research Council
Gibb 2000 ¹⁰⁵	HIV positive vs. negative (n=441)	NR	NR	NR	U.K. Department of
Ireland, U.K.	18.6% (95% CI, 5.8 to 38.6) vs. 6.4% (95% CI, 3.5				Health
Fair	to 10.3)				
	OR=3.8 (95% CI, 0.92 to 13.2)				
	Adjusted for: breastfeeding and HIV status				

Appendix B Table 3. Key Question 5: Evidence Table Interventions to Prevent Mother-to-Infant Transmission of HCV Infection – Additional Study Outcomes

Author year					
Country Study Name	Transmission by other	Transmission by other	Subaroup	Adverse	Funding
Quality	risk factors (maternal)	risk factors (child)	analyses	events	source
Mast 2005 ¹⁰⁴ U.S. (Houston & Honolulu) <i>Good</i>	Maternal HCV/RNA status at delivery positive vs. negative 9/190 (4.6%) vs. 0/54, RR undefined Remaining results are for HCV/RNA+ mothers (n=190) maternal HIV statuspositive vs. negative 2/8 (25%) vs. 7/182 (3.8%), RR 6.5 (95% CI, 1.6 to 26.4) Maternal HCV RNA level, genome copies/mL \leq 106 vs. >106, <107 vs. \geq 107 1/61 (1.6%) vs. 2/87 (2.3%) vs. 4/34 (11.8%), p=0.03 (results continued in next 2 columns)	Results for infants born to HCV/RNA+ mothers: (n=190) Sex Male vs. female 2/85 (2.3%) vs. 5/96 (5.2%), RR 0.45 (95% CI, 0.09 to 2.27), p=0.45 Gestational age <37 vs. ≥37 0/27 vs. 7/155 (4.5%), RR undefined, p=0.6 Birth weight <2500g vs. ≥2500g 1/22 (4.6%) vs. 6/160 (3.8%), RR 1.2 (95% CI, 0.2 to 9.6), p=1 Apgar score at 5 minutes ≤8 vs. >8 0/21 vs. 7/161 (4.4%),	NR	NR	Centers for Disease Control
Resti 2002 ¹⁰⁷ Italy <i>Good</i>	Maternal HCV RNA status positive vs. negative (n=1284): 97/897 (10.8%) vs. 1/387 (0.3%); p=0.00001; OR (95% CI): 6.83 (5.85 to 7.81) Maternal HIV Status positive vs. negative (n=1372): 75/1178 (6.4%) vs. 23/194 (11.9%); p=0.007; OR (95% CI): 1.41 (1.16 to 1.66); AOR (95% CI): 1.13 (0.85 to 1.51); p=0.38 (results continued in next 2 columns)	Infant birth weight <2500 g vs. >2500 g (n=1187): $8/145$ (5.5%) vs. $78/1042$ (7.5%); p=0.39; OR (95% CI): 1.17 (0.44 to 1.90) Gestational age <36 vs. >36 weeks (n=1149): $7/107$ (6.5%) vs. $86/1127$ (7.6%); p=0.68; OR (95% CI): 1.08 (0.69 to 1.47)	NR	NR	Italian Ministero della Ricerca Scientifica & Azienda Ospedaliera A. Meyer Research Department

Abbreviations: ALT = alanine aminotransferase; AOR = adjusted odds ratio; CI = confidence interval; HCV = hepatitis C virus; IVDU = injection drug use; NR = not reported; OR = odds ratio; RNA = ribonucleic acid; RR = relative risk; U.K. = United Kingdom; U.S. = United States.

Appendix B Table 4. Key Question 5: Quality Assessment of Studies of Interventions to Prevent Mother-to-Infant Transmission of HCV Infection

Author year	(1) Did the study attempt to enroll all (or a random sample of) patients meeting inclusion criteria, or a random sample (inception cohort)?	(2) Were the groups comparable at baseline on key prognostic factors (e.g., by restriction or matching)?	(3) Did the study maintain comparable groups through the study period?	(4) Did the study use accurate methods for ascertaining exposures and potential confounders?	(5) Were outcome assessors and/or data analysts blinded to the exposure being studied?	(6) Did the article report attrition?	(7) Is there important differential loss to followup or overall high loss to followup?	(8) Did the study perform appropriate statistical analyses on potential confounders?	(9) Were outcomes pre-specified and defined, and ascertained using accurate methods?	Overall Quality
Ceci 2001 ¹⁰⁸	Yes	Unclear	Unclear	Yes	No	Yes	Unclear	Yes	Yes	Fair
European Paediatric Hepatitis C Virus Network 2005 (Tovo) ¹⁰⁶	Yes	Unclear	Unclear	Yes	Unclear	No	No	Yes	Yes	Good
Gibb 2000 ¹⁰⁵	Unclear	Unclear	Unclear	Yes	No	No	No	Yes	Yes	Fair
Mast 2005 ¹⁰⁴	Yes	Unclear	Unclear	Yes	No	Yes	No	Yes	Yes	Good
Resti 2002 ¹⁰⁷	Yes	Unclear	Unclear	Yes	Unclear	Yes	No	Yes	Yes	Good

Author year			Treatment						
Country		Dates of	duration	Inclusion				_	Funding
Quality	Type of study	enrollment	Followup	criteria	Intervention(s)	N	Population	Outcomes	source
Butt 2019 ¹⁶⁹	Retrospective	NR	Treatment	Adults with HCV	A. DAA regimen	34,206	(A + B) vs. C	A vs. B vs. C	Gilead
U.S.	cohort		duration: NR	infection	(sofosbuvir +		Mean age 59 vs. 58	CVD event (acute MI, unstable,	
Fair			Followup ≥5	included in the	simeprevir,		years	angina, congestive heart	
			years	ERCHIVES	ledipasvir, or		4% vs. 4% female	failure, peripheral vascular	
			Group A: 3.7%	database	daclatasvir +/-		56% vs. 56% white;	disease, percutaneous	
			Group B: 82%	Excluded: HBV,	ribavirin;		24% vs. 24% black;	transluminalcoronary	
			Group C: 43%	HIV coinfection	paritaprevir +		3% vs. 3% Hispanic;	angioplasty, CABG, stroke):	
					ritonavir +		17% vs. 17%	3.4% (435/12,667) vs. 18.1%	
					ombitasvir +		other/unknown	(804/4,436) vs. 13.8%	
					dasabuvir +/-		Fibrosis stage: <1.25:	(2,361/17,103); A vs. C: aHR	
					ribavirin; elbasvir		23% vs. 33%; 1.26 to	0.57 (95% CI, 0.51 to 0.65); B	
					+ grazoprevir +/-		3.25: 56% vs. 50%;	vs. C: aHR 0.78 (95% CI, 0.71	
					ribavirin)		>3.25: 21% vs. 17%	to 0.85)	
					(n=12,667)		Statin use: 22% vs.	Incidence rate/1,000 person-	
					B. Pegylated		26%	years of followup: 16.3 (95%	
					IFN + ribavirin			CI, 14.7 to 18) vs. 23.5 (95%	
					(n=4,436)			CI, 21.8 to 25.3) vs. 30.4 (95%	
					C. Matched,			CI, 29.2 to 31.7); A vs. C:	
					untreated			p<0.001; B vs. C: p<0.001	
					controls				
					(n=17,103)				

Author									
year			Treatment						
Country		Dates of	duration	Inclusion					Funding
Quality	Type of study	enrollment	Followup	criteria	Intervention(s)	Ν	Population	Outcomes	source
Carrat	Prospective	Aug 2012 to	Treatment	Patients with	A. DAA regimen	6,850	Total study	A vs. B (noncirrhotics only)	French
2019 ¹⁶⁸	cohort	Dec 2015	duration: NR	chronic HCV	(sofosbuvir +		population, including	All-cause mortality: 0.8%	National
France				infection	simeprevir +/-		additional 3,045	(35/4,521) vs. 2.1% (48/2,329);	Agency for Aids
Fair			Followup:	recruited from	ribavirin;		patients with cirrhosis	aHR: 0.74 (95% CI, 0.43 to	and Viral
			median 33.4	32 hepatology	sofosbuvir +		A vs. B	1.28)	Hepatitis
			months (IQR	centers in	daclatasvir +/-		Mean age: 57 vs. 54	Liver-related mortality: 0.1%	Research;
			24.0 to 40.7	France.	ribavirin;		Female: 44% vs. 54%	(6/4,521) vs. 0.3% (6/2,329);	French
			months)	Excluded: HBV,	sofosbuvir +		Race NR	unadjusted HR: 1.33 (95% CI,	National
				HIV coinfection,	ledipasvir +/-		Fibrosis stage: F0,	0.46 to 3.84)	Agency of
				previous HCC	ribavirin;		F1, or F2: 41% vs.	HCC: 0.5% (21/4,521) vs. 0.6%	Research;
				diagnosis,	sofosbuvir +		84%	(14/2,329); AHR: 1.02 (95% CI,	French Ministry
				history of	ribavirin;		F3: 17% vs. 6%	0.40 to 2.61)	of Social Affairs
				decompensated	sofosbuvir + IFN		F4: 42% vs. 10%	Decompensated cirrhosis: 0.2%	and Health;
				cirrnosis, liver	alpha + ribavirin;		Genotype:	(7/4,521) VS. 0.2% (4/2,329);	Merck Sharp &
				transplant	sofosbuvir +		GT1: 67% VS. 64%;		Donme;
				recipient	velpatasvir +/-		GT2: 6% VS. 10%;	0.66 to 19.5)	Janssen;
					voxilaprevir,		GT3: 13% VS. 9%;		ADD VIE, Bristol-
					ritopovir		G14. 13% VS. 14%,		Nivers Squibb,
					ombitoovir 1/		G15-7. 2% VS. 5%		Roche
					dasabuvir ±/-				
					ribavirin: elbasvir				
					$\pm arazonrevir \pm /-$				
					ribavirin)				
					(n=4.521 non-				
					cirrhosis only				
					B. Untreated				
					patients				
					(n=2,329, non-				
					cirrhosis only)				

Author year			Treatment						
Country		Dates of	duration	Inclusion					Funding
Quality	Type of study	enrollment	Followup	criteria	Intervention(s)	N	Population	Outcomes	source
Li 2018 ¹⁷⁰	Retrospective	2002 to	Treatment	Adults with HCV	A. Pegylated	17,836	A vs. B vs. C	A vs. B vs. C	NR
U.S.	cohort	2016	duration: ≥28	infection	IFN + ribavirin		Mean age 54 vs. 62	HCC: 5.6% (196/3,534) vs.	
Fair			days	included in the	(n=3,534)		vs. 58 years	0.9% (50/5,834) vs. 5.0%	
			Followup: 7.4	ERCHIVES	B. DAA regimen		4% vs. 3% vs. 3%	(436/8,468)	
			years (group	database	(sofosbuvir +		female	Incidence rate/1,000 person-	
			A); 1.1 year	Excluded: HBV,	simeprevir +/-		67% vs. 51% vs. 50%	years/followup: -Total cohort:	
			(group B)	HIV coinfection;	ribavirin;		white; 17% vs. 31%	7.48 (95% CI, 6.50 to 8.61) vs.	
				HCC diagnosis	sofosbuvir +		vs. 35% black; 6% vs.	7.92 (95% CI, 6.00 to 10.45) vs.	
					ledipasvir +/-		3% vs. 6% Hispanic;	10.90 (95% CI, 9.92 to 11.97);	
					ribavirin;		11% vs. 15% vs. 9%	A vs. B: p=0.72; A vs. C:	
					sofosbuvir +		other	p<0.001	
					daclatasvir +/-		Fibrosis stage: <1.45:		
					ribavirin;		46% vs. 37% vs.		
					ombitasvir +		49%; 1.45 to 3.50:		
					paritaprevir +		41% vs. 43% vs.		
					ritonavir +		37%; >3.5: 13% vs.		
					dasabuvir +/-		20% vs. 15%		
					ribavirin)				
					(n=5,834)				
					C. No antiviral				
					treatment				
					(n=8,468)				

Author									
year			Treatment						
Country		Dates of	duration	Inclusion					Funding
Quality	Type of study	enrollment	Followup	criteria	Intervention(s)	Ν	Population	Outcomes	source
Younossi	Retrospective	October	Treatment	Treatment-naïve	A. Sofosbuvir +	706	Population with	A vs. B	Gilead
2015 ¹³⁵	cohort	2012 to	duration: 8 to	or experienced	ledipasvir		no/mild fibrosis, NR	Quality of life score, mean	
ION 1-3		June 2013	24 weeks	with chronic	(n=420)		by intervention group	change from baseline SF-36	
Multinationa			Followup: 12	HCV infection	B. Sofosbuvir +		Mean age 54 years	physical component score	
I (U.S.,			weeks post-	enrolled in ION-	ledipasvir +		33% female	(scale 0 to 100): 1.70 (SD 5.85;	
Europe)			treatment	1, 2 or 3 trials	ribavirin (n=286)		77% white	p<0.05*) vs. 1.92 (SD 6.17;	
Fair							97% U.Sbased	p<0.05*)	
							population	SF-36 mental component score	
							Treatment-naïve: 71%	(scale 0 to 100): 2.51 (SD 7.95;	
							Treatment-	p<0.05*) vs. 2.18 (SD 8.09;	
							experienced: 29%	p<0.05)	
								FACIT-F fatigue score (scale 0	
								to 52): 4.18 (SD 8.90; p<0.05)	
								vs. 4.34 (SD 9.21; p<0.05)	
								FACIT-F total score (scale 0 to	
								160): 10.27 (SD 19.57; p<0.05)	
								vs. 10.75 (SD 20.02; p<0.05)	
								CLDQ-HCV total score (scale 1	
								to 7): 0.61 (SD 0.88; p<0.05)	
								vs. 0.50 (SD 0.85; p<0.05)	
								WPAI:SHP work productivity	
								impairment score (scale 0-1): -	
								0.032 (SD 0.210; p<0.05) vs	
								0.076 (SD 0.238; p<0.05)	
								WPAI:SHP activity impairment	
								score (scale 0-1): -0.082 (SD	
								0.240; p<0.05) vs0.093 (SD	
								0.230; p<0.05)	
								SF-6D health utility score (0.2-	
								1): 0.052 (SD 0.130; p<0.05)	
								vs. 0.042 (SD 0.124; p<0.05)	

Author									
year			Treatment						
Country		Dates of	duration	Inclusion					Funding
Quality	Type of study	enrollment	Followup	criteria	Intervention(s)	N	Population	Outcomes	source
Younossi	Retrospective	July 2014 to	Treatment	Chronic HCV	A. Sofosbuvir +	1,112	Population with no	A vs. B	Gilead
2017 ¹³⁶	cohort	December	duration: 12 to	infection with no	velpatasvir		cirrhosis, NR by	Mean improvement in patient-	
ASTRAL 1-		2014	24 weeks	cirrhosis or	(n=813)		intervention group	reported outcomes (composite	
4			Followup: 12	compensated	B. Sofosbuvir +/-		Mean age 52 years	SF-36, FACIT-F, CLDQ-HCV,	
Multinationa			weeks post-	cirrhosis	velpatasvir +		41% female	WPAI:SHP; scale 0-100): 5.5	
I (U.S.,			treatment	enrolled in	ribavirin (n=299)		84% white; 6% black;	(SD NR; p>0.05*) vs. 6.1 (SD	
Canada,				ASTRAL-1, 2 or			8% Asian	NR; p>0.05*)	
Europe,				3 trials			42% U.Sbased		
Hong Kong)				(ASTRAL-4			population		
Fair				enrolled only			Treatment-naïve: 80%		
				patients with			Treatment-		
				decompensated			experienced: 20%		
				cirrhosis)					

* Within group difference from baseline

Abbreviations: aHR = adjusted hazard ratio; CABG = coronary artery bypass graft; CI = confidence interval; CLDQ-HCV = Chronic Liver Disease Questionnaire-Hepatitis C Version; CVD = cardiovascular disease; DAA = direct acting antiviral; ERCHIVES = Electronically Retrieved Cohort of HCV-Infected Veterans; FACIT-F = Functional Assessment of Chronic Illness Therapy-Fatigue; HBV = hepatitis B virus; HCC = hepatocellular carcinoma; HCV = hepatitis C virus; HR = hazard ratio; IFN = interferon; IQR = interquartile range; MI = myocardial infarction; NR = not reported; SD = standard deviation; SF-36 = Short Form 36; SF-6D = Short Form 6D; U.S. = United States; WPAI:SHP = Work Productivity and Activity Impairment Questionnaire: Specific Health Problem. Study names are not acronyms.

	Did the study attempt to enroll all (or a random sample of) patients meeting inclusion criteria, or a random sample	Were the groups comparable at baseline on key prognostic factors (e.g., by restriction or	Did the study use accurate methods for ascertaining exposures and potential	Were outcome assessors and/or data analysts blinded to the exposure being	Did the article report	Did the study perform appropriate statistical analyses on potential	Is there important differential loss to follow-up or overall high loss to follow-	Were outcomes pre-specified and defined, and ascertained using accurate	
Author year	(inception cohort)?	matching)?	confounders?	studied?	attrition?	confounders?	up?	methods?	Quality rating
Butt 2019 ¹⁶⁹	Yes	Yes	Yes	Unclear	No	Yes	Unclear	Yes	Fair
Li 2018 ¹⁷⁰	Yes	Yes	Yes	Unclear	No	Yes	Unclear	Yes	Fair
Carrat 2019 ¹⁶⁸	Yes	No	Yes	Unclear	Yes	Yes	No	Yes	Fair
Younossi 2017b ¹³⁶	Yes	NA	Yes	Unclear	No	Yes	Unclear	Yes	Fair
Younossi 2015 ¹³⁵	Yes	NA	Yes	Unclear	No	Yes	Unclear	Yes	Fair

Abbreviation: NA = not applicable.

Appendix B Table 7. Key Questions 6-8: Direct Acting Antiviral Therapy in Adolescents – Study Characteristics

Author year Country <u>Quality</u> Abdel Ghaffar 2019 ²⁰¹ Egypt <i>Fair</i>	Eligibility Age Fibrosis stage Genotype(s) HBV status Age 8 to 18 years Patients with cirrhosis excluded Genotype 4 Patients with HBV infection excluded	Study Recruitment Dates December 2016 to February 2018	Sample Size 40	Baseline Characteristics Age, Sex, Race/ethnicity, Fibrosis stage/ METAVIR score (mean/median if breakdown is NR), Genotype breakdown Mean age 12 years (45% <12 years) 38% female Race NR Fibrosis stage F0: 35%; F1: 38%; F2 and F3: 15% Genotype 4: 100% (mixed 4 and 1a: 13%; mixed 4 and 1b: 15%) Treatment naïve: 100%	Loss to Followup 3% (1/40)	Definition of SVR HCV RNA <lloq< th=""></lloq<>
Balistreri 2017 ¹⁷⁵ and Younossi 2018 ¹⁷² Australia, U.K., U.S. <i>Fair</i>	Age 12 to <18 years Patients with cirrhosis permitted; liver biopsy not required Genotype 1 Patients with HBV infection excluded	November 2014 to October 2015	100	Mean age 15 years 63% female 90% white; 7% black; 2% Asian; 1% NR Fibrosis stage F0-F3: 42%; F4:1%; NR/unknown: 57% Genotype 1a: 81%; 1b: 19% Treatment naïve: 80% Treatment experienced 20% (prior treatment unclear; presumably IFN or pegylated IFN + ribavirin)	2% (2/100)	HCV RNA <15 IU/mL
EI-Karaksy 2018 ²⁰² Egypt <i>Fair</i>	Age 12 to <18 years Fibrosis stage NR; fibrosis stage assessed by FibroScan Genotype 4 Patients with HBV infection excluded	NR	40	Mean age 14 years 35% female Race NR Fibrosis stage F0: 55%; F0 and F1: 13%; F1: 13%; F1 and F2: 5%; F3: 10%; F4: 5% (>100% due to rounding) Genotype 4: 100% Treatment-naïve: 75% Treatment-experienced: 25% (IFN +/- ribavirin)	0% (0/40)	Negative HCV RNA
Jonas 2019 ¹⁷¹ DORA Multinational <i>Fair</i>	Age 12 to <18 years Patients with decompensated cirrhosis excluded; compensated cirrhosis allowed Genotype 1 to 6 Patients with HBV excluded	March 2017 to present (study is ongoing)	48	Median age 14 years 55% female 75% white; 9% black; 13% Asian; 4% mixed race Fibrosis stage F0-F1: 96%; F2: 2%; F3: 2% Genotype 1a: 51%; 1b: 28%; 2: 6%; 3: 9%; 4: 6%; no genotype 5 or 6 enrolled HIV coinfection: 4% Treatment-naïve: 77% Treatment-experienced: 23% (pegylated IFN + ribavirin)	2% (1/48; patient was not treated and excluded from analysis)	HCV RNA <15 IU/mL

Appendix B Table 7. Key Questions 6-8: Direct Acting Antiviral Therapy in Adolescents – Study Characteristics

Author year Country <i>Quality</i> Leung 2018 ²⁰³ ZIRCON Multinational <i>Fair</i>	Eligibility Age Fibrosis stage Genotype(s) HBV status Age 12 to 17 years Patients with cirrhosis permitted, based on liver biopsy, FibroTest or FibroScan Genotype 1 or 4 Patients with HBV infection excluded	Study Recruitment Dates November 2015 to July 2016	Sample Size 38	Baseline Characteristics Age, Sex, Race/ethnicity, Fibrosis stage/ METAVIR score (mean/median if breakdown is NR), Genotype breakdown Median age 15 years 66% female 76% white; 13% black; 8% Asian; 3% mixed race Fibrosis stage (30/38 patients): F0 and F1: 90%; F2: 3%; F3: 3%; F4: 3% Genotype 1a: 42%; 1b: 40%; 4: 18% Treatment naïve: 66% Treatment experienced: 34% (IFN +/- ribavirin)	Loss to Followup 0% (0/38)	Definition of SVR HCV RNA <lloq< th=""></lloq<>
Wirth 2017 ¹⁷³ and Younossi 2018 ¹⁷⁴ Australia, Belgium, Germany, Italy, New Zealand, Russia, U.K., U.S. <i>Fair</i>	Age 12 to <18 years Patients with cirrhosis permitted; liver biopsy not required Genotype 2 or 3 Patients with HBV infection excluded	October 2014 to June 2016	52	Median age 15 years 40% female 90% white; 4% black; 2% Asian; 2% Hawaiian/Pacific Islander; 2% other Fibrosis stage NR; 40% no cirrhosis; 60% cirrhosis presence unknown Genotype 2: 25% Genotype 3: 75% Treatment-naive: 83% Treatment-experienced: 17% (prior treatment unclear; 6% prior nonresponder; 2% prior relapse; 1% IFN intolerant) PedsQL-4.0-SF-15 score (post-hoc analysis; n=50): 73.54 (SD 2.16)	2% (1/52)	HCV RNA <15 IU/mL
Yakoot 2018 ¹⁷⁶ Egypt <i>Good</i>	Age 12 to 17 years Fibrosis stage NR; FibroScan >12.5 kPa and/or APRI >2.0 excluded Genotype 4 Patients with HBV infection excluded	February 2017 to NR	30	Mean age 13 years 43% female Race NR Fibrosis stage F0: 17%; F1: 53%; F2: 27%; F3: 3% Genotype 4: 100% Treatment naïve: 73% Treatment experienced: 27% (prior treatment unclear)	3% (1/30)	HCV RNA <lloq< td=""></lloq<>

Abbreviations: APRI = aspartate amino transferase to platelet ratio index; HBV = hepatitis B virus; HCV = hepatitis C virus; IFN = interferon; LLOQ = lower limit of quantification; NR = not reported; PedsQL = Pediatric Quality of Life Inventory; RNA = ribonucleic acid; SD = standard deviation; SF = short form; SVR = sustained virologic response; U.K. = United Kingdom; U.S. = United States. Study names are not acronyms.

Appendix B Table 8. Key Questions 6-8: Direct Acting Antiviral Therapy in Adolescents – Effectiveness and Harms

Author year	Treatment Regimen (1x/day	Traction to Duration	Efficiency				
Country Quality	noted)	and Assessments	Results	Subgroup Efficacy Results	Clinical Outcomes	Adverse Events	Funding Source
Abdel Ghaffar 2019 ²⁰¹ Egypt <i>Fair</i>	Sofosbuvir 200- 400 mg + daclatasvir 30-60 mg	Treatment duration: 12 weeks Timing of assessments: 12 weeks post treatment	SVR: 98% (39/40)	NR	NR	Any adverse event: NR Serious adverse events: NR Withdrawal due to adverse events: NR Headache: 3% (1/40) Fatigue: 5% (2/40)	The Egyptian Cure Bank non-governmental organization; Society of Friends of Liver Patients in the Arab World
Balistreri 2017 ¹⁷⁵ and Younossi 2018 ¹⁷² Australia, U.K., U.S. <i>Fair</i>	Ledipasvir 90 mg + sofosbuvir 400 mg	Treatment duration: 12 weeks Timing of assessments: 12 weeks post treatment	SVR: 98% (98/100)	Treatment-naïve: 98% (78/80) Treatment-experienced: 100% (20/20)	Mortality: 0% (0/100) PedsQL-4.0-SF-15 Score, mean change from baseline at post- treatment week 24 (scale 0-100, positive mean change = improvement in quality of life): Physical functioning: caregiver report: 2.14, p=0.49, self-report: - 0.49, p=0.97 Emotional functioning: caregiver report 9.32, p<0.001; self-report 3.66, p=0.04 Social functioning: caregiver report 4.79, p=0.18; self-report 3.02 p=0.33 School functioning: caregiver report 4.79, p=0.18; self-report 3.02 p=0.33 Total score: caregiver report: 5.25, p=0.009; self-report: 1.89, p=0.12	Any adverse event: 71% (71/100) Serious adverse events: 0% (0/100) Withdrawals due to adverse events: 0% (0/100) Headache: 27% (27/100) Fatigue: 13% (13/100) Nausea: 11% (11/100) Vomiting: 11% (11/100)	Gilead

Author year Country <i>Quality</i>	Treatment Regimen (1x/day unless otherwise noted)	Treatment Duration and Assessments	Efficacy Results	Subgroup Efficacy Results	Clinical Outcomes	Adverse Events	Funding Source
El-Karaksy 2018 ²⁰² Egypt <i>Fair</i>	Ledipasvir 90 mg + sofosbuvir 400 mg	Treatment duration: 12 weeks Timing of assessments: 12 weeks post treatment	SVR: 100% (40/40)	NR	NR	Headache: 48% (19/40) Fatigue: 53% (21/40) Nausea: 28% (11/40) Diarrhea: 23% (9/40) Insomnia: 23% (9/40)	NR; described as "treatment provided by charity"
Jonas 2019 ¹⁷¹ DORA Multinational <i>Fair</i>	Glecaprevir 300 mg + pibrentasvir 120 mg	Treatment duration: 8 to 16 weeks (94% of study population treated for 8 weeks) Timing of assessments: 12 weeks post treatment	SVR: 100% (47/47)	NR	PedsQL total score, mean change from baseline (N=44): 2.3 (SD 7.7); p=NR	Any adverse event: 87% (41/47) Serious adverse events: 0% Withdrawal due to adverse events: 0% Headache: 17% (8/47) Fatigue: 11% (5/47)	AbbVie
Leung 2018 ²⁰³ ZIRCON Multinational <i>Fair</i>	Ombitasvir 25 mg + paritaprevir 150 mg + ritonavir 100 mg + dasabuvir 250 mg 2x/day + weight-based ribavirin	Treatment duration: 12 weeks Timing of assessments: 12 weeks post treatment	SVR: 100% (38/38)	Genotype 1a: 100% (16/16) Genotype 1b: 100% (15/15) Genotype 4: 100% (7/7) Treatment naïve: 100% (25/25) Treatment experienced: 100% (13/13)	NR	Any adverse event: 84% (32/38) Serious adverse events: 0% (0/38) Withdrawal due to adverse events: 0% (0/38) Headache: 21% (8/38) Fatigue: 18% (7/38)	AbbVie
Wirth 2017 ¹⁷³ and Younossi 2018 ¹⁷⁴ Australia, Belgium, Germany, Italy, New Zealand, Russia, U.K., U.S. <i>Fair</i>	Sofosbuvir 400 mg + weight- based ribavirin	Treatment duration: 12 (genotype 2) or 24 (genotype 3) weeks Timing of assessments: 12 weeks post treatment	SVR: 98% (51/52)	Genotype 2: 100% (13/13) Genotype 3: 97% (38/39)	Mortality: 0% (0/52) PedsQL-4.0-SF-15 Score, mean change from baseline at post- treatment week 24 (positive mean change=improvement in quality of life): 7.26 (SD 2.99); p=0.01	Any adverse event: 81% (41/52) Serious adverse events: 2% (1/52) Withdrawal due to adverse events: 0% (0/52) Headache: 23% (12/52) Fatigue: 12% (6/52) Nausea: 27% (14/52) Diarrhea: 6% (3/52)	Gilead

Appendix B Table 8. Key Questions 6-8: Direct Acting Antiviral Therapy in Adolescents – Effectiveness and Harms

Author year	Treatment Regimen (1x/day						
Country	unless otherwise	Treatment Duration	Efficacy				
Quality	noted)	and Assessments	Results	Subgroup Efficacy Results	Clinical Outcomes	Adverse Events	Funding Source
Yakoot 2018 ¹⁷⁶	Weight-based	Treatment duration:	SVR: 97%	NR	Mortality: 0% (0/30)	Any adverse event: 27%	NR
Egypt	sofosbuvir +	12 weeks	(29/30)			(8/30)	
Good	daclatasvir					Serious adverse events:	
		Timing of				0% (0/30)	
		assessments: 12				Withdrawal due to	
		weeks post treatment				adverse events: 0%	
						(0/30)	
						Headache: 10% (3/30)	
						Fatigue: 13% (4/30)	
						Nausea: 10% (3/30)	

Abbreviations: NR = not reported; PedsQL = Pediatric Quality of Life Inventory; SD = standard deviation; SF = short form; SVR = sustained virologic response; U.K. = United Kingdom; U.S. = United States. Study names are not acronyms.

Author year	Single or multi- arm study?	Non- randomized studies: Enrolled all (or a random sample of) patients meeting inclusion criteria?	Randomized studies: Randomization adequate?	Randomized studies: Allocation concealment adequate?	Groups similar at base- line?	Eligibility criteria specified?	Primary outcome pre- specified and reported?	Outcome assessors masked?	Care provider masked?	Patient masked?	Attrition and with- drawals reported?	Loss to followup: differential (>10%)/ high (>20%)?	Analyze people in the groups in which they were random- ized?	Quality
Abdel Ghaffar 2019 ²⁰¹	Single	Unclear	NA	NA	NA	Yes	Yes	NA	No	No	Yes	No	Yes	Fair
Balistreri 2017 ¹⁷⁵	Single	Unclear	NA	NA	NA	Yes	Yes	NA	No	No	Yes	No	Yes	Fair
El-Karaksy 2018 ²⁰²	Single	Unclear	NA	NA	NA	Yes	Yes	NA	No	No	Yes	No	Yes	Fair
Jonas 2019 ¹⁷¹	Single	Unclear	NA	NA	NA	Yes	Yes	NA	No	No	Yes	No	Yes	Fair
Leung 2018 ²⁰³	Single	Unclear	NA	NA	NA	Yes	Yes	NA	No	No	Yes	No	Yes	Fair
Wirth 2017 ¹⁷³	Single	Unclear	NA	NA	NA	Yes	Yes	NA	No	No	Yes	No	Yes	Fair
Yakoot 2018176	Single	Yes	NA	NA	NA	Yes	Yes	NA	No	No	Yes	No	Yes	Good

Abbreviation: NA = not applicable.

Appendix B Table 10. Key Questions 6-8: Direct Acting Antiviral Therapy in Adults – Study Characteristics

	Eligibility					
Authorizant	Age	Study				
Author year		Desmuitment			1	
Country	Genotype(s)	Recruitment	Commis Cine	Deceline Characteristics	LOSS to	Definition of CVD
Quality	HBV status	Dates	Sample Size	Baseline Characteristics	Followup	Definition of SVR
Abergel 2016a ¹⁴²	Adults>18	March 2014 to	22 (treatment-naïve	Mean age 52 years	0% (0/22)	HCV RNA level <15
France	Patients with cirrhosis	November 2014	population only)	50% female		IU/mL
Fair	were eligible for inclusion,			86% white; 14% black		
	based on liver biopsy,			Fibrosis stage NR; cirrhosis: 5%		
	FibroScan >12.5 kPa, or			Genotype 4: 100%		
	FibroTest >0.75 + APRI			Treatment-naïve: 100%		
	>2					
	Genotype 4					
	Treatment-naïve arm only					
	Patients with HBV					
	infection excluded					
Abergel 2016b ¹⁴¹	Adults>18	March 2014 to	21 (treatment-naïve	Mean age 61 years	0% (0/21)	HCV RNA level <15
France	Patients with cirrhosis	June 2014	population only)	48% female		IU/mL
Good	were eligible for inclusion,			100% white		
	based on liver biopsy,			Fibrosis stage NR; cirrhosis: 14%		
	FibroScan >12.5 kPa, or			Genotype 5: 100%		
	FibroTest >0.75 + APRI			Treatment-naïve: 100%		
	>2					
	Genotype 5					
	Treatment-naïve arm only					
	Patients with HBV					
	infection excluded					

A	Eligibility Age	Otracha				
Author year Country <i>Quality</i>	Genotype(s) HBV status	Recruitment Dates	Sample Size	Baseline Characteristics	Loss to Followup	Definition of SVR
Afdahl 2014 ¹⁸⁵ ION-1 U.S. and Europe <i>Fair</i>	Age >18 years 20% of population could have cirrhosis based on liver biopsy, Fibroscan >12.5kPa, or FibroTest >0.75 and APRI >2 Genotype 1 Patients with HBV infection excluded	October 2012 to May 2013	431 A=214 B=217	A vs. B <u>12-week intervention group (n=214)</u> Mean age 52 vs. 52 years 41% vs. 41% female 87% vs. 87% white; 11% vs. 12% black; <1% vs. 0% Asian; 1% vs. 1% other Fibrosis stage NR; cirrhosis: 16% vs. 15% Genotype 1a: 67%; 1b: 31%, Other 2% Treatment-naive: 100% vs. 100% <u>24-week intervention group (n=217)</u> Mean age 53 vs. 53 years 36% vs. 45% female 82% vs. 84% white; 15% 12% black; 2% vs. 2% Asian; 1% vs. 1% other Fibrosis stage NR; cirrhosis: 15% vs. 17% Genotype 1a: 67% vs. 66%; 1b: 31% vs. 33%, Other 1% vs. 1% Treatment-naive: 100% vs. 100%	0.9% (4/431)	HCV RNA <25 IU/mL
Ahmed 2018 ¹⁹⁵ Egypt <i>Fair</i>	Age ≥18 years Fibrosis/cirrhosis NR; Child-Pugh >8 excluded Genotype 4 Treatment-naïve HBV status NR	January 2015 to NR	100	Mean age 51 years 35% female Race/ethnicity NR Fibrosis stage NR Genotype 4: 100% Treatment-naïve: 100%	0% (0/100)	HCV RNA <15 IU/mL
Andreone 2014 ¹⁸⁶ PEARL-II Austria, Belgium, Italy, The Netherlands, Portugal, Puerto Rico, Sweden, Switzerland, U.S. <i>Fair</i>	Age 18 to 70 years Fibrosis stage NR; patients were required to have no cirrhosis Genotype 1b Prior failure of pegylated IFN + ribavirin treatment Patients with HBV infection excluded	August 2012 to January 2014	186 A=91 B=88	A vs. B Mean age 54 vs. 54 years 40% vs. 50% female 91% vs. 92% white; 6% vs. 3% black; 2% vs. 4% Hispanic Fibrosis stage F0 and F1: 64% vs. 70%; F2: 22% vs. 14%; F3: 13% vs. 14% Genotype 1b: 100% vs. 100% Treatment-naïve: 0% vs. 0% Treatment-experienced: 100% vs. 100% (pegylated IFN + ribavirin)	0.5% (1/186)	HCV RNA <25 IU/mL

	Eligibility					
Author year Country Quality	Age Fibrosis stage Genotype(s) HBV status	Study Recruitment Dates	Sample Size	Baseline Characteristics	Loss to Followup	Definition of SVR
Asselah 2018 ¹⁹⁶ SURVEYOR II Part 4, Multinational (Asia, Europe, U.S. [specific countries NR]) <i>Fair</i>	Age >18 years No cirrhosis based on liver biopsy, FibroScan <12.5 kPa or FibroTest ≤0.48 and APRI <1 Genotype 2, 4, 5 or 6 Treatment naïve or experienced Patients with HBV	October 2014 to October 2016	203 (8-week intervention groups only)	Mean age 52 years 52% female 75% white; 10% black; 11% Asian Fibrosis stage F0 and F1: 84%; F2: 6%; F3: 10% Genotype 2: 71%; 4: 23%; 5: 1%; 6: 5% Treatment-naïve: 87% Treatment-experienced (IFN or peg IFN, with ribavirin, with or without sofosbuvir):	0.5% (1/203)	HCV RNA <lloq< td=""></lloq<>
Asselah 2019 ¹⁴³ ENDURANCE-5 Multinational (Australia, Belgium, Canada, France, New Zealand, Singapore, South Africa, Vietnam, U.S.) <i>Fair</i>	Age ≥18 years Cirrhosis allowed based on liver biopsy, FibroScan <12.5 kPa or FibroTest ≤0.48 Genotype 5 Treatment naïve or experienced Patients with HBV excluded	January 2017 to December 2017	23	Mean age 68 years 57% female 91% white; 4% Asian, 4% black Fibrosis stage F0 and F1: 74%; F2: 13%; F3: 0%; F4 (cirrhosis): 13% Genotype 5: 100% Treatment-naïve: 83% Treatment-experienced (IFN or peg IFN): 17%	0% (0/23)	HCV RNA <15 IU/mL
Asselah 2019 ¹⁴³ ENDURANCE-6 (same publication as ENDURANCE-5) <i>Fair</i>	Age ≥18 years Cirrhosis allowed based on liver biopsy, FibroScan <12.5 kPa or FibroTest ≤0.48 Genotype 6 Treatment naïve or experienced Patients with HBV excluded	See Asselah 2019 ENDURANCE-5	61	Mean age 54 years 52% female 7% white; 92% Asian, 0% black; 1% other Fibrosis stage F0 and F1: 74%; F2: 2%; F3: 15%; F4 (cirrhosis): 10% Genotype 6: 100% Treatment-naïve: 93% Treatment-experienced (IFN or peg IFN): 7%	0% (0/61)	See Asselah 2019 ENDURANCE-5
Brown 2018 ¹⁴⁴ C-SCAPE (Genotype 4 only) Multinational (Australia, Belgium, France, Israel, Spain, U.K., U.S.) <i>Fair</i>	Age ≥18 years No cirrhosis based on liver biopsy, FibroScan <12.5 kPa or FibroTest ≤0.48 Genotype 2, 4, 5 or 6 Treatment-naive Patients with HBV excluded	October 2013 to December 2014	20 (Genotype 4 only; total population n=38)	Total population (genotypes 2, 4, 5, 6) A vs. B Mean age 52 vs. 53 years 58% vs. 37% female 74% vs. 68% white; 26% vs. 32% other race Fibrosis stage F0 to F2: 79% vs. 90%; F3: 21% vs. 5%; unknown: 0% vs. 5% Treatment-naïve: 100% vs. 100%	0% (0/20)	HCV RNA <25 IU/mL

	Eligibility					
	Age					
Author year	Fibrosis stage	Study				
Country	Genotype(s)	Recruitment			Loss to	
Quality	HBV status	Dates	Sample Size	Baseline Characteristics	Followup	Definition of SVR
Chayama 2018 ¹⁹⁷	Age ≥18 years	February 2016 to	129	Median age 64 years	0.8% (1/129)	HCV RNA <15 IU/mL
CERTAIN-1 (Arm A only)	No cirrhosis based on	June 2016		64% female		
Japan	liver biopsy or FibroScan			Race/ethnicity NR		
Fair	<12.5 kPa or FibroTest			Fibrosis stage NR		
	>0.73 and APRI ≤2			Genotype 1: 100%		
	Genotype 1			Treatment-naïve: 73%		
	Treatment naïve or			Treatment-experienced (IFN with/without		
	experienced			ribavirin): 27%		
	Patients with HBV					
	excluded					
Chuang 2016 ¹⁴⁵	Age ≥20 years	December 2013	85	Mean age 55 years	0% (0/85)	HCV RNA <25 IU/mL
Taiwan	≤20% enrolled participants	to March 2014		58% female		
Fair	could meet cirrhosis			100% Asian		
	criteria, based on Metavir			Fibrosis stage: NR		
	score 4, Ishak score ≥5,			Genotype: 1: 1%; 1a: 12%; 1b: 87%		
	or Fibroscan >12.5 kPa			Cirrhosis: 11%		
	Genotype 1			Treatment-naïve: 100%		
	Patients with HBV					
	infection excluded					
Dore 2016 ¹³⁷	Age 18 to 65 years	March to	309	A vs. B vs. C vs. D vs. E	0% (0/311)	HCV RNA <25 IU/mL
MALACHITE-1	No cirrhosis, based on	November 2014	<u>Genotype 1a</u>	Mean age 46 vs. 45 vs. 46 vs. 47 vs. 46		
Australia, Canada, Europe,	FibroTest ≤0.72 and APRI		A=69	years		
South America	≤2; or FibroScan <9.6		B=34	39% vs. 59% vs. 55% vs. 52% vs. 59%		
Good	kPa; or liver biopsy within			female		
	24 months		Genotype 1b	17% vs. 9% vs. 14% vs. 18% vs. 7%		
	Genotype 1		C=84	Hispanic/Latino; other race/ethnicity NR		
	Treatment-naïve		D=83	Fibrosis stage F0 and F1: 72% vs. 71%		
	Patients with HBV		E=41	vs. 83% vs. 72% vs. 76%; F2: 18% vs.		
	infection excluded			21% vs. 8% vs. 13% vs. 10%; F3: 10% vs.		
				9% vs. 8% vs. 14% vs. 15%		
				Treatment-naive: 100% across all groups		

	Eligibility					
	Age					
Author year	Fibrosis stage	Study				
Country	Genotype(s)	Recruitment			Loss to	
Quality	HBV status	Dates	Sample Size	Baseline Characteristics	Followup	Definition of SVR
Dore 2016 ¹³⁷	Age 18 to 65 years	March to	148	A vs. B	0% (0/148)	HCV RNA <25 IU/mL
MALACHITE-2	No cirrhosis, based on	November 2014	A=101	Mean age 47 vs. 45		
Australia, Canada, Europe,	FibroTest ≤0.72 and APRI		B=47	46% vs. 40% female		
South America	≤2; or FibroScan <9.6			100% vs. 100% white		
Good	kPa; or liver biopsy within			12% vs. 4% Hispanic/Latino		
	24 months			Fibrosis F0 and F1: 78% vs. 68%; F2:		
	Genotype 1			17% vs. 23%; ≥F3: 5% vs. 9%		
	I reatment-experienced			Treatment-naive: 0%		
	(pegylated IFN + ribavirin)			Treatment-experienced: 100%		
	Patients with HBV			(peginterferon and ribavirin)		
Exercise 2045 (Dent A)146	Infection excluded	A	077		00((0/077)	
Everson 2015 (Part A)	Age ≥18 years	August 2013 to	3//	A VS. B VS. C VS. D VS. E VS. F	0% (0/377)	HCV RNA <lloq 12<="" td=""></lloq>
U.S.	FIDROSIS Stage: INR;	August 2014	A=27 P_29	Iviean age 49 vs. 49 vs. 52 vs. 50 vs. 48		weeks post-treatment
9000	participants could not		D=20 C=27	1200×10^{-10}		
	have cirriosis, based on.		D = 28	40% VS. 39% VS. 33% VS. 37% VS. 20%		
	of scrooping: or EibroTost		E-23	85% vs 80% vs 81% vs 96% vs 83%		
	<0.48 and AST platelet		E-20 F-22	vs. 73% white: 15% vs. 4% vs. 15% vs.		
	index <1 during screening:			0% vs 9% vs 5% black: 0% vs 7% vs		
	or Fibroscan <12.5 kPa			4% vs. 4% vs. 9% vs. 23% other		
	within 6 months of			Fibrosis/METAVIR score: NR		
	baseline			Groups A & B: Genotype 1; Groups C &		
	Genotype 1-6			D: Genotype 3; Groups E & F: Genotypes		
	Treatment naïve			2; 4 to 6		
	Patients with HBV			Treatment naive: 100% across all groups		
	infection excluded					
Feld 2014 ¹⁸⁷	Adults >18	November 2012	477	Mean age 49	0.4% (2/477)	HCV RNA level <25
SAPPHIRE-1	Fibrosis Stage NR	to May 2013		43% female		IU/mL
Australia, New Zealand;	Genotype 1			91% white; 6% black; 4% other		
Austria, France, Germany,	Treatment naïve or			METAVIR score F0 or F1: 77%; F2: 15%;		
Hungary, Great Britain,	experienced			F3: 8.4%		
Italy, Spain, Sweden,	Patients with HBV			Genotype 1a: 69% Genotype 1b: 32%		
Switzerland; Canada, U.S.	infection excluded			Treatment-naive: 68%		
Good				Treatment-experienced: 32% (9.0%		
				protease inhibitor, peginterferon, and		
				Iribavirin; 20% pegylated IFN and ribavirin;		
				3.7% nonpegylated IFN with or without		
				ribavirin)		

	Eligibility					
Author year Country <i>Quality</i>	Age Fibrosis stage Genotype(s) HBV status	Study Recruitment Dates	Sample Size	Baseline Characteristics	Loss to Followup	Definition of SVR
Feld 2015 ¹³⁹ ASTRAL-1 U.S., Canada, Europe, Hong Kong <i>Good</i>	Age ≥18 years Fibrosis stage NR; up to 20% could have cirrhosis based on: liver biopsy (Metavir stage 4 or Ishak score 5 or 6), FibroTest score >0.75, AST:platelet ratio >2, or FibroScan >12.5 kPa) Genotype 1, 2, 4, 5, 6 Treatment-naive or experienced Patients with HBV infection excluded	July 2014 to December 2014	706 A=624 B=116	A vs. B Mean age 54 vs. 53 years 60% vs. 59% female 79% vs. 78% white; 8% vs. 9% black; 10% vs. 9% Asian; 2% vs. 3% other Fibrosis stage/METAVIR score NR Genotype 1a: 34% vs. 40%; 1b: 19% vs. 16%; 2: 17% vs. 18%; 4: 19% vs. 19%; 5: 6% vs. 0%; 6: 7% vs. 7% Compensated cirrhosis: 19% vs. 18% Treatment-naive: 72% vs. 68% Treatment-experienced: 28% vs. 32% (5% vs. 9% protease inhibitor, peginterferon, and ribavirin; 21% vs. 20% pegylated IFN and ribavirin; 3% vs. 4% nonpegylated IFN with or without ribavirin)	0.1% (1/706)	HCV RNA level <15 IU/mL at 12 weeks post-treatment
Ferenci 2014 ¹⁸⁸ PEARL III Austria, Belgium, Hungary, Israel, Italy, Poland, Portugal, Romania, Russia, Spain, U.S. <i>Good</i> Same publication as PEARL IV	Age 18 to 70 years No cirrhosis based on liver biopsy with 24 months, Fibro Scan (NR) or FibroTest (NR) Genotype 1b Patients with HBV infection excluded	NR	419 A=209 B=210	A vs. B Mean age 49 vs. 48 years 59% vs. 49% female 94% vs. 94% white; 5% vs. 5% black; 1% vs. 1% other; 2% vs. 1% Hispanic Fibrosis score F0 or F1: 68% vs. 71%; F2: 23% vs. 18%; F3: 10% vs. 11% Treatment-naïve: 100% vs. 100%	0% (0/419)	HCV RNA <25 IU/mL
Ferenci 2014 ¹⁸⁸ PEARL IV Canada, U.K., U.S. <i>Good</i> Same publication as PEARL III	Age 18 to 70 years No cirrhosis based on liver biopsy with 24 months, Fibro Scan (NR) or FibroTest (NR) Genotype 1a Treatment naive Patients with HBV infection excluded	NR	305 A=205 B=100	A vs. B Mean age 51 vs. 52 years 37% vs. 30% female 83% vs. 86% white; 13% vs. 10% black; vs. 4% 4% other; 11% vs. 11% Hispanic Fibrosis score F0 and F1: 64% vs. 63%; F2: 17% vs. 21%; F3: 19% vs. 16% Treatment-naïve: 100% vs. 100%	1% (3/305)	HCV RNA <25 IU/mL

	Eligibility					
	Age					
Author year	Fibrosis stage	Study				
Country	Genotype(s)	Recruitment			Loss to	
Quality	HBV status	Dates	Sample Size	Baseline Characteristics	Followup	Definition of SVR
Foster 2015 ¹⁴⁷	Age ≥18 years	October 2014 to	269	A vs. B	0.4% (1/269)	HCV RNA <15 IU/mL
ASTRAL-2	Fibrosis stage NR; up to	December 2014	A=134	Mean age 57 vs. 57 years		
U.S.	20% could have		B=132	36% vs. 45% female		
Fair	compensated cirrhosis			93% vs. 84% white; 4% vs. 9% black; 1%		
	based on: liver biopsy			vs. 4% Asian; 2% vs. 3% other		
	(Metavir stage 4 or Ishak			Fibrosis stage NR; 14% vs. 14% cirrhosis		
	score 5 or 6), FibroTest			Genotype 2: 100% vs. 100%		
	score >0.75, AST:platelet			Treatment-naïve: 86% vs. 85%		
	ratio >2, or FibroScan			Treatment experienced: 14% vs. 15%		
	>12.5 kPa)			(IFN-containing regimen)		
	Genotype 2					
	Patients with HBV					
	infection excluded					
Foster 2015 ¹⁴⁷	Age ≥18 years	Same as Foster	558	A vs. B	1.4% (4/280)	Same as Foster 2015
ASTRAL-3	Fibrosis stage NR; up to	2015 ASTRAL-2	A=278	Mean age 49 vs. 50 years		ASTRAL-2
U.S.	20% could have		B=280	39% vs. 37% female		
Fair	compensated cirrhosis			90% vs. 87% white; 1% vs. <1% black;		
	based on: liver biopsy			8% vs. 11% Asian; <1% vs. 2% other		
Same publication as	(Metavir stage 4 or Ishak			FIDROSIS Stage NR; 29% VS. 30% CITTOSIS		
ASTRAL-2	score 5 or 6), Fibro l est			Genotype 3: 100% vs. 100%		
	score >0.75, AST:platelet			Treatment experienced, 26% vo. 26%		
	12.5 kPa			(IEN containing regimen)		
	> 12.5 KPd					
	Betients with HBV					
	infection excluded					
Gane 2015 ¹⁴⁸	Age >18 years	April 2013 to	25	Mean age 51 years	0% (0/25)	HCV RNA <15 II I/ml
New Zealand (Genotype 6	I In to 40% of enrolled	October 2014	20	36% female	070 (0/20)	
subset)	patients could have			16% white: 84% Asian		
Fair	cirrhosis diagnosis based			Fibrosis stage NR		
	on liver biopsy. Fibroscan			Cirrhosis: 8%		
	>12.5 kPa, or FibroTest			Genotype 6c-1: 68%; 6a or 6b: 32%		
	>0.75 and APRI >2			Treatment-naïve: 92%		
	Genotype 6			Treatment-experienced: 8% (previous		
	Patients with HBV			treatment not described)		
	infection excluded					

	Eligibility						
	Age						
Author year	Fibrosis stage	Study					
Country	Genotype(s)	Recruitment				Loss to	
Quality	HBV status	Dates	Sample	Size	Baseline Characteristics	Followup	Definition of SVR
Grebely 2018 ¹⁵⁰	Age ≥18 years	March 2016 to	103		Mean age 48 years	2% (2/103)	HCV RNA <lloq< th=""></lloq<>
SIMPLIFY	Cirrhosis allowed, based	October 2016			28% female		
Multinational (Australia.	on Fibroscan >14.6 kPa				Race/ethnicity NR		
Canada, New Zealand,	Genotype 1 to 6				Fibrosis stage F0 and F1: 61%: F2 and		
Norway Switzerland UK	Treatment-naïve (DAA				F3: 28%: F4 (cirrhosis): 9%		
	only: prior IFN treatment				Genotype 1a: 34%: 1b: 1%: 2: 5%: 3:		
Fair	NR)				58% 4.2%		
	IVDU within 6 months of				No IVDLL in last 30 days: 26% less than		
	study entry				daily IVDLL in last 30 days: 48% at least		
	Patients with HBV				daily IVDU in the last 30 days: 26%		
	excluded				Injection drugs used in the last 30 days.		
	o, loid do d				55% heroin 13% cocaine 30%		
					methamphetamine 21% other opioids 7%		
					other drugs		
					History of opioid substitution therapy: 82%		
Grebely 2018 ¹⁴⁹	Age >18 years	June 2016 to	87		Mean age 48 years	1% (1/87)	HCV RNA <i loq<="" td=""></i>
D3FFAT	Cirrhosis allowed based	February 2017	0.		23% female	.,	
Multinational (Australia	on FibroScan >14.6 kPa				Race/ethnicity NR		
Canada, France, New	or FIB-4 >3.25				Fibrosis stage F0 and F1: 77%: F2 and		
Zealand, Norway,	Genotype 1				F3: 13%: F4 (cirrhosis): 8%		
Switzerland)	Treatment naive				Genotype 1a: 90%: 1b: 10%		
Fair	IVDU within 6 months of				Treatment-naïve: 100%		
	study entry or use of				IVDU in last 6 months: 61%		
	opioid substitution therapy				Non-IVDU in last 6 months: 43%		
	Patients with HBV				History of opioid substitution therapy: 85%		
	excluded						
Hezode 2015 ¹⁸⁹	Age 18 to 70 years	August 2012 to	42		Mean age 44 years	0% (0/42)	HCV RNA <25 IU/mL
PEARL I (Treatment-naïve	No cirrhosis based on	March 2014			33% female		
population)	liver biopsy in the past 24				Race/ethnicity NR: 86% European: 14%		
France, Hungary, Italy,	months or FibroTest ≤0.72				North American		
Poland, Romania, Spain,	or APRI ≤2 or FibroScan				Fibrosis stage F0 and F1: 79%; F2: 14%;		
Turkey, U.S.	<9.6 kPa				F3: 7%		
Good	Genotype 4				Genotype 4: 100%		
	Patients with HBV				Treatment-naïve: 100%		
See also Lawitz 2015 ¹⁵⁵	infection excluded						
(PEARL I - Genotype 1b)							

	Eligibility					
Author year	Age Eibrosis stago	Study				
Country	Genotype(s)	Recruitment			Loss to	
Quality	HBV status	Dates	Sample Size	Baseline Characteristics	Followup	Definition of SVR
Hezode 2015 ¹⁸⁹ PEARL I (Treatment experienced population) France, Hungary, Italy, Poland, Romania, Spain, Turkey, U.S. <i>Good</i> See also Lawitz 2015 ¹⁵⁵ (PEARL 1 - Genotype 1b)	Same as Hezode 2015 (Treatment naïve population)	Same as Hezode 2015 (Treatment naïve population)	49	Mean age 51 years 26% female Race/ethnicity NR; 86% European; 14% North American Fibrosis stage F0 and F1: 67%; F2: 22%; F3: 10% Genotype 4: 100% Treatment-naïve: 0%	0% (0/49)	Same as Hezode 2015 (Treatment naïve population)
Kowdley 2014a ¹⁹⁰ ION-3 U.S. <i>Fair</i>	Age ≥18 years No cirrhosis based on liver biopsy in the past 24 months or FibroTest ≤0.48 and APRI ≤1 Genotype 1 Patients with HBV infection excluded	May 2013 to June 2013	431	8-week intervention group (n=215) Mean age 53 years 40% female 76% white; 21% black; 3% other; 6% Hispanic; 93% non-Hispanic; 1% NR Fibrosis stage F0 to F2: 59%; F3: 13%; 28% NR Genotype 1a: 80%; 1b: 20%; unconfirmed subtype: 0.5% Treatment-naive: 100% <u>12-week intervention group (n=216)</u> Mean age 53 years 41% female 77% white; 19% black; 3% other; 6% Hispanic; 94% non-Hispanic Fibrosis stage F0 to F2: 59%; F3: 13%; 28% NR Genotype 1a: 80%; 1b: 20% Treatment-naive: 100%	2% (8/431)	HCV RNA <25 IU/mL

	Eligibility					
	Age					
Author year	Fibrosis stage	Study				
Country	Genotype(s)	Recruitment			Loss to	
Quality	HBV status	Dates	Sample Size	Baseline Characteristics	Followup	Definition of SVR
Kowdley 2014b ¹⁹¹	Age 18 to 70 years	October 2011 to	158	A vs. B	2.5% (4/158)	HCV RNA <25 IU/mL
AVIATOR	FibroTest ≤0.72 and APRI	April 2012	A=79	Mean age 48 vs. 50 years		24 weeks after the
Australia, Canada, France,	≤2 at screening; or		B=79	43% vs. 44% female		end of treatment
Germany, New Zealand,	FibroScan <9.6 kPa, or			17% vs. 16% black; other races NR; 9%		Primary efficacy
Puerto Rico, Spain, U.K.,	the absence of cirrhosis			vs. 8% Hispanic		endpoint; 12-week
U.S.	based on a liver biopsy			Fibrosis score F2 or F3: 25% vs. 32%		post-treatment results
Good	within 36 months			Genotype 1a: 67% vs. 69%		reported in online
	Genotype 1			Treatment-naïve: 100% vs. 100%		supplement
	Patients with HBV					
	infection excluded					
Kumada 2017 (Part 2	Age 20 to 80 years	August 2014 to	Part 2 only	Mean age 61 years	NR	HCV RNA
only) ¹⁵²	Fibrosis stage NR;	October 2015	227	62% female		undetectable
Japan	patients with cirrhosis			100% Asian (Japanese)		
Good	were eligible for study			Fibrosis stage/METAVIR score NR		
	inclusion			Genotype 1a: 2%; 1b: 98%		
	Genotype 1			Treatment-naïve: 66%		
	Patients with HBV			Treatment-experienced: 34% (IFN-		
	infection excluded			containing regimen)		
Kumada 2015 ¹⁵¹	Age 18 to 75 years	December 2013	321	A vs. B	0% (0/321)	HCV RNA <lloq 12<="" td=""></lloq>
GIFI-1 (Substudy 1)	Liver biopsy within 24	through 2014	A=215	Mean age 61 vs. 62 years		weeks post-treatment
Japan	months of study with		B=106	63% vs. 56% female		
Fair	METAVIR or New					
	Inuyama Score ≤3; or if no			Fibrosis stage: F0 and F1: 60% vs. 74%;		
	biopsy Fibro lest score of			F2: 21% vs. 3%; F3: 20% vs. 23%; NR:		
	≤ 0.72 and APRI ≤ 2 ,			62% VS. 71%		
	screening transient			Genotype 1b: 100%		
	elastography (e.g.,			Treatment-haive: 65% vs. 64%		
	FIDroScan) <12.5 kPa; or			l reatment-experienced: 35% VS. 36%		
	Screening			(IFIN-containing regimen)		
	Discriminant Score <0					
	Intection excluded				1	

Appendix B Table 10. Key Questions 6-8: Direct Acting Antiviral Therapy in Adults – Study Characteristics

	Eligibility					
	Age					
Author year	Fibrosis stage	Study				
Country	Genotype(s)	Recruitment			Loss to	
Quality	HBV status	Dates	Sample Size	Baseline Characteristics	Followup	Definition of SVR
Kwo 2016 ¹⁵³	Age 18 to 70 years	April 2014 to	155	Mean age 56 years	0% (0/310)	HCV RNA <25 IU/mL
OPTIMIST-1	FibroScan ≤12.5 kPa	January 2015		47% female		or undetectable
Canada, U.S.	within 6 months of			78% white; 20% black; 1% Asian; <1%		
Fair	screening or between			other		
	screening and day 1; or,			METAVIR Score F0 to F2: 43%; F3: 10%;		
	FibroTest ≤0.48 +			NR: 47%		
	AST:platelet ratio index ≤1			Genotype 1a: 75%; 1b: 25%		
	at screening; or, liver			Treatment-naive: 74%		
	biopsy within 2 years of			Treatment-experienced: 26% (IFN-		
	screening or between			containing regimen)		
	screening and day 1			5 5 ,		
	Genotype 1					
	Patients with HBV					
	infection excluded					
Lalezari 2015 ¹⁹²	Age 18 to 70 years	April 2013 to	38	Mean age 48 years	0% (0/38)	HCV RNA <15 IU/mL
U.S.	Fibrosis stage NR: no	December 2013		34% female	· · · ·	
Fair	cirrhosis (undefined)			95% white: 3% Hispanic/Latino		
	Genotype 1			Fibrosis stage F0-F1: 79%: F2: 16%: F3:		
	Patients with HBV			5%		
	infection excluded			Genotype 1a: 84%: other subgenotypes		
	Stable opioid replacement			NR		
	therapy with either			Opioid replacement therapy, methadone:		
	methadone or			50%: buprenorphine: 50%		
	buprenorphine			Treatment-naïve: 95%		
				Treatment-experienced: 5% (pegylated		
				IFN + ribavirin)		
Lawitz 2014a ¹⁵⁴	Age ≥18 years	November 2011	41	A vs. B	0% (0/41)	HCV RNA <25 IU/mL
COSMOS	METAVIR F0-F2; previous	to January 2014	A=14	Median age 56 vs. 55 years	~ /	
U.S.	nonresponders to	,	B=27	42% vs. 26% female		
Fair	peginterferon and ribavirin			79% vs. 70% white; 21% vs. 30%		
	Genotype 1			black/African American; 14% vs. 15%		
	Patients with HBV			Hispanic/Latino		
	infection excluded			Fibrosis stage F0 and F1: 57% vs. 41%;		
				F2: 43% vs. 59%		
				Genotype 1a: 71% vs. 78%; 1b: 29% vs.		
				22%		
				Treatment-naive: 0% vs. 0%		
				Treatment-experienced: 100% vs. 100%		

Author year Country	Eligibility Age Fibrosis stage Genotype(s)	Study			Loss to	
Quality	HBV status	Dates	Sample Size	Baseline Characteristics	Followup	Definition of SVR
Lawitz 2014b ¹⁹³ LONESTAR (Cohort A) U.S. <i>Fair</i>	Age >18 years No cirrhosis, based on liver biopsy Genotype 1 Patients with HBV infection excluded	November 2012 to December 2012	60 A=20 B=19 C=21	A vs. C <u>8-week intervention group</u> Mean age 48 vs. 50 years 30% vs. 43% female 20% vs. 0% black; 80% vs. 100% non- black 15% vs. 57% Hispanic; 85% vs. 43% non- Hispanic Fibrosis stage NR; cirrhosis: 0% vs. 0% Genotype 1a: 85% vs. 90%; 1b: 15% vs. 10% Treatment-naive: 100% vs. 100% B <u>12-week intervention group</u> Mean age 46 years 42% female 5% black; 95% non-black 47% Hispanic; 53% non-Hispanic Fibrosis stage NR; cirrhosis: 0% Genotype 1a: 89% Treatment-naive: 100%	2% (1/60)	HCV RNA <25 IU/mL or undetectable
Lawitz 2015 ¹⁵⁵ PEARL-1 France, Hungary, Italy, Poland, Puerto Rico, Romania, Spain, Turkey, U.S. <i>Fair</i>	Age 18 to 70 years No cirrhosis, based on liver biopsy or FibroScan ≥14.6 kPa Genotype 1b Patients with HBV infection excluded	August 2012 to March 2014	82 (without cirrhosis; 42 treatment naïve, 40 prior null responder)*	Mean age 55 years 51% female 80% white; 15% black; 5% Asian; <1% American Indian/Alaska Native Fibrosis stage F0 and F1: 63%; F2: 23%; F3: 14% Genotype 1b: 100% Treatment naïve: 51%Treatment- experienced: 49% (pegylated IFN + ribavirin)	1% (1/82)	HCV RNA <25 IU/mL

Appendix B Table 10. Key Questions 6-8: Direct Acting Antiviral Therapy in Adults – Study Characteristics

Author year Country Quality Lim 2016 ¹⁵⁶ Korea Fair	Eligibility Age Fibrosis stage Genotype(s) HBV status Age ≥18 years Up to 20%of enrolled patients could have cirrhosis, based on liver biopsy Treatment-naïve arm only	Study Recruitment Dates NR	Sample Size	Baseline Characteristics Mean age 54 years 61% female 100% Asian Fibrosis stage NR; 9% cirrhosis Genotype 1a: 4%; 1b: 96% Treatment-naïve: 100%	Loss to Followup 0% (0/46)	Definition of SVR HCV RNA <25 IU/mL
Nelson 2015 ¹⁵⁷ ALLY-3 U.S. <i>Fair</i>	Patients with HBV infection excluded Age ≥18 years Fibrosis stage NR; patients with compensated cirrhosis were eligible for inclusion Genotype 3 Patients with HBV infection excluded	NR	101 (treatment-naïve population only)	Mean age 53 years 43% female 91% white; 4% black; 5% Asian FibroTest F0 to F3: 76%; F4: 22% Genotype 3: 100% Cirrhosis: 19% Treatment-naïve: 100%	0% (0/101)	HCV RNA <25 IU/mL
Pianko 2015 ¹⁵⁸ Australia, New Zealand, U.S. <i>Fair</i>	Age ≥18 years No cirrhosis, based on liver biopsy, FibroTest >0.75 and APRI >2.0, or FibroScan >12.5 kPa Genotype 3 Treatment experienced (IFN + ribavirin) Patients with HBV infection excluded	June 2013 to August 2014	53 A=27 B=26	A vs. B Mean age 55 vs. 56 33% vs. 35% female 93% vs. 92% white; 0% vs. 4% black Fibrosis stage NR; 0% vs. 0% cirrhosis Genotype 3: 100% vs. 100% Treatment naïve: 0% vs. 0% Treatment-experienced: 100% vs. 100%	0% (0/53)	HCV RNA <lloq< td=""></lloq<>
Poordad 2017 ¹⁹⁴ MAGELLAN-1 U.S. <i>Fair</i>	Age 18 to 70 years Liver biopsy with 24 months, FibroScan <12.5 kPa, or FibroTest ≤0.48 and APRI <1 Genotype 1 Prior DAA treatment failure Patients with HBV infection excluded	NR	50 A=6 B=22 C=22	A vs. B vs. C Mean age 59 vs. 59 vs. 56 years 50% vs. 18% vs. 9% female 33% vs. 45% vs. black; other race/ethnicity NR Fibrosis stage F0-F1: 67% vs. 50% vs. 77%; F2: 17% vs. 27% vs. 0%; F3: 17% vs. 23% vs. 23% Genotype 1a: 67% vs. 82% vs. 91%; 1b: 33% vs. 18% vs. 9% Treatment-experienced: 100% vs. 100% vs. 100%	0% (0/50)	HCV RNA <15 IU/mL

	Eligibility					
Author year Country <i>Quality</i>	Age Fibrosis stage Genotype(s) HBV status	Study Recruitment Dates	Sample Size	Baseline Characteristics	Loss to Followup	Definition of SVR
Pott-Junior 2019 (Group A daclatasvir/ sofosbuvir arm) ¹⁵⁹ Brazil <i>Good</i>	Age ≥18 years Fibrosis stage 3 based on liver biopsy or FibroScan ≥9.6 but <12.5; no cirrhosis Genotype 1 Treatment-naïve or experienced Patients with HBV excluded	NR	65	Mean age 56 years 53% female Race/ethnicity NR Mean FibroScan 9.9 kPa Genotype 1: 100% Treatment-naïve: 60% Treatment-experienced (pegylated IFN): 40%	0% (0/65)	HCV RNA <lloq< td=""></lloq<>
Pott-Junior 2019 (Group B · simeprevir/ sofosbuvir arm) ¹⁵⁹ Brazil <i>Good</i>	See Pott-Junior 2019 Group A	See Pott-Junior 2019 Group A	60	Mean age 53 years 48% female Race/ethnicity NR Mean FibroScan 10.2 kPa Genotype 1: 100% Treatment-naïve: 60% Treatment-experienced (pegylated IFN): 40%	0% (0/60)	See Pott-Junior 2019 Group A
Sperl 2016 ¹⁹⁸ and Ng 2018 ¹³⁸ C-EDGE Head-2-Head (elbasvir/grazoprevir arm only) Multinational (Europe, Turkey) <i>Fair</i>	Age NR Cirrhosis allowed; criteria NR Genotype 1, 4 or 6 Treatment naïve or experienced Patients with HBV excluded	NR	129	Mean age 48 years 57% female 99% white; other races NR Fibrosis stage NR; 17% cirrhosis Genotype 1a: 14%; 1b: 81%; 4: 5% Treatment-naïve: 78% Treatment-experienced (peg IFN + ribavirin): 22%	0.8% (1/129)	HCV RNA <15 IU/mL
Sulkowski 2014 ¹⁶¹ A1444040 Study U.S. <i>Fair</i>	Age 18 to 70 years No cirrhosis based on liver biopsy within 24 months or FibroTest ≤0.72 and APRI ≤2 Genotype 1, 2 or 3 Patients with HBV infection excluded	June 2011 to November 2012	82 A=41 B=41	A vs. B Median age 55 vs. 54 years 51% vs. 49% female 80% vs. 80% white; 12% vs. 17% black; 7% vs. 2% other Fibrosis stage F0 and F1: 37% vs. 32%; F2 and F3:46% vs. 54%; F4: 15% vs. 12% Genotype 1a: 83% vs. 80%; 1b: 17% vs. 20% Treatment-naïve; 100% vs. 100%	0% (0/82)	HCV RNA <25 IU/mL

	Eligibility					
	Age					
Author year	Fibrosis stage	Study				
Country	Genotype(s)	Recruitment			Loss to	
Quality	HBV status	Dates	Sample Size	Baseline Characteristics	Followup	Definition of SVR
Sulkowski 2015 ¹⁶⁰	Age ≥18 years	February 2013 to	129	A vs. B	0% (0/129)	HCV RNA <25 IU/mL
C-WORTHY	Fibrosis stage NR;	July 2014	A=44	Mean age 52 vs. 51 years		
Australia, Canada,	patients with HCC or		B=85	48% vs. 53% female		
Denmark France, Hungary,	decompensated liver			82% vs. 95% white; 18% vs. 5% non-		
Israel, New Zealand, Puerte	disease excluded			white; 11% vs. 9% Hispanic		
Rico, Spain, Sweden,	Genotype 1			Fibrosis stage F0 to F2: 89% vs. 95%; F3:		
Turkey, U.S.	Patients with HBV			11% vs. 5%		
Fair	infection excluded			Genotype 1a: 68% vs. 61%; 1b: 32% vs.		
				37%		
				Treatment-naïve: 100% vs. 100%		
Toyoda 2018 ¹⁹⁹	Age ≥18 years	February 2016 to	90 (Arm A only)	Mean age 57 years	1% (1/90)	HCV RNA <15 IU/mL
CERTAIN-2 (Arm A only)	No cirrhosis based on	July 2016		53% female		
Japan	liver biopsy, FibroScan			Race/ethnicity NR		
Fair	<12.5 kPa or FibroTest			Median fibrosis stage 1.6		
	≤0.72			Genotype 2a: 72%; 2b: 28%		
	Genotype 2			Treatment-naive: 83%		
	Patients with HBV			I reatment-experienced (IFN): 17%		
400	excluded					
Waked 2016 ¹⁶²	Age ≥18 years	November 2014	100 (treatment-naïve	Mean age 49 years	0% (0/100)	HCV RNA <lloq< td=""></lloq<>
AGATE-II	No cirrhosis based on	to March 2015	population only)	30% female		
Egypt	liver biopsy in the past 24			98% white; 2% black		
Good	months or Fibro lest ≤0.72			Fibrosis F0 and F1: 68%; F2: 11%; F3:		
	or APRI ≤2 or FibroScan			19%; F4: 2%		
	>12.5 kPa			Genotype 4: 100%		
	Genotype 4			Treatment-naive: 100%		
	Patients with HBV					
M/-: 0040163		Marco 004.0 ta luke	000		00((0/000)	
	Age ≥20 years	May 2016 to July	206	Mean age 47 years	0% (0/206)	HCV RNA <lloq< td=""></lloq<>
China	Cirrnosis allowed, based	2017		50% female		
Fair	Children Soon + 12 5 kDo			Race/ethnicity NR		
	FIDIOSCAII > 12.5 KPa			The stage NR, 16% cithosis		
	Treatment noïve or			Treatment experienced: 49%		
	avparianced			rreatment-experienced. 40%		
	Pationte with URV					
	excluded					

	Eligibility					
	Age					
Author year	Fibrosis stage	Study				
Country	Genotype(s)	Recruitment			Loss to	
Quality	HBV status	Dates	Sample Size	Baseline Characteristics	Followup	Definition of SVR
Wei 2019a ¹⁶⁴	Age >18 years	March 2015 to	486 (efficacy; 435	Mean age 48 years	0.2% (1/486)	HCV RNA <lloq< td=""></lloq<>
C-CORAL (Genotype 1 and	Cirrhosis allowed, based	September 2016	excluding Genotype 6);	56% female		
4 only)	on liver biopsy or		609 (harms)	72% Asian, 28% white, 0.2% other		
Multinational (Australia,	FibroScan >12.5 kPa			Fibrosis stage F0 to F2: 70%; F3: 11%;		
China, Korea, Russia,	Genotype 1 or 4			F4: 19%		
Taiwan, Thailand, Vietnam)	Treatment naïve			Genotype 1a: 8%; 1b: 80%; other type 1:		
Good	Patients with HBV			1%; 4: 0.6%		
	excluded			Treatment-naïve: 100%		
Wei 2019b ¹⁶⁵	Age ≥18 years	April 2016 to	375	Median age 45 years	0.3% (1/375)	HCV RNA <15 IU/mL
Multinational (China,	Cirrhosis allowed, based	June 2017		47% female		
Malaysia, Singapore,	on liver biopsy or			Race/ethnicity NR		
Thailand, Vietnam)	FibroScan or FibroTest			Fibrosis stage NR; 18% cirrhosis		
Fair	and APRI			Genotype 1: 34%; 2: 17%; 3: 22%; 6: 26%		
	Genotype 1-6			Treatment-naïve: 82%		
	Treatment naïve or			Treatment-experienced (primarily IFN or		
	experienced			peg IFN + ribavirin): 18%		
	Patients with HBV					
	excluded					
Zeuzem 2015 ¹⁶⁶	Age >18 years	June 2014 to	246 (immediate	Total population (n=316; 22% cirrhosis)	Total	HCV RNA
C-EDGE	Fibrosis stage NR; 20%	March 2015	treatment group only,	Mean age 52 years	population	unquantifiable
Multinational (Australia,	cirrhosis planned		without cirrhosis)	46% female	0.6% (2/316)	
Czech Republic, France,	enrollment			17% Asian; 19% black; 60% white; 4%		
Germany, Israel, Puerto	Genotype 1, 4 or 6; 15%			other		
Rico, South Korea, Taiwan,	genotype 4 or 6 planned			Fibrosis F0 to F2: 67%; F3: 11%; F4: 22%		
U.S.)	enrollment			Genotype 1a: 50%; 1b: 42%; 4: 6%; 6: 3%		
Good	Patients with HBV			Treatment-naïve: 100%		
	infection excluded					
Appendix B Table 10. Key Questions 6-8: Direct Acting Antiviral Therapy in Adults – Study Characteristics

	Eligibility					
Authorizon	Age	Study				
Author year		Bocruitmont			Loss to	
Quality	HBV status	Dates	Sample Size	Baseline Characteristics	Followup	Definition of SVR
Zeuzem 2018 ¹⁶⁷		October 2015 to	667	8-week intervention group (n=351)	0.3%(1/351)	HCV RNA <15 IU/ml
ENDURANCE-1	No cirrhosis based on	May 2016	001	Median age 53 years	0.070 (17001)	
Multinational (Australia.	liver biopsy serum	11149 2010		52% female		
Austria, Belgium, Canada,	markers or transient			4% black: 82% white: other race/ethnicity		
Chile, France, Germany,	elastography			NR		
Hungary, Israel, Italy,	Genotype 1			Fibrosis stage F0 or F1: 85%; F2: 6%; F3:		
Lithuania, Mexico, New	Treatment naïve or			9%		
Zealand, Poland, Portugal,	experienced (IFN or			Genotype 1a: 43%		
Puerto Rico, Romania,	sofosbuvir)			Treatment-naïve: 62%		
Russian Federation, Spain,	Patients with HBV			Treatment- experienced: 38% (99% IFN;		
South Korea, Sweden,	infection excluded			1% sofosbuvir)		
Switzerland, Taiwan, U.K.,				People who inject drugs: 28%		
U.S.)				Opioid substitution therapy: 3%		
Fair				HIV coinfection: 4%		
				12-week intervention group (n=352)		
				Median age 52 years		
				50% female		
				4% black; 86% white; other race/ethnicity		
				NR		
				Fibrosis stage F0 or F1: 85%; F2: 7%; F3:		
				17%		
				Genotype 1a: 41%		
				I reatment-naive: 62% I reatment-		
				experienced: 38% (99% IFN; 1%		
				SOTOSDUVIĽ)		
				People who inject drugs: 28%		
				Upiola substitution therapy. 5%		

Appendix B Table 10. Key Questions 6-8: Direct Acting Antiviral Therapy in Adults – Study Characteristics

	Eligibility					
Author year	Fibrosis stage	Study				
Country	Genotype(s)	Recruitment			Loss to	
Quality	HBV status	Dates	Sample Size	Baseline Characteristics	Followup	Definition of SVR
Zeuzem 2018 ¹⁶⁷	Age ≥18 years	Same as Zeuzem	505	A vs. B vs. C	0.6% (3/505)	Same as Zeuzem
ENDURANCE-3 (same	No cirrhosis based on	2018	A=157	Median age 47 vs. 48 vs. 49 years		2018
publication as	liver biopsy, serum		B=233	41% vs. 48% vs. 55% female		
ENDURANCE-1)	markers or transient		C=115	2% vs. 2% vs. 3% black; 85% vs. 8*% vs.		
Fair	elastography			90% white; other race/ethnicity NR		
	Genotype 3			Fibrosis stage F0 or F1: 78% vs. 86% vs.		
	Treatment naïve or			84%; F2: 5% vs. 5% vs. 7%; F3: 17% vs.		
	experienced (IFN or			9% vs. 9%		
	sofosbuvir)			Genotype 3: 100% vs. 100% vs. 100%		
	Patients with HBV			Treatment-naïve: 100% vs. 100% vs.		
	infection excluded			100%		
				People who inject drugs: 66% vs. 64% vs.		
				63%		
				Opioid substitution therapy: 20% vs. 16%		
				vs. 15%		

Note: *Excluding patients who withdrew or were lost to follow up.

Abbreviations: APRI = aspartate amino transferase to platelet ratio index; AST = aspartate amino transferase; DAA = direct acting antiviral; HBV = hepatitis B virus; HCV = hepatitis C virus; HCC = hepatocellular carcinoma; IFN = interferon; IVDU = injection drug use; LLOQ = lower limit of quantification; NR = not reported; RNA = ribonucleic acid; SVR = sustained virologic response; U.K. = United Kingdom; U.S. = United States. Study names are not acronyms.

Author year	Treatment Regimen	Treatment			
Country	(1x/day unless	Duration and	Overall SVR	Genotype SVR	
Quality	otherwise noted)	Assessments	Results	Results	Other Subgroup SVR Results
Abergel 2016a ¹⁴² France <i>Fair</i>	Ledipasvir 90 mg + sofosbuvir 400 mg	Treatment duration: 12 weeks Timing of assessment: 12 weeks post- treatment	SVR: 96% (21/22)	Genotype 4: 96% (21/22)	NR
Abergel 2016b ¹⁴¹ France <i>Good</i>	Ledipasvir 90 mg + sofosbuvir 400 mg	Treatment duration: 12 weeks Timing of assessment: 12 weeks post- treatment	SVR: 95% (20/21)	Genotype 5: 95% (20/21)	NR
Afdahl 2014 ¹⁸⁵ ION-1 U.S. and Europe <i>Fair</i>	A. Ledipasvir 90 mg + sofosbuvir 400 mg B. Ledipasvir 90 mg + sofosbuvir 400 mg + ribavirin	Treatment duration: 12 to 24 weeks Timing of assessment: 12 weeks post- treatment	A vs. B <u>12-week intervention</u> group SVR: 99% (211/214) vs. 97% (211/217) <u>24-week intervention</u> group SVR: 98% (212/217) vs. 99% (215/217)	A vs. B <u>SVR, 12-week</u> <u>intervention group*</u> Genotype 1a: 99% (141/142) vs. 100% (143/143) Genotype 1b: 100% (66/66) vs. 100% (67/67) Other: 100% (4/4) vs. 100% (1/1) <u>SVR, 24-week</u> <u>intervention group*</u> Genotype 1a: 100% (143/143) vs. 100% (143/143) vs. 100% (141/141) Genotype 1b: 97% (66/68) vs. 100% (71/71) Other: 100% (3/3) vs. 100% (3/3)	A vs. B <u>SVR, 12-week intervention group</u> * <65 years: 99% (196/197) vs. 100% (189/189) ≥65 years: 100% (15/15) vs. 100% (22/22) Male: 99% (125/126) vs. 100% (124/124) Female: 100% (86/86) vs. 100% (87/87) Black: 100% (24/24) vs. 100% (26/26) Non-Black: 99.5% (187.188) vs. 100% (184/184) Hispanic: 100% (26/26) vs. 100% (19/19) Non-Hispanic: 99.5% (184/185) vs. 100% (192/192) No cirrhosis: 100% (179/179) vs. 100% (178/178) Cirrhosis: 97% (32/33) vs. 100% (33/33) <u>SVR, 24-week intervention group</u> * <65 years: 99.5% (191/192) vs. 100% (202/202) ≥65 years: 96% (21/22) vs. 100% (13/13) Male: 99% (136/138) vs. 100% (118/118) Female: 100% (76/76) vs. 100% (97/97) Black: 94% (29/31) vs. 100% (26/26) Non-Black: 100% (183/183) vs. 100% (188/188) Hispanic: 100% (29/29) vs. 100% (26/26) Non-Hispanic: 100% (183/183) vs. 100% (179/179) Cirrhosis: 97% (31/32) vs. 100% (36/36)

Author year	Treatment Regimen	Treatment			
Country	(1x/day unless	Duration and	Overall SVR	Genotype SVR	
Quality	otherwise noted)	Assessments	Results	Results	Other Subgroup SVR Results
Ahmed 2018 ¹⁹⁵ Egypt <i>Fair</i>	Ledipasvir 90 mg + sofosbuvir 400 mg	Treatment duration: 12 weeks Timing of	SVR: 99% (99/100)	Genotype 4: 99% (99/100)	NR
		assessments: 12 weeks post- treatment			
Andreone 2014 ¹⁸⁶ PEARL-II Austria, Belgium, Italy, The Netherlands, Portugal, Puerto Rico, Sweden, Switzerland, U.S. <i>Fair</i>	A. Ombitasvir 25 mg + paritaprevir 150 mg + ritonavir 100 mg + dasabuvir 250 mg 2x/day B. Ombitasvir 25 mg + paritaprevir 150 mg + ritonavir 100 mg + dasabuvir 250 mg 2x/day + ribavirin	Treatment duration: 12 weeks Timing of assessment: 12 weeks post- treatment	A vs. B SVR: 100% (91/91) vs. 97% (85/88)	A vs. B Genotype 1b: 100% (91/91) vs. 97% (85/88)	A vs. B Male: 100% (54/54) vs. 95% (41/43) Female: 100% (37/37) vs. 98% (44/45) Black: 100% (5/5) vs. 100% (3/3) Other: 100% (86/86) vs. 97% (82/85)
Asselah 2018 ¹⁹⁶ SURVEYOR II Part 4, Multinational (Asia, Europe, U.S. [specific countries NR]) <i>Fair</i>	Glecaprevir 300 mg + pibrentasvir 120 mg	Treatment duration: 8 weeks Timing of assessments: 12 weeks post- treatment	SVR: 97% (196/203)	Genotype 2: 98% (142/145) Genotype 4: 93% (43/46) Genotype 5: 100% (2/2) Genotype 6: 90% (9/10)	NR
Asselah 2019 ¹⁴³ ENDURANCE-5 Multinational (Australia, Belgium, Canada, France, New Zealand, Singapore, South Africa, Vietnam, U.S.) <i>Fair</i>	Glecaprevir 300 mg + pibrentasvir 120 mg	Treatment duration: 8 weeks Timing of assessments: 12 weeks post- treatment	SVR: 96% (22/23)	Genotype 5: 96% (22/23)	NR (reported for combined genotypes only)
Asselah 2019 ¹⁴³ ENDURANCE-6 (same publication as ENDURANCE-5) <i>Fair</i>	See Asselah 2019 ENDURANCE-5	See Asselah 2019 ENDURANCE- 5	SVR: 98% (60/61)	Genotype 6: 98% (60/61)	See Asselah 2019 ENDURANCE-5

Author year	Treatment Regimen	Treatment			
Country	(1x/day unless	Duration and	Overall SVR	Genotype SVR	
Quality	otherwise noted)	Assessments	Results	Results	Other Subgroup SVR Results
Brown 2018 ¹⁴⁴	A. Elbasvir 50 mg +	Treatment	A vs. B	NR	NR
C-SCAPE (Genotype 4	grazoprevir 100 mg	duration: 12	SVR: 90% (9/10) vs.		
only)	(n=10)	weeks	100% (10/10)		
Multinational (Australia,	B. Elbasvir 50 mg +	Timing of			
Belgium, France, Israel,	grazoprevir 100 mg +	assessments:			
Spain, U.K., U.S.)	ribavirin (n=10)	12 weeks post-			
Fair		treatment			
Chayama 2018 ¹⁹⁷	Glecaprevir 300 mg +	Treatment	SVR: 99% (128/129)	Genotype 1: 99%	NR
CERTAIN-1 (Arm A only)	pibrentasvir 120 mg	duration: 8		(128/129)	
Japan <i>Fair</i>		weeks			
		Timing of			
		assessments:			
		12 weeks post-			
		treatment			
Chuang 2016 ¹⁴⁵	Ledipasvir 90 mg +	Treatment	SVR: 98% (83/85)	Genotype 1: 98%	Treatment-naïve: 100% (42/42)
Taiwan	sofosbuvir 400 mg	duration: 12		(83/85)	Treatment experienced: 95% (41/43)
Fair		weeks			
		Timing of			
		assessment:			
		12 weeks post-			
		treatment			

Author year	Treatment Regimen	Treatment			
		Duration and	Overall SV/P	Conotype SVP	
Country	(TX/day unless	Duration and	Dverall SVR	Beaulta	Other Subgroup SVD Booulto
Quality	otherwise noted)	Assessments	Results	Results	Other Subgroup SVR Results
Dore 2016 ¹³⁷	Genotype 1a	Ireatment	Genotype 1a	<u>Genotype 1a</u>	NR
MALACHITE-1	A. Ombitasvir 25 mg +	duration: 12	A vs. B	A vs. B	
Australia, Canada, Europe,	paritaprevir 150 mg +	weeks; some	SVR: 97% (67/69)	SVR: 97% (67/69) vs.	
South America	ritonavir 100 mg +	patients in	vs. 82% (28/34)	82% (28/34)	
Good	dasabuvir 250 mg	groups B and	Genotype 1b	Genotype 1b	
	2x/day + weight-based	D received up	C vs. D vs. E	C vs. D vs. E	
	ribavirin	to 48 weeks of	SVR: 99% (83/84)	SVR: 99% (83/84) vs.	
	B. Telaprevir 750 mg	pegylated IFN /	vs. 98% (81/83) vs.	98% (81/83) vs. 78%	
	3x/day + subcutaneous	ribavirin	78% (32/41)	(32/41)	
	pegylated IEN 180 ug	Timing of	1070 (02/41)	(02/41)	
	1/wook i woight based	accoccmont:			
	ribovirin	12 weeks post			
	nbavinn Caratura 1h	12 weeks post-			
	Genotype Tb	treatment			
	C. Ombitasvir 25 mg +				
	paritaprevir 150 mg +				
	ritonavir 100 mg +				
	dasabuvir 250 mg				
	2x/day + weight-based				
	ribavirin				
	D. Ombitasvir 25 mg +				
	paritaprevir 150 mg +				
	ritonavir 100 mg +				
	dasabuvir 250 mg				
	2v/day				
	E Telaprevir 750 mg				
	SX/Uay + Subcularieous				
	1/week + weight-based				
	ribavirin				
Dore 2016 ¹³⁷	A. Ombitasvir 25 mg +	Treatment	A vs. B	A vs. B	NR
MALACHITE-2	paritaprevir 150 mg +	duration: 12	SVR: 99% (100/101)	Genotype 1a: 100%	
Australia, Canada, Europe,	ritonavir 100 mg +	weeks; some	vs. 66% (31/47)	(19/19) vs. 57% (4/7)	
South America	dasabuvir 250 mg	patients in		Genotype 1b: 99%	
Good	2x/day + weight-based	group B and D		(81/82) vs. 68%	
	ribavirin	received up to		(27/40)	
	B. Telaprevir 750 mg	48 weeks of		· · · · ·	
	3x/dav + subcutaneous	pegylated IFN /			
	pegylated IEN 180 ug	ribavirin			
	1/week + weight-based	Timing of			
	ribovirin	accoccmont:			
		12 wooko poot			
		1∠ weeks post-			
	1	treatment			

Author year	Treatment Regimen	Treatment	Overall SVP	Genotype SVP	
Quality	otherwise noted)		Results	Results	Other Subgroup SVR Results
Everson 2015 (Part A) ¹⁴⁶ U.S. <i>Good</i>	Part A (trial phase) A. Sofosbuvir 400 mg + velpatasvir 25 mg (Genotype 1) B. Sofosbuvir 400 mg + velpatasvir 100 mg (Genotype 1) C. Sofosbuvir 400 mg + velpatasvir 25 mg (Genotype 3) D. Sofosbuvir 400 mg + velpatasvir 100 mg (Genotype 3) E. Sofosbuvir 400 mg + velpatasvir 25 mg (Genotype 2; 4-6) F. Sofosbuvir 400 mg + velpatasvir 100 mg (Genotype 2; 4-6)	Treatment duration: 12 weeks Timing of assessment: 12 weeks post- treatment	A vs. B vs. C vs. D vs. E vs. F SVR: 96% (26/27) vs. 100% (28/28) vs. 93% (25/27) vs. 93% (25/27) vs. 96% (22/23) vs. 95% (21/22)	A vs. B vs. C vs. D vs. E vs. F Genotype 1, Group A: 96% (26/27) Genotype 1, Group B: 100% (28/28) Genotype 3, Group C: 93% (25/27) Genotype 3, Group D: 93% (25/27) Genotype 2 or 4-6, Group E: 96% (22/23) Genotype 2 or 4-5, Group F: 95% (21/22)	NR
Feld 2014 ¹⁸⁷ SAPPHIRE-1 Australia, New Zealand; Austria, France, Germany, Hungary, Great Britain, Italy, Spain, Sweden, Switzerland; Canada, U.S. <i>Good</i>	A. Ombitasvir 25 mg + paritaprevir 150 mg + ritonavir 100 mg + dasabuvir 250 mg 2x day + weight-based ribavirin B. Placebo for 12 weeks followed by open-label ombitasvir 25 mg + paritaprevir 150 mg + ritonavir 100 mg + dasabuvir 250 mg 2x day + weight-based ribavirin	Treatment duration: 12 weeks Timing of assessment: 12 weeks post- treatment	SVR: 96% (455/473)	Genotype 1a: 95% (307/322) 1b: 98% (148/151)	Age <55 years: 97% (95% CI, 94.5 to 98.7); (280/290) Age ≥55 years: 96% (95% CI, 92.7 to 98.6); (175/183) Male: 95% (95% CI, 92.7 to 97.8); (258/271) Female: 98% (95% CI, 95.4 to 99.7); (197/202) Black: 96% (95% CI, 89.6 to 100.0); (27/28) Non-Black: 96% (95% CI, 94.4 to 98.0); (428/445) F0 or F1: 97% (95% CI, 95.2 to 98.7); (352/363) F2: 94% (95% CI, 88.9 to 99.7); (66/70) F3: 93% (95% CI, 84.3 to 100.0); (37/40) History of diabetes: 100% (95% CI, 100.0-100.0); (19/19) No history of diabetes: 96% (95% CI, 94.2 to 97.8); (436/454)
Feld 2015 ¹³⁹ ASTRAL-1 U.S., Canada, Europe, Hong Kong <i>Good</i>	A. Sofosbuvir 400 mg + velpatasvir 100 mg B. Placebo	Treatment duration: 12 weeks Timing of assessment: 12 weeks post- treatment	A vs. B SVR: 99% (618/624) vs. 0% (0/116)	Group A only Genotype 1: 99% (323/328) 1a: 98% (206/210) 1b: 99% (117/118) 2: 100% (104/104) 4: 100% (116/116)	Group A only Age <65 years: 99% (530/536) -Genotype 1: 98% (287/292); Genotype 2: 100% (79/79); Genotype 4: 100% (116/116); Genotype 5: 95% (18/19); Genotype 6: 100% (41/41) Age ≥65 years: 100% (88/88)

Author year	Treatment Regimen	Treatment			
Country	(1x/day unless	Duration and	Overall SVR	Genotype SVR	
Quality	otherwise noted)	Assessments	Results	Results	Other Subgroup SVR Results
Feld 2015 ¹³⁹					-Genotype 1: 100% (36/36); Genotype 2: 100% (25/25);
ASTRAL-1					Genotype 4: 100% (11/11); Genotype 5: 100% (16/16);
U.S., Canada, Europe,					Genotype 6: 0/0
Hong Kong					Male: 99% (369/374)
Good					-Genotype 1: 98% (193/197); Genotype 2: 100% (57/57);
(cont'd)					Genotype 4: 100% (86/86); Genotype 5: 93% (13/14); Genotype
					6: 100% (21/21)
					Female: 99.6% (249/250)
					-Genotype 1: 99% (130/131); Genotype 2: 100% (47/47);
					Genotype 4: 100% (30/30); Genotype 5: 100% (21/21);
					Genotype 6: 100% (21/21)
					White: 99% (488/493)
					-Genotype 1: 99% (275/279); Genotype 2: 100% (82/82);
					Genotype 4: 100% (96/96); Genotype 5: 97% (34/35); Genotype
					6: 100% (1/1)
					Black: 98% (51/52)
					-Genotype 1: 96% (24/25); Genotype 2: 100% (13/13); Genotype
					4: 100% (14/14); Genotype 5 & 6: 0/0
					Other: 100% (76/76)
					-Genotype 1: 100% (22/22); Genotype 2: 100% (8/8); Genotype
					4: 100% (6/6); Genotype 5 & 6: 0/0
					NO CITTIOSIS. 99% (490/301) Capatype 1: 08% (251/255): Capatype 2: 100% (02/02):
					C_{0}
					6: 100% (25/25)
					$\begin{array}{c} 0.100\% (33/33) \\ \text{Cirrbosic: } 0.0\% (120/121) \\ \end{array}$
					Genotype 1: 99% (72/73): Genotype 2: 100% (10/10): Genotype
					4: 100% (27/27): Genotype 5: 100% (5/5): Genotype 6: 100%
					(6/6)
					Treatment-naïve: 99% (418/423)
					-Genotype 1: 98% (214/218: Genotype 1a: 97% [128/132]:
					Genotype 1b: 100% [86/86]): Genotype 2: 100% (79/79):
					Genotype 4: 100% (64/64): Genotype 5: 96% (23/24): Genotype
					6: 100% (38/38)
					Treatment-experienced: 99.5% (200/201)
					-Genotype 1: 99% (109/110: Genotype 1a: 100% [78/78]:
					Genotype 1b: 97% [31/32]): Genotype 2: 100% (25/25):
					Genotype 4: 100% (52/52); Genotype 5: 100% (11/11):
					Genotype 6: 100% (3/3)

Author year	Treatment Regimen	Treatment			
Country	(1x/day unless	Duration and	Overall SVR	Genotype SVR	
Quality	otherwise noted)	Assessments	Results	Results	Other Subgroup SVR Results
Ferenci 2014 ¹⁸⁸	A. Ombitasvir 25 mg +	Treatment	SVR: 99% (207/209)	Genotype 1b: 99%	NR
PEARL III	paritaprevir 150 mg +	duration: 12	vs. 99.5% (209/210)	(207/209) vs. 99.5%	
Austria, Belgium, Hungary,	ritonavir 100 mg +	weeks		(209/210)	
Israel, Italy, Poland,	dasabuvir 250 mg	Timing of			
Portugal, Romania, Russia,	2x/day	assessment:			
Spain, U.S.	B. Ombitasvir 25 mg +	12 weeks post-			
Good	paritaprevir 150 mg +	treatment			
	ritonavir 100 mg +				
Same publication as	dasabuvir 250 mg				
PEARL IV	2x/day + ribavirin				
Ferenci 2014 ¹⁸⁸	A. Ombitasvir 25 mg +	Treatment	A vs. B	Genotype 1a: 90%	NR
PEARL IV	paritaprevir 150 mg +	duration: 12	SVR: 90% (185/205)	(185/205) vs. 97%	
Canada, U.K., U.S.	ritonavir 100 mg +	weeks	vs. 97% (97/100)	(97/100)	
Good	dasabuvir 250 mg				
	2x/day	Timing of			
Same publication as	B. Ombitasvir 25 mg +	assessment:			
PEARL III	paritaprevir 150 mg +	12 weeks post-			
	ritonavir 100 mg +	treatment			
	dasabuvir 250 mg				
	2x/day + ribavirin				
Foster 2015 ¹⁴⁷	A. Sofosbuvir 400 mg +	Treatment	A vs. B	Genotype 2: SVR: 99%	NR
ASTRAL-2	velpatasvir 100 mg	duration: 12	SVR: 99% (133/134)	(133/134) vs. 94%	
U.S.	B. Sofosbuvir 400 mg +	weeks	vs. 94% (124/132)	(124/132)	
Fair	ribavirin				
		Timing of			
		assessment:			
		12 weeks post-			
		treatment			

Author year	Treatment Regimen	Treatment			
Country	(1x/day unless	Duration and	Overall SVR	Genotype SVR	
Quality	otherwise noted)	Assessments	Results	Results	Other Subgroup SVR Results
Foster 2015 ¹⁴⁷ ASTRAL-3 U.S. <i>Fair</i> <i>Same publication as</i> <i>ASTRAL-2</i>	Same as Foster 2015 ASTRAL-2	Treatment duration: 12 (group A) or 24 (group B) weeks Timing of assessment: 12 weeks post- treatment	A vs. B SVR: 95% (264/277) vs. 80% (221/275)	A vs. B Genotype 3: 95% (264/277) vs. 80% (221/275)	A vs. B Age <65 years: 95% (257/270) vs. 81% (210/261) Age ≥65 years: 100% (7/7) vs. 79% (11/14) Male: 94% (159/170) vs. 76% (132/175) Female: 98% (105/107) vs. 88% (89/101) Black: 100% (3/3) vs. 100% (1/1) White: 95% (238/250) vs. 78% (187/239) Other: 96% (23/24) vs. 94% (32/34) No cirrhosis: 97% (191/197) vs. 87% (163/187) Cirrhosis: 91% (73/80) vs. 66% (55/83) Missing data: 0% vs. 60% (3/5) Treatment-naive: 97% (200/206) vs. 86% (176/204) Treatment-experienced: 90% (64/71) vs. 63% (45/71) No cirrhosis + treatment-naive: 98% (160/163) vs. 90% (141/156) No cirrhosis + treatment-experienced: 91% (31/34) vs. 71% (22/31)
Gane 2015 ¹⁴⁸ New Zealand (Genotype 6 subset) <i>Fair</i>	Ledipasvir 90 mg + sofosbuvir 400 mg	Treatment duration: 12 weeks Timing of assessment: 12 weeks post- treatment	SVR: 96% (24/25)	Genotype 6: 96% (24/25)	NR
Grebely 2018 ¹⁵⁰ SIMPLIFY Multinational (Australia, Canada, New Zealand, Norway, Switzerland, U.K., U.S.) <i>Fair</i>	Sofosbuvir 400 mg + velpatasvir 100 mg	Treatment duration: 12 weeks Timing of assessments: 12 weeks post- treatment	SVR: 94% (97/103)	NR	Male: 92% (68/74) Female: 100% (29/29) Age ≤41 years: 93% (26/28) Age >41 years: 95% (71/75) F0 and F1: 97% (57/59) F2 and F3: 93% (25/27) Cirrhosis: 78% (7/9) Current opioid substitution therapy: 96% (43/45) No current opioid substitution therapy: 93% (54/58) Recent IVDU: 95% (72/76) No recent IVDU: 93% (25/27)

Author year	Treatment Regimen	Treatment			
Country	(1x/day unless	Duration and	Overall SVR	Genotype SVR	
Quality	otherwise noted)	Assessments	Results	Results	Other Subgroup SVR Results
Grebely 2018 ¹⁴⁹	Ombitasvir 25 mg +	Treatment	SVR: 91% (79/87)	Genotype 1: 91%	Male: 91% (61/67)
D3FEAT	paritaprevir 150 mg +	duration: 12		(79/87)	Female: 90% (18/20)
Multinational (Australia,	ritonavir 100 mg +	weeks			Age ≤54 years: 89% (59/66)
Canada, France, New	dasabuvir 250 mg +				Age >54 years: 95% (20/21)
Zealand, Norway,	1000 to 1200 mg	Timing of			F0 and F1: 90% (61/68)
Switzerland)	ribavirin	assessments:			F2 and F3: 100% (12/12)
Fair		12 weeks post-			Cirrhosis: 86% (6/7)
		treatment			Recent IVDU: 93% (39/42)
					No recent IVDU: 89% (40/45)
Hezode 2015 ¹⁸⁹	Ombitasvir 25 mg +	Treatment	SVR: 100% (42/42)	Genotype 4: 100%	NR
PEARL I (Treatment-naïve	paritaprevir 150 mg +	duration: 12		(42/42)	
population)	ritonavir 100 mg +	weeks			
France, Hungary, Italy,	ribavirin (weight-based;				
Poland, Romania, Spain,	dose NR)	Timing of			
Turkey, U.S.		assessment:			
Good		12 weeks post-			
		treatment			
See also Lawitz 2015 ¹⁵⁵					
(PEARL I - Genotype 1b)					
Hezode 2015 ¹⁸⁹	Same as Hezode 2015	Same as	SVR: 100% (49/49)	Genotype 4: 100%	NR
PEARL I (Treatment	(Treatment naïve	Hezode 2015		(49/49)	
experienced population)	population)	(Treatment			
France, Hungary, Italy,		naïve			
Poland, Romania, Spain,		population)			
Turkey, U.S.					
Good					
See also Lawitz 2015 ¹⁵⁵					
(PFARL I - Genotype 1b)					
(FEARLI - Genolype ID)					

Author year	Treatment Regimen	Treatment			
Country	(1x/day unless	Duration and	Overall SVR	Genotype SVR	
Quality	otherwise noted)	Assessments	Results	Results	Other Subgroup SVR Results
Kowdley 2014a ¹⁹⁰	Ledipasvir 90 mg +	Treatment	8-week intervention	8-week intervention	8-week intervention group
ION-3	sofosbuvir 400 mg	duration: 8 to	group	group	<65 years: 94% (185/196)
U.S.	_	12 weeks	SVR: 94% (202/215)	Genotype 1a: 93%	≥65 years: 90% (17/19)
Fair		Timing of		(159/171)	Male: 92% (119/130)
		assessment:	12-week intervention	Genotype 1b: 98%	Female: 98% (83/85)
		12 weeks post-	group	(42/43)	Black: 91% (41/45)
		treatment	SVR: 95% (206/216)	Unconfirmed subtype:	Non-black: 95% (161/170)
				100% (1/1)	Hispanic: 100% (13/13)
					Non-Hispanic: 94% (187/200)
				12-week intervention	
				group	12-week intervention group
				Genotype 1a: 95%	<65 years: 95% (189/199)
				(163/172)	≥65 years: 100% (17/17)
				Genotype 1b: 98%	Male: 95% (122/128)
				(43/44)	Female: 96% (84/85)
					Black: 95% (40/42)
					Non-black: 95% (165/173)
					Hispanic: 93% (13/14)
					Non-Hispanic: 96% (193/202)
Kowdley 2014b ¹⁹¹	A. Ombitasvir 25 mg +	Treatment	A vs. B	A vs. B	A vs. B
AVIATOR	paritaprevir 150 mg +	duration:	SVR, 12 weeks post-	Genotype 1a +	Black: 100% (13/13) vs. 100% (13/13)
Australia, Canada, France,	ritonavir 150 mg +	12 weeks	treatment: 91%	treatment naive: 83%	Non-black: 86% (57/66) vs. 96% (63/66)
Germany, New Zealand,	dasabuvir 800 mg		(72/79) vs. 99%	(43/52) vs. 94%	
Puerto Rico, Spain, U.K.,	B. Ombitasvir 25 mg +	Timing of	(78/79)	(51/54)	
U.S.	paritaprevir 150 mg +	assessment:	SVR, 24 weeks post-	Genotype 1b +	
Good	ritonavir 100-150 mg +	24 weeks post-	treatment: 89%	treatment naive: 100%	
	dasabuvir 800 mg +	treatment	(70/79) vs. 96%	(25/25) vs. 100%	
	ribavirin 1000-1200 mg		(76/79)	(25/25)	
Kumada 2017 (Part 2	Elbasvir 50 mg +	Treatment	SVR: 97% (219/227)	Genotype 1a: 100%	<65 years: 99% (122/123)
only) ¹⁵²	grazoprevir 100 mg	duration: 12		(4/4)	65-74 years: 93% (70/75)
Japan		weeks		Genotype 1b: 96%	≥75 years: 93% (27/29)
Good		Timing of		(215/223)	Male: 98% (85/87)
		assessment:			Female: 96% (134/140)
		12 weeks post-			Treatment-naïve: 97% (144/149)
		treatment			Treatment-experienced: 96% (75/78)

Author year	Treatment Regimen	Treatment			
Country	(1x/day unless	Duration and	Overall SVR	Genotype SVR	
Quality	otherwise noted)	Assessments	Results	Results	Other Subgroup SVR Results
Kumada 2015 ¹⁵¹ GIFT-1 (Substudy 1) Japan <i>Fair</i>	A. Ombitasvir 25 mg + paritaprevir 150 mg + ritonavir 100 mg (double-blind treatment) B. Placebo for 12 weeks, followed by ombitasvir 25 mg + paritaprevir 150 mg + ritonavir 100 mg (open- label treatment)	Treatment duration: 12 weeks Timing of assessment: 12 weeks post- treatment	A vs. B SVR: 95% (204/215) vs. 98% (104/106)	A vs. B Genotype 1b: 95% (204/215) vs. 98% (104/106)	A vs. B Treatment-naïve: 94.2% (131/139) vs. 98/5% (67/68) Treatment-experienced: 96.1% (73/76) vs. 97.4% (37/38)
Kwo 2016 ¹⁵³ OPTIMIST-1 Canada, U.S. <i>Fair</i>	Simeprevir 150 mg + sofosbuvir 400 mg	Treatment duration: 12 weeks Timing of assessment: 12 weeks post- treatment	SVR: 97% (150/155)	Genotype 1a: 97% (112/116) Genotype 1b: 97% (38/39)	Treatment-naïve: 97% (112/115) Treatment experienced: 95% (38/40)
Lalezari 2015 ¹⁹² U.S. <i>Fair</i>	Ombitasvir 25 mg + paritaprevir 150 mg + ritonavir 100 mg + dasabuvir 250 mg 2x/day + ribavirin 1000- 1200 mg	Treatment duration: 12 weeks Timing of assessment: 12 and 24 weeks post treatment	SVR, 12 weeks: 97.4% (37/38) SVR, 24 weeks: 97.4% (37/38)	Genotype 1, 12 weeks: 97.4% (37/38) Genotype 1, 24 weeks: 97.4% (37/38)	NR
Lawitz 2014a ¹⁵⁴ COSMOS U.S. <i>Fair</i>	A. Simeprevir 150 mg + sofosbuvir 400 mg B. Simeprevir 150 mg + sofosbuvir 400 mg + ribavirin	Treatment duration: 12 weeks Timing of assessment: 12 weeks post treatment	SVR: 92.9% (13/14) vs. 96% (26/27)	Genotype 1: 92.9% (13/14) vs. 96% (26/27)	Treatment-naïve: (4/4) vs. (5/6)

Author year	Treatment Regimen	Treatment			
Country	(1x/day unless	Duration and	Overall SVR	Genotype SVR	
Quality	otherwise noted)	Assessments	Results	Results	Other Subgroup SVR Results
Lawitz 2014b ¹⁹³ LONESTAR (Cohort A) U.S. <i>Fair</i>	A. Ledipasvir 90 mg + sofosbuvir 400 mg, 8 weeks B. Ledipasvir 90 mg + sofosbuvir 400 mg, 12 weeks C. Ledipasvir 90 mg + sofosbuvir 400 mg + ribavirin	Treatment duration: 8 and 12 weeks Timing of assessment: 12 weeks post treatment	A vs. C 8-week intervention group SVR: 95% (19/20) vs. 100% (21/21) B 12-week intervention group SVR: 95% (18/19)	A vs. C <u>8-week intervention</u> <u>group</u> Genotype 1: 95% (19/20) vs. 100% (21/21) B <u>12-week intervention</u> <u>group</u> Construe 1: 05%	NR
Lawitz 2015 ¹⁵⁵ PEARL-1 France, Hungary, Italy, Poland, Puerto Rico, Romania, Spain, Turkey, U.S. <i>Fair</i>	Ombitasvir 25 mg + paritaprevir 150 mg + ritonavir 100 mg	Treatment duration: 12 weeks Timing of assessment: 12 weeks post- treatment	SVR: 92.7% (76/82)	Genotype 1: 95% (18/19) Genotype 1b: 92.7% (76/82)	Treatment-naïve: 95.2% (40/42) Treatment-experienced: 90.0% (36/40)
Lim 2016 ¹⁵⁶ Korea <i>Fair</i>	Ledipasvir 90 mg + sofosbuvir 400 mg	Treatment duration: 12 weeks Timing of assessments: 12 weeks post treatment	SVR: 100% (46/46)	Genotype 1: 100% (46/46)	Age <65 years: 100% (33/33) Age ≥65 years: 10% (13/13)
Nelson 2015 ¹⁵⁷ ALLY-3 U.S. <i>Fair</i>	Daclatasvir 60 mg + sofosbuvir 400 mg	Treatment duration: 12 weeks Timing of assessment: 12 weeks post- treatment	SVR: 90% (91/101)	Genotype 3: 90% (91/101)	Age <65 years: 90% (128/142) [†] Age ≥65 years: 70% (7/10) [†] Male gender: 86% (77/90) [†] Female gender: 94% (58/62) [†] F0-F3: 95% (72/76) F4: 73% (16/22) Treatment-naïve: 97% (73/75) Treatment-experienced: 94% (32/34)

Author year Treatment Regimen Treatment	
Country (1x/day unless Duration and Overall SVR Genotype SVR	
Quality otherwise noted) Assessments Results Results Other Subgroup SVR Results	
Pianko 2015 ¹⁵⁸ A. Sofosbuvir 400 mg + Treatment A vs. B A vs. B NR	
Australia, New Zealand, velpatasvir 100 mg duration: 12 SVR: 100% (27/27) Genotype 3: 100%	
U.S. (Group 3) weeks vs. 100% (26/26) (27/27) vs. 100%	
Fair B. Sofosbuvir 400 mg + (26/26)	
velpatasvir 100 mg + Timing of	
ribavirin (Group 4) assessment:	
12 weeks post-	
treatment	
Poordad 2017 ¹⁹⁴ A. Glecapravir 200 mg Treatment A vs. B vs. C A vs. B vs. C NR	
MAGELLAN-1 + pibrentasvir 80 mg duration: 12 SVR: 100% (6/6) vs. Genotype 1: 100%	
U.S. B. Glecapravir 200 mg weeks 86% (19/22) vs. 95% (6/6) vs. 86% (19/22)	
Fair + pibrentasvir 120 mg Timing of (21/22) vs. 95% (21/22)	
C. Glecapravir 200 mg assessment:	
+ pibrentasvir 120 mg + 12 weeks post-	
ribavirin treatment	
Pott-Junior 2019 (Group A - Daclatasvir 60 mg + Treatment SVR: 100% (65/65) Genotype 1a: 100% Treatment-naïve: 100% (39/39)	
daclatasvir/ sofosbuvir sofosbuvir 400 mg duration: 12 (27/27) Treatment-experienced: 100% (26/26)	
arm) ¹⁵⁹ Weeks Genotype 1b: 100%	
Brazil (35/35)	
Good assessments:	
12 weeks post-	
treatment	
Pott-Junior 2019 (Group B - Simeprevir 150 mg + See Pott- SVR: 93% (56/60) Genotype 1a: 90% Treatment-naive: 97% (35/36)	
simeprevir/sorosbuvir sorosbuvir 400 mg Junior 2019 (28/31) [reatment-experienced: 88% (21/24)	
arm) ¹³⁹ Group A Genotype 1b: 96%	
6000 Enert 2016 ¹⁹⁸ and Na	
Spen 2016 ¹⁰⁰ and Ng Elbasvir 50 mg + (Treatment SVR: 99% (128/129) Genotype 1a: 100% (Male: 100% (55/55)) 2019^{138} (10/48)	
2010^{100} (16/16) [Grazoprevir 100 mg uuration. 12] (16/16) [Grazoprevir 2010 [Grazoprevir 100 mg uuration. 12] [Grazoprevir 2010 [Gra	
C-EDGE nead-2-nead Weeks Genotype 10. 99% Age 540 years 100% (377)	
$\begin{bmatrix} (104/105) \\ (104/105) \end{bmatrix} = \begin{bmatrix} (104/105) \\ (104/105) \end{bmatrix} = \begin{bmatrix} (104/105) \\ (104/105) \end{bmatrix} = \begin{bmatrix} (104/105) \\ (104/105) \\ (104/105) \end{bmatrix} = \begin{bmatrix} (104/105) \\ (104/105) \\ (104/105) \\ (104/105) \end{bmatrix} = \begin{bmatrix} (104/105) \\ (104/1$	
Multinational (Europe 12 weeks post- 12 we	
$\frac{12}{12} = \frac{1000}{100} = \frac{1000}$	
Fair (100/107)	
Treatment-experienced: 100% (29/29)	

Author year	Treatment Regimen	Treatment			
Country	(1x/day unless	Duration and	Overall SVR	Genotype SVR	
Quality	otherwise noted)	Assessments	Results	Results	Other Subgroup SVR Results
Sulkowski 2014 ¹⁶¹	A. Sofosbuvir 400 mg +	Treatment	A vs. B	NR	NR
A1444040 Study	daclatasvir 60 mg	duration: 12	SVR: 100% (41/41)		
U.S.	B. Sofosbuvir 400 mg +	weeks	vs. 95% (39/41)		
Fair	daclatasvir 60 mg +	Timing of			
	ribavirin	assessment:			
		12 weeks post-			
		treatment			
Sulkowski 2015 ¹⁶⁰	A. Grazoprevir 100 mg	Treatment	A vs. B	A vs. B	NR
C-WORTHY	+ elbasvir 50 mg	duration: 12	SVR: 98% (43/44)	Genotype 1: 98%	
Australia, Canada,	B. Grazoprevir 100 mg	weeks	vs. 93% (79/85)	(43/44) vs. 93%	
Denmark France, Hungary,	+ elbasvir 50 mg +	Timing of		(79/85)	
Israel, New Zealand, Puerto	ribavirin	assessment:			
Rico, Spain, Sweden,		12 weeks post-			
Turkey, U.S.		treatment			
Fair					
Toyoda 2018 ¹⁹⁹	Glecaprevir 300 mg +	Treatment	SVR: 98% (88/90)	Genotype 2: 98%	NR
CERTAIN-2 (Arm A only)	pibrentasvir 120 mg	duration: 8		(88/90)	
Japan		weeks			
Fair		I iming of			
		assessments:			
		12 Weeks post-			
Wakad 2016 ¹⁶²	Ombitaavir 25 mg I	Treatment	SV/D: 049/ (04/100)	Construct 1: 049/	ND
		duration: 12	3VR. 94% (94/100)	Genotype 4. 94%	
	ritopovir 100 mg + 1000			(94/100)	
Good	to 1200 mg ribavirin	Timing of			
0000	to 1200 mg nbavinin	accessment.			
		12 wooks nost-			
		treatment			
Wei 2018 ¹⁶³	l edinasvir 90 ma +	Treatment	SVR: 100%	Genotype 1: 100%	Treatment-naïve: 100% (106/106)
China	sofosbuvir 400 mg +	duration: 12	(206/206)	(206/206)	Treatment-experienced: 100% (100/100)
Fair	conception rooming r	weeks	(200,200)	200,200)	
		Timing of			
		assessments.			
		12 weeks post-			
		treatment			

Author year	Treatment Regimen	Treatment			
Country	(1x/day unless	Duration and	Overall SVR	Genotype SVR	
Quality	otherwise noted)	Assessments	Results	Results	Other Subgroup SVR Results
Wei 2019a ¹⁶⁴	A. Elbasvir 50 mg +	Treatment	SVR-12: 94%	SVR-12	SVR-12
C-CORAL (Genotype 1 and	grazoprevir 100 mg	duration: 12	(459/486)	Genotype 1a: 92%	Male: 96% (207/216)
4 only)	(n=326)	weeks	SVR-24: 94%	(34/37)	Female: 93% (252/270)
Multinational (Australia,	B. Placebo (n=123;	Timing of	(458/486)	Genotype 1b: 98%	Asian: 93% (325/350)
China, Korea, Russia,	harms assessment	assessments:		(382/389)	White: 99% (133/135)
Taiwan, Thailand, Vietnam)	only)	12 weeks post-		Genotype 1-other:	Other: 1005 (1/1)
Good		treatment		100% (6/6)	Hispanic/Latino: 100% (5/5)
				Genotype 4: 100%	Non-Hispanic/Latino: 94% (454/481)
				(3/3)	Age <65 years: 95% (420/444)
				SVR-24	Age ≥65 years: 93% (39/42)
				Genotype 1a: 92%	No cirrhosis: 95% (375/396)
				(34/37)	Cirrhosis: 93% (84/90)
				Genotype 1b: 98%	SVR-24
				(381/389)	Male: 95% (206/216)
				Genotype 1-other:	Female: 93% (252/270)
				100% (6/6)	Asian: 93% (324/350)
				Genotype 4: 100%	White: 99% (133/135)
				(3/3)	Other: 1005 (1/1)
					Hispanic/Latino: 100% (5/5)
					Non-Hispanic/Latino: 94% (453/481)
					Age <65 years: 95% (420/444)
					Age ≥65 years: 91% (38/42)
					No cirrhosis: 95% (375/396)
					Cirrhosis: 93% (84/90)
Wei 2019b ¹⁶⁵	Sofosbuvir 400 mg +	Treatment	SVR: 97% (362/375)	Genotype 1a: 100%	Male: 94% (186/197)
Multinational (China,	velpatasvir 100 mg	duration: 12		(22/22)	Female: 99% (176/178)
Malaysia, Singapore,		weeks		Genotype 1b: 100%	Age <65 years: 96% (340/353)
Thailand, Vietnam)		Timing of		(107/107)	Age ≥65 years: 100% (22/22)
Fair		assessments:		Genotype 2: 100%	No cirrhosis: 98% (302/308)
		12 weeks post-		(64/64)	Cirrhosis: 90% (60/67)
		treatment		Genotype 3a and	Treatment-naive: 97% (297/307)
				unconfirmed subtype:	Treatment-experienced: 96% (65/68)
				95% (40/42)	
				Genotype 3b: 76%	
				(32/42)	
				Genotype 6: 99%	
				(97/98)	

Author year	Treatment Regimen	Treatment			
Country	(1x/day unless	Duration and	Overall SVR	Genotype SVR	Other Subgroup SV/D Depute
Quality	otherwise noted)	Assessments	Results	Results	Other Subgroup SVR Results
Zeuzem 2018 ¹⁶⁷	A. Glecaprevir 300 mg	Treatment	A vs. B vs. C	Genotype 3a: 95%	Male: 93% (86/92) vs. 93% (112/121) vs. 92% (48/52)
ENDURANCE-3 (same	+ pibrentasvir 120 mg, 8	duration: 8 to	SVR-12: 95%	(148/156) vs. 96%	Female: 97% (63/65) vs. 98% (110/112) vs. 100% (63/63)
publication as	weeks	12 weeks	(149/157) vs. 95%	(220/230) vs. 97%	
ENDURANCE-1)	B. Glecaprevir 300 mg		(222/233) vs. 97%	(111/115)	Black race: 100% (3/3) vs. 100% (4/4) vs. 75% (3/4)
Fair	+ pibrentasvir 120 mg,	Timing of	(111/115)	Other genotype 3:	Not Black race: 95% (146/154) vs. (218/229) vs. 97% (108/111)
	12 weeks	assessments:	SVR-24: 91%	100% (1/1) vs. 67%	
	3. Sofosbuvir 400 ma +	12 and 24	(143/157) vs. 92%	(2/3) vs. NA	Age <65 years: 95% (144/152) ys. 95% (213/224) ys. 96%
	daclatasvir 60 mg. 12	weeks post-	(214/233) vs. 96%	() -	(107/111)
	weeks	treatment	(110/115)		Age >65 years: 100% (5/5) vs. 100% (9/9) vs. 100% (4/4)
			(
					People who inject drugs (recent or history): 94% (98/104) vs
					$93\% (139/149) v_{S} 96\% (70/73)$
					Not people who inject drugs: 96% (51/53) vs. 90% (83/84) vs.
					$\alpha_{8\%}$ (41/42)
					50 /8 (41/42)
					No current opioid substitution therapy: 94% (119/126) vs. 96%
					(188/105) vg 0.6% (0.1/08)
					(100, 130) vs. 30/0 (34/30) Current onicid substitution thereps: $070((20/21))$ vs. $000((24/20))$
					Current opioid substitution therapy: 97% (30/31) VS. 90% (34/38)
					vs. 100% (1//1/)

*Excluding patients who withdrew or were lost to follow up.

†Based on total study population (treatment naïve and experienced combined).

Abbreviations: IFN = interferon; IVDU = injection drug use; NR = not reported; SVR = sustained virologic response; U.K. = United Kingdom; U.S. = United States. Study names are not acronyms.

Author year	Treatment Regimen			E
Country	(1X/day unless otherwise noted)	Clinical Outcomes	Adverse Events	Funding
Abergel 2016a ¹⁴² France <i>Fair</i>	Ledipasvir 90 mg + sofosbuvir 400 mg	Mortality: 0% (0/21)	Adverse EventsEntire study cohort (n=44; 23% cirrhosis)Any adverse event: 71% (31/44)Serious adverse events: 0%Withdrawal due to adverse events: 0%Headache: 25% (11/44)Fatigue: 20% (9/44)Nausea: 9% (4/44)Diarrhea: 9% (4/44)Hemoglobin 10.0 to 10.9 g/dL: 2% (1/44)Adverse Events: 0%	Gilead
Abergel 2016b ¹⁴¹ France <i>Good</i>	Ledipasvir 90 mg + sofosbuvir 400 mg	Mortality: 0% (0/22)	Bilirubin >1.0-1.5x ULN: 5% (2/44) Entire study cohort (n=41; 22% cirrhosis) Any adverse event: 80% (33/41) Serious adverse events: 2% (1/41; worsening depression) Withdrawal due to adverse events: 0% Headache: 27% (11/41) Fatigue: 10% (4/41) Diarrhea: 7% (3/41) Hemoglobin 100-109 g/dL: 2% (1/41) Bilirubin >1.0-1.5 ULN: 10% (4/41)	Gilead

Author year	Treatment Regimen			
Country	(1x/day unless			Funding
Quality	otherwise noted)	Clinical Outcomes	Adverse Events	Source
Afdahl 2014 ¹⁸⁵	A. Ledipasvir 90 mg +	NR	A vs. B	Gilead
ION-1	sofosbuvir 400 mg		12-week intervention group	
U.S. and Europe	B. Ledipasvir 90 mg +		Any adverse event: 79% (169/214) vs. 85% (185/217)	
Fair	sofosbuvir 400 mg +		Serious adverse event*: 0.5% (1/214) vs. 3% (7/217)	
	ribavirin		Withdrawal due to adverse events: 0% vs. 0%	
			Headache: 25% (53/214) vs. 23% (49/217)	
			Fatigue: 21% (44/214) vs. 36% (79/217)	
			Nausea: 11% (24/214) vs. 17% (37/217)	
			Diarrhea: 11% (24/214) vs. 8% (18/217)	
			Insomnia: 8% (17/214) vs. 21% (45/217)	
			Anemia: 0% vs. 12% (25/217)	
			Rash: 7% (16/214) vs. 10% (21/217)	
			24-week intervention group	
			Any adverse event: 82% (178/217) vs. 92% (200/217)	
			Serious adverse event*: 8% (8% (18/217) vs. 3% (7/217)	
			Withdrawal due to adverse events: 2% (4/217) vs. 3% (6/217)	
			Headache: 24% (54/217) vs. 30% (65/217)	
			Fatigue: 24% (24% (53/217) vs. 38% (82/217)	
			Nausea: 13% (29/217) vs. 15% (32/217)	
			Diarrhea: 11% (24/217) vs. 6% (14/217)	
			Insomnia: 12% (26/217) vs. 22% (47/217)	
			Anemia: 0% vs. 10% (22/217)	
			Rash: 7% (16/217) vs. 12% (25/217)	
Ahmed 2018 ¹⁹⁵	Ledipasvir 90 mg +	NR	Any adverse event: 26% (26/100)	NR
Egypt	sofosbuvir 400 mg		Headache: 2% (2/100)	
Fair			Fatigue: 18% (18/100)	
			Nausea: 2% (2/100)	
			Diarrhea: 1% (1/100)	
			Insomnia: 2% (2/100)	
Andreone 2014 ¹⁸⁶	A. Ombitasvir 25 mg +	NR	A vs. B	AbbVie
PEARL-II	paritaprevir 150 mg +		Any adverse event: 77.9% (74/95) vs. 79% (72/91)	
Austria, Belgium, Italy,	ritonavir 100 mg +		Withdrawals due to adverse events: 0% (0/95) vs. 2% (2/91)	
The Netherlands,	dasabuvir 250 mg		Serious adverse events (Pancreatitis, cellulitis, nephrolithiasis, osteoarthritis): 2%	
Portugal, Puerto Rico,	2x/day		(2/95) vs. 2% (2/91)	
Sweden, Switzerland,	B. Ombitasvir 25 mg +		Headache: 23.3% (22/95) vs. 24.2% (22/91)	
U.S.	paritaprevir 150 mg +		Fatigue: 15.8% (15/95) vs. 31.9% (29/91)	
Fair	ritonavir 100 mg +		Nausea: 6.3% (6/95) vs. 20.9% (19/91)	
	dasabuvir 250 mg		Diarrhea: 12.6% (12/95) vs. 13.2 (12/91)	
	2x/day + ribavirin		Anemia: 0% (0/95) vs. 11% (10/91)	
			Rash: 1% (1/95) vs. 9% (8/91)	

Author year	Treatment Regimen			
Country	(1x/day unless			Funding
Quality	otherwise noted)	Clinical Outcomes	Adverse Events	Source
Asselah 2018 ¹⁹⁶ SURVEYOR II Part 4, Multinational (Asia, Europe, U.S. [specific countries NR]) <i>Fair</i>	Glecaprevir 300 mg + pibrentasvir 120 mg	NR	Any adverse event: 63% (128/203) Serious adverse events (cholecystitis, urosepsis): 1% (2/203) Withdrawal due to adverse events: 0% (0/203) Headache: 18% (37/203) Fatigue: 14% (28/203) Nausea: 11% (23/203)	AbbVie
Asselah 2019 ¹⁴³ ENDURANCE-5 Multinational (Australia, Belgium, Canada, France, New Zealand, Singapore, South Africa, Vietnam, U.S.) <i>Fair</i>	Glecaprevir 300 mg + pibrentasvir 120 mg	Mortality: 0% (0/23)	Total population (n=84, genotype 5 and 6 combined) Any adverse event: 55% (46/84) Serious adverse events (gastric ulcer, pyelonephritis, giardiasis and depression, pulmonary tuberculosis, viral infection): 6% (5/84) Withdrawal due to Adverse events: 0% (0/84) Headache: 13% (11/84) Fatigue:13% (11/84)	AbbVie
Asselah 2019 ¹⁴³ ENDURANCE-6 (same publication as ENDURANCE-5) <i>Fair</i>	See Asselah 2019 ENDURANCE-5	Mortality: 0% (0/61)	See Asselah 2019 ENDURANCE-5	See Asselah 2019 ENDURANCE-5
Brown 2018 ¹⁴⁴ C-SCAPE (Genotype 4 only) Multinational (Australia, Belgium, France, Israel, Spain, U.K., U.S.) <i>Fair</i>	A. Elbasvir 50 mg + grazoprevir 100 mg (n=10) B. Elbasvir 50 mg + grazoprevir 100 mg + ribavirin (n=10)	Mortality: 0% (0/20)	Total population (genotypes 2, 4, 5, 6) Any adverse event: 79% (15/19) vs. 95% (18/19) Serious adverse events: 0% (0/19) vs. 0% (0/19) Withdrawal due to adverse events: 5% (1/19) vs. 0% (0/19) Headache: 26% (5/19) vs. 32% (6/19) Fatigue: 16% (3/19) vs. 26% (5/19) Nausea: 5% (1/19) vs. 11% (2/19) Asthenia: 21% (4/19) vs. 16% (3/19)	Merck
Chayama 2018 ¹⁹⁷ CERTAIN-1 (Arm A only) Japan <i>Fair</i>	Glecaprevir 300 mg + pibrentasvir 120 mg	NR	Any adverse event: 57% (74/129) Serious adverse events: 0% (0/129) Withdrawal due to adverse events: 0% (0/129) Headache: 5% (6/129) Rash: 2% (3/129)	AbbVie
Chuang 2016 ¹⁴⁵ Taiwan <i>Fair</i>	Ledipasvir 90 mg + sofosbuvir 400 mg	Total population (treatment-naïve and treatment-experienced) Mortality: 0% (0/85)	Total population (treatment-naïve and treatment-experienced) Any adverse event: 60% (51/60) Withdrawals due to adverse events: 1% (1/85) Headache: 14% (12/85) Fatigue: 9% (8/85) Nausea: 6% (5/85)	Gilead

Author year	Treatment Regimen			Funding
Country Quality	otherwise noted)	Clinical Outcomes	Adverse Events	Source
Dore 2016 ¹³⁷ MALACHITE-1 Australia, Canada, Europe, South America <i>Good</i>	Genotype 1a A. Ombitasvir 25 mg + paritaprevir 150 mg + ritonavir 100 mg + dasabuvir 250 mg 2x/day + weight-based ribavirin B. Telaprevir 750 mg 3x/day + subcutaneous pegylated IFN 180 ug 1/week + weight-based ribavirin Genotype 1b C. Ombitasvir 25 mg + paritaprevir 150 mg + ritonavir 100 mg + dasabuvir 250 mg 2x/day + weight-based ribavirin D. Ombitasvir 25 mg + paritaprevir 150 mg + ritonavir 100 mg + dasabuvir 250 mg 2x/day + weight-based ribavirin D. Ombitasvir 25 mg + paritaprevir 150 mg + ritonavir 100 mg + dasabuvir 250 mg 2x/day E. Telaprevir 750 mg 3x/day + subcutaneous pegylated IFN 180 ug 1/week + weight-based ribavirin	Genotype 1a A vs. B SF-36 mental component score, mean change from baseline at 12 weeks post- treatment: -1.1 (SD 12) vs2.1 (SD 10.1) SF-36 physical component score, mean change from baseline at 12 weeks post- treatment: 3.1 (SD 8.7) vs. 0.7 (SD 7.6) Genotype 1b C vs. D vs. E SF-36 mental component score, mean change from baseline at 12 weeks post- treatment: 1.9 (SD 9.6) vs. 1.4 (SD 8.1) vs0.3 (SD 10.3) SF-36 physical component score, mean change from baseline at 12 weeks post- treatment: 2.3 (SD 5.3) vs. 2.5 (SD 5.7) vs. 1.0 (SD 8.4)	(A + C [with ribavirin]) vs. D (without ribavirin) vs. (B + E [telaprevir]) Any adverse event: 75% (115/153) vs. 49% (41/83) vs. 99% (74/75); (A+C) vs. (B+E): RR 0.76 (95% CI, 0.69 to 0.84); D vs. (B+E): RR 0.50 (95% CI, 0.40 to 0.62) Withdrawals due to adverse events: 1% (1/153) vs. 0% (0/83) vs. 8% (6/75); (A+C) vs. (B+E): RR 0.08 (95% CI, 0.01 to 0.67) Serious adverse events (one each: prostate cancer, overdose, anemia, cough, chest pain, hematochezia, retinopathy, toxic skin eruption, cellulitis): 1% (1/153) vs. 0% (0/83) vs. 12% (9/75); (A+C) vs. (B+E): RR 0.05 (95% CI, 0.007 to 0.42); D vs. (B+E): RR 0.05 (95% CI, 0.003 to 0.80) Headache: 27% (41/153) vs. 19% (16/83) vs. 31% (23/75); (A+C) vs. (B+E): RR 0.87 (95% CI, 0.57 to 1.34); D vs. (B+E): RR 0.63 (95% CI, 0.007 to 0.42); D vs. (B+E): RR 0.87 (95% CI, 0.57 to 1.34); D vs. (B+E): RR 0.63 (95% CI, 0.06 to 0.43) Nausea: 21% (32/153) vs. 8% (7/83) vs. 40% (30/75); (A+C) vs. (B+E): RR 0.45, (95% CI, 0.35 to 0.79); D vs. (B+E): RR 0.21 (95% CI, 0.10 to 0.43) Nausea: 21% (32/153) vs. (B+E): RR 0.3 (95% CI, 0.10 to 0.45) Anemia: 7% (10/153) vs. (B+E): RR 0.3 (95% CI, 0.10 to 0.45) Anemia: 7% (10/153) vs. (B+E): RR 0.3 (95% CI, 0.004 to 0.19) Rash: 8% (12/153) vs. 0% vs. 23% (17/75); (A+C) vs. (B+E): RR 0.37 (95% CI, 0.19 to 0.73); D vs. (B+E): RR 0.03 (95% CI, 0.000 to 0.42)	AbbVie

Author year	Treatment Regimen			E
	(1x/day unless	Clinical Outcomos	Advorso Evonts	Funding
Doro 2016 ¹³⁷	A Ombitasvir 25 mg i			
	A. Offibilasvii 25 filg +	SE-36 mental	A vs. D Any adverse event: 62% (63/101) vs. 01% (/3//7): PR 0.68 (05% CL 0.57 to 0.81)	ADDVIE
Australia Canada	ritopavir 100 mg +	component score, mean	Sarious adverse events (enilensy, anemia [2 neonle], addeminal nain, infectious	
Europe South America	dasabuvir 250 mg	component score, mean	diarrhea, staphylococcal : 1% (1/101) vs. 5% (11/17): PR 0.04 (05% CL 0.006 to	
Good	$2x/day \pm weight-based$	12 weeks nost-	0 32)	
8000	ribavirin	treatment: 0.8 (SD 8.0)	Withdrawal due to adverse events: 0% (0/101) vs. 11% (5/47); RR 0.04 (05% CI	
	B Telaprevir 750 mg	$v_{\rm S} = 1.5 (SD 7.5)$	0.002 to 0.76	
	3x/day + subcutaneous	SF-36 physical	Headache: 29% (29/101) vs. 45% (21/47): RR 0.64 (95% CL 0.41 to 1.00)	
	pegylated IFN 180 ug	component score mean	Eatique: 12% (12/101) vs. 26% (12/47): RR 0.47 (95% CL 0.23 to 0.96)	
	1/week + weight-based	change from baseline at	Nausea: 10% (10/101) vs. 43% (20/47); RR 0.23 (95% CL 0.12 to 0.46)	
	ribavirin	12 weeks post-	Insomnia: 6% (6/101) vs. 21% (10/47): RR 0.28 (95% CL 0.11 to 0.72)	
		treatment: 3.0 (SD 6.4)	Anemia: 3% (3/101) vs. 34% (16/47): RR 0.09 (95% CL 0.03 to 0.28)	
		vs1.3 (5.3)	Rash: 3% (3/101) vs. 17% (8/47); RR 0.06 (95% CI, 0.02 to 0.21)	
Everson 2015 (Part A) ¹⁴⁶	Part A (trial phase)	A vs. B vs. Ć vs. D vs. E	(A + C + E) vs. (B + D + F)	Gilead
U.S.	A. Sofosbuvir 400 mg +	vs. F	Any adverse event: 68% (52/77) vs. 70% (54/77)	
Good	velpatasvir 25 mg	Mortality: 0% (0/27) vs.	Withdrawal due to adverse events: 0% (0/77) vs. 0% (0/77)	
	(Genotype 1)	0% (0/28) vs. 0% (0/27)	Serious adverse events (not described): 3% (2/77) vs. 1% (1/77)	
	B. Sofosbuvir 400 mg +	vs. 0% (0/27) vs. 4%	Headache: 21% (16/77) vs. 18% (14/77)	
	velpatasvir 100 mg	(1/23) vs. 0% (0/22)	Fatigue: 25% (19/77) vs. 18% (14/77)	
	(Genotype 1)		Nausea: 13% (10/77) vs. 10% (8/77)	
	C. Sofosbuvir 400 mg +		Diarrhea: 6% (5/77) vs. 9% (7/77)	
	velpatasvir 25 mg		Constipation: 12% (9/77 vs. 8% (6/77)	
	(Genotype 3)		Insomnia: 4% (3/77) vs. 6% (5/77)	
	D. Sofosbuvir 400 mg +		Hemoglobin <100g/L: 0% vs. 0%	
	velpatasvir 100 mg		Bilirubin >2.5x ULN: 0% vs. 0%	
	(Genotype 3)		Rash: 5% (4/77) vs. 5% (4/77)	
	E. Sofosbuvir 400 mg +			
	veipatasvir 25 mg			
	(Genotype 2; 4-6)			
	F. Sotosbuvir 400 mg +			
	veipatasvir 100 mg			
	(Genotype 2; 4-6)			

Author year	Treatment Regimen			
Country	(1x/day unless			Funding
Quality	otherwise noted)	Clinical Outcomes	Adverse Events	Source
Feld 2014 ¹⁸⁷	A. Ombitasvir 25 mg +	NR	A vs. B	AbbVie
SAPPHIRE-1	paritaprevir 150 mg +		Any adverse event: 86% (414/473) vs. 73% (116/158); RR 1.19 (95% CI, 1.08 to	
Australia, New Zealand;	ritonavir 100 mg +		1.32)	
Austria, France,	dasabuvir 250 mg 2x		Withdrawal due to adverse event: 0.6% (3/473) vs. 0.6% (1/158); RR 1.00 (95% CI,	
Germany, Hungary,	day + weight-based			
Great Britain, Italy,	ribavirin		Serious adverse events (appendicitis, lobar pneumonia, cholecystitis, lumbar	
Spain, Sweden,	B. Placebo for 12 weeks		vertebral fracture in one patient each; aortic stenosis and postoperative wound	
Switzerland; Canada,	followed by open-label		infection in one; overdose and encephalopathy in one; mediastinal mass and non-	
U.S.	ombitasvir 25 mg +		small-cell lung cancer in one; acute respiratory failure and hypoxemia in one;	
Good	paritaprevir 150 mg +		abdominal pain, sinus tachycardia, diarrhea, chills, vomiting, nausea, and ventricular	
	ritonavir 100 mg +		extrasystoles in one; and anemia and noncardiac chest pain in one): 2% (10/4/3) vs.	
	dasabuvir 250 mg 2x		0%; RR 7.04 (95% CI, 0.42 to 120)	
	day + weight-based		Diarrnea: 14% (65/4/3) vs. 7% (11/158); RR 1.97 (95% CI, 1.07 to 3.64)	
	ribavirin		[Fatigue: 35% (164/4/3) vs. 29% (45/158); RR 1.22 (95% CI, 0.92 to 1.60)	
			Headache: 33% (156/4/3) vs. 2/% (42/158); RR 1.24 (95% CI, 0.93 to 1.66)	
			Nausea: 24% (112/4/3) vs. 13% (21/158); RR 1.78 (95% CI, 1.16 to 2.74)	
			Insomnia: 14% (66/4/3) vs. 8% (12/158); RR 1.84 (95% CI, 1.02 to 3.31)	
—			Rash: 11% (51/4/3) vs. 6% (9/158); RR 1.89 (95% CI, 0.95 to 3.76)	<u>.</u>
Feld 2015 ¹³⁹	A. Sofosbuvir 400 mg +	A vs. B		Gilead
ASTRAL-1	velpatasvir 100 mg	Mortality: 0.2% (1/624)	Any adverse event: 78% (485/624) vs. 77% (89/116); RR 1.01, 95% CI, 0.91 to 1.13	
U.S., Canada, Europe,	B. Placebo	vs. 0% (0/116)	Serious adverse events (19 events in 15 patients: abscess limb, acute myocardial	
Hong Kong		Mean change from	infarction, appendicitis, bronchitis, cellulitis, chronic obstructive pulmonary disease,	
Good		baseline in patient-	epilepsy, extremity necrosis, gastroenteritis, influenza, ligament sprain, lung cancer,	
		reported outcomes	mania, palpitations, rotatorcuff syndrome, small intestinal obstruction, sudden death	
		(composite SF-36,	from unknown cause, upper limb fracture, and vestibular neuronitis): 2% (15/624) vs.	
		FACIT-F, CLDQ-HCV,	0% (0/116); RR 5.80, 95% CI, 0.35 to 96	
		WPAI:SHP; scale 0 to	Withdrawals due to adverse events: 0.2% (1/624) vs. 2% (2/116); RR 0.09 (95% CI,	
		100), 24-weeks post-		
		treatment: 5.4; p<0.05	[Headache: 29% (182/624) vs. 28% (33/116); RR 1.03 (95% CI, 0.75 to 1.40)	
		for all individual	Fatigue: 20% (126/624) vs. 20% (23/116); RR 1.02 (95% CI, 0.68 to 1.52)	
		components except	Nausea: 12% (75/624) vs. 11% (13/116)	
		WPAI:SHP WORK	Diarrnea: 8% (48/624) vs. 7% (8/116); RR 1.12 (95% CI, 0.54 to 2.30)	
		productivity and	Insomnia: 8% (50/624) vs. 9% (11/116); RR 0.84 (95% CI, 0.45 to 1.57)	
		WPAI:SHP absenteeism	Hemoglobin <10 g/dL: 0.4% (2/624) vs. 0% (0/116); RR 2.21 (95% CI, 0.11 to 46)	

Author year	Treatment Regimen			
Country	(1x/day unless			Funding
Quality	otherwise noted)	Clinical Outcomes	Adverse Events	Source
Ferenci 2014 ¹⁸⁸	A. Ombitasvir 25 mg +	NR	A vs. B	AbbVie
PEARL III	paritaprevir 150 mg +		Any adverse event: 67.0% (140/209) vs. 80% (168/210)	
Austria, Belgium,	ritonavir 100 mg +		Serious adverse events (coronary artery disease, atrial fibrillation, nephrolithiasis,	
Hungary, Israel, Italy,	dasabuvir 250 mg		epididymitis, arthritis, breast lesion, uterine polyp, myalgia): 2% (4/209) vs. 2%	
Poland, Portugal,	2x/day		(4/210)	
Romania, Russia, Spain,	B. Ombitasvir 25 mg +		Withdrawal due to adverse events: none	
U.S.	paritaprevir 150 mg +		Headache: 23% (49/209) vs. 24% (51/210)	
Good	ritonavir 100 mg +		Fatigue: 23% (48/209) vs. 21% (45/210)	
	dasabuvir 250 mg		Nausea: 4% (9/209) vs. 23% (11/210)	
Same publication as	2x/day + ribavirin		Diarrhea: 6% (13/209) vs. 4% (9/210)	
PEARL IV			Rash: 3% (8/209) vs. 6% (12/210)	
Ferenci 2014 ¹⁸⁸	A. Ombitasvir 25 mg +	NR	Any adverse event: 82% (169/205) vs. 92.0% (92/100)	AbbVie
PEARL IV	paritaprevir 150 mg +		Serious adverse events (pancreatitis, anemia, intestinal obstruction, diverticulitis):	
Canada, U.K., U.S.	ritonavir 100 mg +		0.5% (1/205) vs. 3.0% (3/100)	
Good	dasabuvir 250 mg		Withdrawal due to adverse events: none	
	2x/day		Headache: 28% (58/205) vs. 25.0% (25/100)	
Same publication as	B. Ombitasvir 25 mg +		Fatigue: 35% (72/205) vs. 46.0% (46/100)	
PEARL III	paritaprevir 150 mg +		Nausea: 14% (28/205) vs. 21.0% (21/100)	
	ritonavir 100 mg +		Diarrhea: 16.1% (33/205) vs. 14.0% (14/100)	
	dasabuvir 250 mg		Rash: 5% (10/205) vs. 5% (5/100)	
	2x/day + ribavirin			
Foster 2015 ¹⁴⁷	A. Sofosbuvir 400 mg +	A vs. B	A vs. B	Gilead
ASTRAL-2	velpatasvir 100 mg	Mortality: 1% (2/134) vs.	Any adverse event: 69% (92/134) vs. 77% (101/132)	
U.S.	B. Sofosbuvir 400 mg +	0% (0/132)	Serious adverse events (pneumonia, enteritis, abdominal pain, arthralgia,	
Fair	ribavirin		depression): 1% (2/134) vs. 2% (2/132)	
			Withdrawals due to adverse events: 1% (1/134) vs. 0% (0/132)	
			Dyspepsia: 1% (1/134) vs. 4% (5/132)	
			Headache: 18% (24/134) vs. 22% (29/132)	
			Fatigue: 15% (20/134) vs. 35% (47/132)	
			Nausea: 10% (14/134) vs. 14% (19/132)	
			Grade 3 or 4 bilirubin elevation: 0% (0/134) vs. 0% (0/132)	
			Insomnia: 4% (6/134) vs. 14% (18/132	

Author year Treatment Regimen	
Country (1x/day unless	Funding
Quality otherwise noted) Clinical Outcomes Adverse Events	Source
Foster 2015 ¹⁴⁷ Same as Foster 2015 A vs. B A vs. B	Gilead
ASTRAL-3 ASTRAL-2 Mortality: 0% (0/278) vs. Any adverse event: 88% (245/277) vs. 95% (260/275)	
U.S. 0.7% (2/280) Serious adverse events (myocardial infarction, bursitis, cellulitis, cardiovasc	ular
Fair accident, cholecystitis, chronic obstructive pulmonary disease, depression, f	food
poisoning, gunshot wound, hematochezia, overdose, intervertebral disc prot	trusion,
Same publication as aneurysm, lung infection, ovarian cyst rupture, stenosis, infection, psychotic	disorder,
ASTRAL-2 rash): 2% (6/277) vs. 5% (15/275)	
Withdrawal due to adverse events: 0% (0/277) vs. 3% (9/275)	
Dyspepsia: 3% (9/277) vs. 11% (30/275)	
Headache: 32% (90/277) vs. 32% (89/275)	
Fatigue: 26% (71/277) vs. 38% (105/275)	
Nausea: 17% (46/277) vs. 21% (58/275)	
Insomnia: 11% (31/277) vs. 27% (74/275)	
Gane 2015 ¹⁴⁸ Ledipasvir 90 mg + Mortality: 0% (0/25) Any adverse event: 84% (21/25)	Gilead
New Zealand (Genotype sofosbuvir 400 mg Serious adverse events (not described): 4% (1/25)	
6 subset) Withdrawal due to adverse events: 0% (0/25)	
Fair Headache: 8% (2/25)	
Fatigue: 24% (6/25)	
Nausea: 0% (0/25)	
Diarrhea: 16% (4/25)	
Gastroenteritis: 0% (0/25)	
Vomiting: 0% (0/25)	
Hemoglibin 7.0 to <9.0 g/dL: 0% (0/25)	
AST elevation >5 to 10x ULIN: 4% (1/25)	
RdSII. 0% (2/23)	Cilaad
Single Lev Single Singl	Glieau
Serious adverse events (Habdoniyolysis, other serious adverse events (K)	. 170
(7/103)	
Norway, Switzerland	
Fair Nausea: 14% (14/103)	
Vomiting: 4% (4/103)	
Diarrhea: 4% (4/103)	
Insomnia: 9% (9/103)	

Author year	Treatment Regimen			
Country	(1x/day unless	Clinical Outcomes	Adverse Events	Funding
	Ombitasvir 25 mg +	Mortality: 3% (3/87)	Auverse event: 61% (53/87)	
	paritaprovir 150 mg +	Mortality. 378 (3/87)	Sarious adverse events (NR): 6% (5/87)	ADD VIE
Multinational (Australia	ritonavir 100 mg +		Withdrawal due to adverse events: 0% (0/87)	
Canada France New	dasabuvir 250 mg +		Headache: 5% (12/87)	
Zealand Norway	1000-1200 mg ribavirin		Fatigue: 10% (25/87)	
Switzerland)			Nausea: 8% (20/87)	
Fair			Vomiting: 4% (11/87)	
, an			Anemia: 5% (12/87)	
			Insomnia: 4% (11/87)	
Hezode 2015 ¹⁸⁹	Ombitasvir 25 mg +	NR	Any adverse event: 88% (37/42)	AbbVie
PEARL I (Treatment-	paritaprevir 150 mg +		Serious adverse events: 0%	
naïve population)	ritonavir 100 mg +		Withdrawal due to adverse events: 0%	
France, Hungary, Italy,	ribavirin (weight-based;		Headache: 33% (14/42)	
Poland, Romania, Spain,	dose NR)		Fatigue: 12% (5/42)	
Turkey, U.S.			Nausea: 17% (7/42)	
Good			Diarrhea: 14% (6/42)	
			Insomnia: 10% (4/42)	
See also Lawitz 2015 ¹⁵⁵			Hemoglobin <100 g/L: 2% (1/42)	
(PEARL I - Genotype 1b)			Total bilirubin, grade 3 elevation: 0%	
			ALT elevation >5x ULN and ≥2x baseline: 0%	
			AST elevation >5x ULN and ≥2x baseline: 0%	
Hezode 2015 ¹⁸⁹	Same as Hezode 2015	NR	Any adverse event: 88% (43/49)	AbbVie
PEARL I (Treatment	(Treatment naïve		Serious adverse events: 0%	
experienced population)	population)		Withdrawal due to adverse events: 0%	
France, Hungary, Italy,			Headache: 29% (14/49)	
Poland, Romania, Spain,			Fatigue: 18% (9/49)	
Turkey, U.S.			Nausea: 12% (6/49)	
Good			Diarrhea: 6% (3/49)	
0 1 1 1 0 0 1 5155			Insomnia: 16% (8/49)	
See also Lawitz 2015 ¹³⁵			Hemoglobin <100 g/L: 2% (1/49)	
(PEARL I - Genotype 1b)			I otal bilirubin, grade 3 elevation: 6% (3/49)	
			ALT elevation >5x ULN and ≥2x baseline: 0%	
			AST elevation >5x ULN and ≥2x baseline: 0%	

Author year	Treatment Regimen			
Country	(1x/day unless			Funding
Quality	otherwise noted)	Clinical Outcomes	Adverse Events	Source
Kowdley 2014a ¹⁹⁰ ION-3 U.S. <i>Fair</i>	Ledipasvir 90 mg + sofosbuvir 400 mg	NR	8-week intervention group Any adverse event: 67% (145/215) Serious adverse events (anaphylaxis, colitis, inadequately controlled diabetes, gastrointestinal hemorrhage, hypertension, pituitary tumor): 2% (4/215) Withdrawal due to adverse events: 0% Headache: 14% (30/215) Fatigue: 21% (45/215) Nausea: 7% (15/215) Diarrhea: 7% (15/215) Insomnia: 5% (11/215) Anemia: 1% (2/215) Rash: 1% (3/215) 12-week intervention group Any adverse event: 69% (149/216) Serious adverse events (abdominal pain, bile duct stone, hemothorax, hypoglycemia, intestinal perforation, mental illness, respiratory failure, rhabdomyolysis, traffic accident, bone injury, lung cancer): 2% (5/216) Withdrawal due to adverse events: 1% (2/216) Headache: 15% (33/216) Fatigue: 23% (49/216) Nausea: 11% (24/216) Diarrhea: 4% (9/216) Insomnia: 7% (15/216) Anemia: 1% (2/216) Rash: 2% (5/216)	Gilead
Kowdley 2014b ¹⁹¹ AVIATOR Australia, Canada, France, Germany, New Zealand, Puerto Rico, Spain, U.K., U.S. <i>Good</i>	A. Ombitasvir 25 mg + paritaprevir 150 mg + ritonavir 150 mg + dasabuvir 800 mg B. Ombitasvir 25 mg + paritaprevir 150 mg + ritonavir 100-150 mg + dasabuvir 800 mg + ribavirin 1000-1200 mg	NR	A vs. B Any adverse event: NR Serious adverse events (affective disorder, animal bite, arthralgia, acute cholecystitis, and facial paresis (occurring in one patient each); increased blood creatinine level and bronchitis occurring in the same patient; the cervicobrachial syndrome, neck pain, and osteoarthritis of the spine occurring in the same patient; lung disorder and pneumonia occurring in the same patient): 3% (2/79) vs. 1% (1/79) Withdrawals due to adverse events: 0% (0/79) vs. 3% (2/79) Headache: 19% (15/79) vs. 27% (21/79) Fatigue: 20% (16/79) vs. 28% (22/79) Nausea: 14% (11/79) vs. 28% (22/79) Diarrhea: 16% (13/79) vs. 13% (10/79) Grade 3 or 4 bilirubin elevation: 0% (0.79) vs. 5% (4/79) Grade 3 or 4 ALT elevation: 0% (0/79) vs. 1% (1/79) Anemia: 1% (1/79) vs. 9% (7/79)	AbbVie

Author year Country <i>Quality</i>	Treatment Regimen (1x/day unless otherwise noted)	Clinical Outcomes	Adverse Events	Funding Source
Kumada 2017 (Part 2 only) ¹⁵² Japan <i>Good</i>	Elbasvir 50 mg + grazoprevir 100 mg	Mortality: 0% (0/227)	Serious adverse events (not described): 5% (11/227) Withdrawal due to adverse events: 1% (3/227) Clinically significant adverse event: 4% (8/227)	Merck
Kumada 2015 ¹⁵¹ GIFT-1 (Substudy 1) Japan <i>Fair</i>	A. Ombitasvir 25 mg + paritaprevir 150 mg + ritonavir 100 mg (double-blind treatment) B. Placebo for 12 weeks, followed by ombitasvir 25 mg + paritaprevir 150 mg + ritonavir 100 mg (open- label treatment)	A vs. B Mortality: 0% (0/255 vs. 0% (0/106)	A vs. B (placebo-controlled phase only) Any adverse event: 68.8% (148/215) vs. 56.6% (60/106); RR 1.22 (95% CI, 1.01 to 1.47) Serious adverse events (not described): 3.3% (7/215) vs. 1.9% (2/106); RR 1.73 (95% CI, 0.36 to 8.16) Withdrawals due to adverse events: 0.9% (2/215) vs. 0% (0/106); RR 2.48 (95% CI, 0.12 to 51) Headache: 8.8% (19/215) vs. 9.4% (10/106); RR 0.94 (95% CI, 0.45 to 1.94) Nausea: 4.3% (9/215) vs. 3.8% (4/106); RR 1.11 (95% CI, 0.35 to 3.52) Hemoglobin <8g/dL: 0% vs. 0%	AbbVie
Kwo 2016 ¹⁵³ OPTIMIST-1 Canada, U.S. <i>Fair</i>	Simeprevir 150 mg + sofosbuvir 400 mg	Mortality: 0% (0/155) Quality of life, mean change from baseline (among 141/155 with SVR) - -HCV-SIQv4 overall body symptom score - 3.9 (SE 0.96) -Fatigue Severity Scale: -0.5 (SE 0.15) -Center for Epidemiologic Studies- Depression Scale: -0.2 (SE 0.73) -EQ-5D VAS: 4.1 (SE 1.4)	Any adverse event: 66% (103/155) Serious adverse events (colitis): 1% (1/155) Withdrawals due to adverse events: 0% (0/155) Nausea: 15% (23/155) Headache: 14% (22/155) Fatigue: 12% (19/155) Increased bilirubin: 1% (1/155) Rash: 6% (10/155)	Janssen
Lalezari 2015 ¹⁹² U.S. <i>Fair</i>	Ombitasvir 25 mg + paritaprevir 150 mg + ritonavir 100 mg + dasabuvir 250 mg 2x/day + ribavirin 1000- 1200 mg	NR	Any adverse event: 92.1% (35/38) Serious adverse events (cerebrovascular accident, sarcoma, acute myeloid leukemia): 7.9% (3/38) Withdrawal due to adverse events: 2.6% (1/38) Headache: 31.6% (12/38) Fatigue: 47.4% (18/38) Nausea: 50% (19/38) Vomiting: 10.5% (4/38) Insomnia: 18.4% (7/38) Anemia: 10.5% (4/38) Rash: 15.8% (6/38)	AbbVie

Author year	Treatment Regimen					
Country	(1x/day unless			Funding		
Quality	otherwise noted)	Clinical Outcomes	Adverse Events	Source		
Lawitz 2014a ¹⁵⁴	A. Simeprevir 150 mg +	Mortality: 0% (0/81)	Any adverse event: 79% (11/14) vs. 89% (24/27)	Janssen		
COSMOS	sofosbuvir 400 mg		Serious adverse events: 0% vs. 0%			
U.S.	B. Simeprevir 150 mg +		Withdrawals due to adverse events: 0% vs. 0%			
Fair	sofosbuvir 400 mg +		Anemia: 0% vs. 0%			
	ribavirin		Rash: 7% (1/14) vs. 22% (6/27)			
Lawitz 2014b ¹⁹³	A. Ledipasvir 90 mg +	NR	8-week intervention group	Gilead		
LONESTAR (Cohort A)	sofosbuvir 400 mg, 8		Any adverse event: 45% (9/20)			
U.S.	weeks		Serious adverse events: 0%			
Fair	B. Ledipasvir 90 mg +		Withdrawal due to adverse events: 0%			
	sofosbuvir 400 mg, 12		Headache: 10% (2/20)			
	weeks		Nausea: 10% (2/20)			
	C. Ledipasvir 90 mg +		Rash: 5% (1/20)			
	sofosbuvir 400 mg +		12-week intervention group			
	ribavirin		Any adverse event: 42% (8/19)			
			Serious adverse events (exacerbation of peptic ulcer disease): 5% (1/19)			
			Withdrawal due to adverse events: 0%			
			Headache: 0%			
			Nausea: 5% (1/19)			
			Rash: 0%			
Lawitz 2015 ¹⁵⁵	Ombitasvir 25 mg +	Mortality: 0% (0/82)	Any adverse event: 76.8% (63/82)	AbbVie		
PEARL-1	paritaprevir 150 mg +		Serious adverse events (unclear; NR according to treatment group): 2.4% (2/82)			
France, Hungary, Italy,	ritonavir 100 mg		Severe adverse events: 2.4% (2/82)			
Poland, Puerto Rico,			Withdrawals due to adverse events: 0% (0/82)			
Romania, Spain, Turkey,			Asthenia: 6.1% (5/82)			
U.S.			Diarrhea: 7.3% (6/82)			
Fair			Dry skin: 8/5% (7/82)			
			Fatigue: 7.2% (6/82)			
			Headache: 29.3% (24/82)			
			Hypertension: 1.2% (1/82)			
			Nausea: 9.8% (8/82)			
			Pruritus: 7.3% (6/82)			
Lim 2016 ¹⁵⁶	Ledipasvir 90 mg +	Includes all patients	Includes all patients (n=93, including treatment experienced, 28% cirrhosis)	Gilead		
Korea	sofosbuvir 400 mg	(n=93, including	Any adverse event: 49% (46/93)			
Fair		treatment experienced,	Serious adverse event (contact dermatitis, erysipelas, inguinal hernia): 3% (3/93)			
		28% cirrhosis)	Withdrawals due to adverse events: (1/93)			
		Mortality: 0% (093)	Headache: 8% (7/93)			
			Fatigue: 6% (6/93)			

Author year	Treatment Regimen			
Country	(1x/day unless			Funding
Quality	otherwise noted)	Clinical Outcomes	Adverse Events	Source
Nelson 2015 ¹⁵⁷	Daclatasvir 60 mg +	Mortality: 0% (0/152)	Any adverse event: NR	Bristol-Myers
ALLY-3	sofosbuvir 400 mg		Serious adverse events (gastrointestinal hemorrhage): 0.7% (1/152)	Squibb
U.S.			Headache: 20% (30/152)	
Fair			Fatigue: 19% (29/152)	
			Nausea: 12% (18/152)	
			Diarrhea: 9% (13/152)	
			Insomnia: 6% (9/152)	
Pianko 2015 ¹⁵⁸	A. Sofosbuvir 400 mg +	Includes Genotype 3	Includes Genotype 3 patients with cirrhosis and Genotype 1 patients (n=80; 41%	Gilead
Australia, New Zealand,	velpatasvir 100 mg	patients with cirrhosis	cirrhosis)	
U.S.	(Group 3)	and Genotype 1 patients	A vs. B	
Fair	B. Sofosbuvir 400 mg +	A vs. B	Any adverse event: 79% (63/80) vs. 86% (69/80)	
	velpatasvir 100 mg +	Mortality: 0% (0/80)	Serious adverse events (group A only: cholecystitis, suicide, rib fracture, contusion;	
	ribavirin (Group 4)		group B not described): 5% (4/80) vs. 4% (3/80)	
			Withdrawal due to adverse events: 0% (0/80) vs. 0% (0/80)	
			Headache: 23% (18/80) vs. 30% (24/80)	
			Fatigue: 24% (19/80) vs. 34% (27/80)	
			Nausea: 9% (7/80) vs. 23% (18/80)	
			Diarrhea: 11% (9/80) vs. 5% (4/80)	
			Insomnia: 8% (6/80) vs. 20% (16/80)	
			Rash: 3% (2/80) vs. 11% (9/80)	
Poordad 2017 ¹⁹⁴	A. Glecapravir 200 mg +	NR	A vs. B vs. C	AbbVie
MAGELLAN-1	pibrentasvir 80 mg		Any adverse event: 83.3% (5/6) vs. 81.8% (18/22) vs. 86.4% (19/22)	
U.S.	B. Glecapravir 200 mg +		Serious adverse events (fracture, breast cancer): 16.7% (1/6) vs. 0% vs. 4.5% (1/22)	
Fair	pibrentasvir 120 mg			
	C. Glecapravir 200 mg +		Headache: 16.7% (1/6) VS. 36.4% (8/22) VS. 22.7% (5/22)	
	pibrentasvir 120 mg +		[Fatigue: 16.7% (1/6) vs. 18.2% (4/22) vs. 36.4% (8/22)	
	ribavirin		Nausea: 16.7% (1/6) VS. 13.6% (3/22) VS. 27.3% (6/22)	
			Insomnia: 0% vs. 0% vs. 27.3% (6/22)	
			ALT > 3X ULIN. 0% VS. 0% VS. 0%	
			AST >3X ULIN. 0% VS. 0% VS. 0%	
			Dilliudiii >3X ULIN. 0% VS. 0% VS. 0% Hemoglabia $_{2}$ 10 g/dl : 0% vs. 0%	
Bett Junier 2010 (Croup	Declatoovir 60 mg J	Mortality: $09(.0/127)$	Hendopho: 15% (10/65)	Fodorol
A declates vir/	Dacialasvii 60 mg	$1001121112 \cdot 0\% (0/127)$	Estimute: 220/ (15/65)	Feueral
cofochuvir arm) ¹⁵⁹	SUIUSDUVII 400 IIIg		raugue. 23/0 (10/00)	São Poulo
Brozil			Nausta, 070 (4703)	Sau Faulu
Good			$\frac{1}{100}$	
9000			Rach: 2% (1/65)	
			rasii. 270 (1703)	

Author year	Treatment Regimen			
Country	(1x/day unless			Funding
Quality	otherwise noted)	Clinical Outcomes	Adverse Events	Source
Pott-Junior 2019 (Group	Simeprevir 150 mg +	See Pott-Junior 2019	Headache: 28% (17/60)	See Pott-Junior
B - simeprevir/ sofosbuvir	sofosbuvir 400 mg	Group A	Fatigue: 28% (17/60)	2019 Group A
arm) ¹⁵⁹	_		Nausea: 13% (8/60)	
Brazil			Vomiting: 5% (3/60)	
Good			Insomnia: 10% (6/60)	
			Rash: 10% (6/60)	
Sperl 2016 ¹⁹⁸ and Ng	Elbasvir 50 mg +	SF-36 physical	Any adverse event: 52% (67/129)	Merck
2018 ¹³⁸	grazoprevir 100 mg	component score, mean	Serious adverse events (type of adverse event NR): 0.8% (1/129)	
C-EDGE Head-2-Head		change from baseline:	Withdrawal due to adverse events: 0%	
(elbasvir/grazoprevir arm		2.0		
only)		SF-36 mental		
Multinational (Europe,		component score, mean		
Turkey)		change from baseline:		
Fair		2.0		
		FACIT-F score, mean		
		change from baseline:		
		1.75		
Sulkowski 2014 ¹⁶¹	A. Sofosbuvir 400 mg +	Mortality: 0% (0/41)	Any adverse event: 93% (38/41)	Bristol-Myers
A1444040 Study	daclatasvir 60 mg		Serious adverse events (psychiatric disorder): 2% (1/41)	Squibb; Gilead
U.S.	B. Sofosbuvir 400 mg +		Withdrawal due to adverse events: 0%	
Fair	daclatasvir 60 mg +		Headache: 34% (14/41)	
	ribavirin		Fatigue: 39% (16/41)	
			Nausea: 20% (8/41)	
			Vomiting: 2% (1/41)	
			Diarrhea: 5% (2/41)	
			Insomnia: 10% (4/41)	
2 ··· ··· · · · · · · · · · · · · · · ·			Grade 3 or 4 lab abnormality: 0%	
Sulkowski 2015 ¹⁶⁰	A. Grazoprevir 100 mg +	Mortality: 0% (0/44)	Any adverse event: NR; drug-related adverse events 56% (24/43 ⁺)	Merck
C-WORTHY	elbasvir 50 mg		Serious adverse events: 0%	
Australia, Canada,	B. Grazoprevir 100 mg +		Withdrawal due to adverse events: 0%	
Denmark France,	elbasvir 50 mg +		Headache: 35% (15/43)	
Hungary, Israel, New	ribavirin		Fatigue: 23% (10/43)	
Zealand, Puerto Rico,			Nausea: 10% (7/43)	
Spain, Sweden, Turkey,			Diarmea: 12% (5/43)	
U.S.			Hemoglobin <8.5 g/dL: U%	
Fair			ALT >2.5X Daseline value: 0%	
			AST >2.5X Daseline value: 0%	
	1		Bilirudin >5x daseline value: 0%	

Author year	Treatment Regimen			
Country	(1x/day unless			Funding
Quality	otherwise noted)	Clinical Outcomes	Adverse Events	Source
Toyoda 2018 ¹⁹⁹	Glecaprevir 300 mg +	NR	Any adverse event: 48% (43/90)	AbbVie
CERTAIN-2 (Arm A only)	pibrentasvir 120 mg		Serious adverse events (pneumothorax, unstable angina): 2% (2/90)	
Japan			Withdrawal due to adverse events: 1% (1/90)	
Fair			Headache: 7% (6/90)	
			Nausea: 3% (3/90)	
			Anemia: 0% (0/90)	
Waked 2016 ¹⁶²	Ombitasvir 25 mg +	Mortality: 1% (1/100)	Any adverse event: 80% (80/100)	AbbVie
AGATE-II	paritaprevir 150 mg +		Serious adverse events (deep venous thrombosis, cardiac arrest): 2% (2/100)	
Egypt	ritonavir 100 mg + 1000-		Headache: 41% (41/100)	
Good	1200 mg ribavirin		Fatigue: 35% (35/100)	
			Dyspepsia: 17% (17/100)	
			Insomnia: 9% (9/100)	
			Grade 2 hemoglobin abnormality: 7% (7/100)	
			Grade ≥2 total bilirubin elevation: 19% (19/100)	
Wei 2018 ¹⁶³	Ledipasvir 90 mg +	Mortality: 0% (0/206)	Any adverse event: 58% (120/206)	Gilead
China	sofosbuvir 400 mg +		Serious adverse events (epicondylitis, asthma, bone contusion): 1% (3/206)	
Fair			Withdrawal due to adverse events: 0% (0/206)	
Wei 2019a ¹⁶⁴	A. Elbasvir 50 mg +	A vs. B	A vs. B	Merck
C-CORAL (Genotype 1	grazoprevir 100 mg	Mortality: 0.2% (1/486)	Any adverse event: 47% (230/486) vs. 50% (62/123)	
and 4 only)	(n=326)	vs. 0% (0/123)	Serious adverse events (suicide, contusion, Evans syndrome, lymphoma, enteritis vs.	
Multinational (Australia,	B. Placebo (n=123;		influenza, fracture): 2% (8/486) vs. 2% (2/123)	
China, Korea, Russia,	harms assessment only)		Withdrawal due to adverse events: 0.6% (3/486) vs. 2% (2/123)	
Taiwan, Thailand,			Headache: 6% (27/486) vs. 5% (6/123)	
Vietnam)			Fatigue: 5% (22/486) vs. 7% (9/123)	
Good				
Wei 2019b ¹⁶⁵	Sofosbuvir 400 mg +	Mortality: 0% (0/375)	Any adverse event: 50% (189/375)	Gilead
Multinational (China,	velpatasvir 100 mg		Serious adverse events (foot infection, pneumonia, ligament rupture): 1% (3/375)	
Malaysia, Singapore,			Withdrawal due to adverse events: 0% (0/375)	
I hailand, Vietnam)			Headache: 5% (18/375)	
Fair				
Zeuzem 2015 ¹⁰⁰	Grazoprevir 100 mg +	Patients without cirrhosis	Patients without cirrhosis only	Merck
C-EDGE	elbasvir 50 mg	only	Any adverse event: 71% (175/246)	
iviultinational (Australia,		wortality: 0.4% (1/246)	Serious adverse events (not described): 3% (7/246)	
Czech Republic, France,			vvitnarawai due to adverse event: 0.8% (2/246)	
Germany, Israel, Puerto				
Rico, South Korea,				
Taiwan, U.S.)				
Good				

Author year	Treatment Regimen			Funding
Quality	otherwise noted)	Clinical Outcomes	Adverse Events	Source
Zeuzem 2018 ¹⁶⁷	Glecaprevir 300 mg +	8-week intervention	8-week intervention group	AbbVie
ENDURANCE-1	pibrentasvir 120 mg	group	Any adverse event: 62% (216/351)	
Multinational (Australia,		Mortality: 0% (0/351)	Serious adverse events (suicide attempt, unstable angina, fracture, uterine	
Austria, Belgium,		12-week intervention	leiomyoma, transient ischemic attack): 1% (5/351)	
Canada, Chile, France,		group	Withdrawal due to adverse events: 0% (0/351)	
Germany, Hungary,		Mortality: 0.3% (1/352)	Headache: 19% (68/351)	
Israel, Italy, Lithuania,			Fatigue: 9% (31/351)	
Mexico, New Zealand,			Nausea: 5% (19/351)	
Poland, Portugal, Puerto			<u>12-week intervention group</u>	
Rico, Romania, Russian			Any adverse event: 66% (234/352)	
Federation, Spain, South			Serious adverse events (irritable bowel syndrome, pneumonia/death, bronchitis, atrial	
Korea, Sweden,			fibrillation): 1% (4/352)	
Switzerland, Taiwan,			Withdrawal due to adverse events: 0.3% (1/352)	
U.K., U.S.)			Headache: 18% (62/352)	
Fair			Fatigue: 12% (43/352)	
			Nausea: 8% (29/352)	
Zeuzem 2018 ¹⁶⁷	A. Glecaprevir 300 mg +	A vs. B vs. C	A vs. B vs. C	Same as
ENDURANCE-3 (same	pibrentasvir 120 mg, 8	Mortality: 0.6% (1/157)	Any adverse event: 62% (98/157) vs. 76% (177/233) vs. 70% (80/115)	Zeuzem 2018
publication as	weeks	vs. 0% (0/233) vs. 0.9%	Serious adverse events (ulcerative keratitis, overdose, substance-abuse	
ENDURANCE-1)	B. Glecaprevir 300 mg +	(1/115)	dependence): 2% (3/157) vs. 2% (5/233) vs. 2% (2/115)	
Fair	pibrentasvir 120 mg, 12		Withdrawal due to adverse events: 0% (0/157) vs. 1% (3/233) vs. 0.9% (1/115)	
	weeks		Headache: 20% (31/157) vs. 26% (60/233) vs. 20% (23/115)	
	3. Sofosbuvir 400 mg +		Fatigue: 13% (20/157) vs. 19% (44/233) vs. 14% (16/115)	
	daclatasvir 60 mg. 12		Nausea: 12% (19/157) vs. 14% (32/233) vs. 13% (15/115)	
	weeks			

*Serious adverse events occurring in more than one person (each occurred in 2 people; NR by intervention group): cellulitis, chest pain, gastroenteritis, hand fracture, noncardiac chest pain, pneumonia.

[†]One patient excluded from analysis due to receiving the wrong intervention.

Abbreviations: ALT = alanine aminotransferase; AST = aspartate amino transferase; CI = confidence interval; CLDQ-HCV = Chronic Liver Disease Questionnaire-Hepatitis C Version; EQ-5D VAS = EuroQoL 5-Dimensions questionnaire visual analog scale; FACIT-F = Functional Assessment of Chronic Illness Therapy-Fatigue; HCV-SIQv4 = Hepatitis C Symptom and Impact Questionnaire; NR = not reported; RR = relative risk; SD = standard deviation; SE = standard error; SF = short form; SVR = sustained virologic response; U.K. = United Kingdom; ULN = upper limit of normal; U.S. = United States; WPAI:SHP = Work Productivity and Activity Impairment Questionnaire: Specific Health Problem. Study names are not acronyms.

		Non-												
		randomized												
		studies:												
		Enrolled all												
		(or a												
		random					Primary					Loss to	Analyze	
	Single-	sample of)	Randomized	Randomized			outcome					followup	neonle in	
	or	nationts	studios	studios			nre-				Attrition	differential	the groups	
	multi	monting	Pandom-	Allocation	Groups	Eligibility	specified	Outcome	Caro		and	(>10%)/	in which	
	arm	inclusion	ization	concealment	similar at	critoria	and	assassors	nrovider	Patient	withdrawals	high	they were	
Author year	study2	critoria?	adequate?	adoquato?	basolino?	specified?	reported?	maskod?	maskod?	maskod?	reported?	(\\20%)2	assigned?	Quality
Abergel	Single	Unclear	NA	NA	NA	Yes	Yes	NA	No	No	Yes	No	Yes	Fair
20168142	<u>.</u>												.,	<u> </u>
Abergel 2016b ¹⁴¹	Single	Yes	NA	NA	NA	Yes	Yes	NA	No	No	Yes	No	Yes	Good
Afdhal 2014 ¹⁸⁵	Multi	NA	Unclear	No (open label)	Yes	Yes	Yes	NA	No	No	Yes	No	Yes	Fair
Ahmed 2018 ¹⁹⁵	Single	Yes	NA	NA	NA	Yes	Yes	NA	No	No	Yes	No	Yes	Fair
Andreone	Multi	NA	Unclear	No (open	Yes	Yes	Yes	NA	No	No	Yes	No	Yes	Fair
2014 ¹⁸⁶				label)										
Asselah 2018 ¹⁹⁶ SURVERYOR	Single	Unclear	NA	NA	NA	Yes	Yes	NA	No	No	Yes	No	Yes	Fair
II Asselah	Single	Unclear	NA	NA	NA	Yes	Yes	NA	No	No	Yes	No	Yes	Fair
2019 ¹⁴³ ENDURANCE-														
5 and 6														
Brown 2018 ¹⁴⁴ C-SCAPE	Single	NA	Unclear	No (open label)	No	Yes	Yes	NA	No	No	Yes	No	Yes	Fair
Chayama 2018 ¹⁹⁷	Single	Yes	NA	NA	NA	Yes	Yes	NA	No	No	Yes	No	Yes	Fair
CERTAIN-1														
Chuang 2016 ¹⁴⁵	Single	Unclear	NA	NA	NA	Yes	Yes	NA	No	No	Yes	No	Yes	Fair
Dore 2016 ¹³⁷	Multi	NA	Yes	No (open	Yes	Yes	Yes	Yes	No	No	Yes	Νο	Yes	Good
MALACHITE 1				label)										
Dore 2016 ¹³⁷	Multi	NA	Yes	No (open	Yes	Yes	Yes	Yes	No	No	Yes	No	Yes	Good
MALACHITE 2				label)										
Everson 2015 ¹⁴⁶	Multi	NA	Yes	No (open label)	Yes	Yes	Yes	NA	No	No	Yes	No	Yes	Good
Feld 2014 ¹⁸⁷	Multi	NA	Yes	Yes	Yes	Yes	Yes	NA	Yes	Yes	Yes	No	Yes	Good
Feld 2015 ¹³⁹	Multi	NA	Yes	Yes	Yes	Yes	Yes	NA	Yes	Yes	Yes	No	Yes	Good
		Non-												
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		randomized												
		Enrolled all												
		circled all												
		random					Brimany					Loss to	Analyzo	
	Single	cample of)	Pandomizod	Pandomizod			outcomo					followup	Analyze	
	or or	sample of	studios:	studios			bro-				Attrition	differential	the groups	
	multi-	meeting	Random-	Allocation	Groups	Fligibility	specified	Outcome	Care		and		in which	
	arm	inclusion	ization	concealment	similar at	criteria	and	assessors	nrovider	Patient	withdrawals	high	they were	
Author year	study?	criteria?	adequate?	adequate?	baseline?	specified?	reported?	masked?	masked?	masked?	reported?	(>20%)?	assigned?	Quality
Ferenci	Multi	NA	Yes	Yes	Yes	Yes	Yes	NA	Yes	Yes	Yes	No	Yes	Good
2014 ¹⁸⁸														
PEARL 3														
Ferenci	Multi	NA	Yes	Yes	Yes	Yes	Yes	NA	Yes	Yes	Yes	No	Yes	Good
PFARI 4														
Foster 2015 ¹⁴⁷	Multi	NA	Unclear	No (open	Yes	Yes	Yes	NA	No	No	Yes	No	Yes	Fair
ASTRAL 2				label)										
Foster 2015 ¹⁴⁷	Multi	NA	Unclear	No (open	Yes	Yes	Yes	NA	No	No	Yes	No	Yes	Fair
ASTRAL 3				label)										
Gane 2015 ¹⁴⁸	Single	Unclear	NA	NA	NA	Yes	Yes	NA	No	No	Yes	No	Yes	Fair
Grebely 2018 ¹⁵⁰	Single	Unclear	NA	NA	NA	Yes	Yes	NA	No	No	Yes	No	Yes	Fair
SIMPLIFY														
Grebely	Single	Unclear	NA	NA	NA	Yes	Yes	NA	No	No	Yes	No	Yes	Fair
2018 ¹⁴⁹														
D3FEAT	N /1 1+:	ΝΙΔ	Vaa	No (onon	Vaa	Vee	Vaa	NIA	No	No	Vaa	No	Vaa	Cood
2015 ¹⁸⁹	wuu	NA	res	label)	res	res	res	NA	INO	NO	res	INO	res	Good
Kowdley	Multi	NA	Unclear	No (open	Yes	Yes	Yes	NA	No	No	Yes	No	Yes	Fair
2014a ¹⁹⁰				label)										
Kowdley 2014b ¹⁹¹	Multi	NA	Yes	No (open label)	Yes	Yes	Yes	NA	No	No	Yes	No	Yes	Good
Kumada 2015 ¹⁵¹	Multi	NA	Unclear	Unclear	Yes	Yes	Yes	NA	Yes	Yes	Yes	No	Yes	Fair
Kumada	Multi	NA	Yes	Yes	Yes	Yes	Yes	NA	Unclear	Yes	Yes	No	Yes	Good
2017 ¹⁵²														
Kwo 2016 ¹⁵³	Multi	NA	Unclear	No (open label)	Yes	Yes	Yes	No	No	No	Yes	No	Yes	Fair
Lalezari	Single	Unclear	NA	NA	NA	Yes	Yes	NA	No	No	Yes	No	Yes	Fair
2015 ¹⁹²	3.5								-					
Lawitz 2014a ¹⁵⁴	Multi	NA	Yes	No (open label)	No	Yes	Yes	NA	No	No	Yes	No	Yes	Fair

	Single- or multi- arm	Non- randomized studies: Enrolled all (or a random sample of) patients meeting inclusion	Randomized studies: Random- ization	Randomized studies: Allocation concealment	Groups similar at	Eligibility	Primary outcome pre- specified and	Outcome	Care	Patient	Attrition and withdrawals	Loss to followup: differential (>10%)/ high	Analyze people in the groups in which they were	
Author year	study?	criteria?	adequate?	adequate?	baseline?	specified?	reported?	masked?	masked?	masked?	reported?	(>20%)?	assigned?	Quality
Lawitz 2014b ¹⁹³	Multi	NA	Yes	No (open label)	No	Yes	Yes	NA	No	No	Yes	No	Yes	Fair
Lawitz 2015 ¹⁵⁵	Single	Unclear	NA	NA	NA	Yes	Yes	NA	No	No	Yes	No	Yes	Fair
Lim 2016 ¹⁵⁶	Single	Unclear	NA	NA	NA	Yes	Yes	NA	No	No	Yes	No	Yes	Fair
Nelson 2015 ¹⁵⁷	Single	Unclear	NA	NA	NA	Yes	Yes	NA	No	No	Yes	No	Yes	Fair
Pianko 2015 ¹⁵⁸	Multi	NA	Yes	No (open label)	Yes	Yes	Yes	NA	No	No	Yes	No	Yes	Fair
Poordad 2017 ¹⁹⁴	Multi	NA	Unclear	No (open label)	No	Yes	Yes	NA	No	No	Yes	No	Yes	Fair
Pott-Junior 2019 ¹⁵⁹	Multi	NA	Yes	Yes	Yes	Yes	Yes	NA	No	No	Yes	No	Yes	Good
Sperl 2016 ¹⁹⁸ C-EDGE	Single	Yes	NA	NA	NA	Yes	Yes	No	No	No	Yes	No	Yes	Fair
Sulkowski 2014 ¹⁶¹	Multi	NA	Unclear	No (open label)	Yes	Yes	Yes	NA	No	No	Yes	No	Yes	Fair
Sulkowski 2015 ¹⁶⁰	Multi	NA	Unclear	No (open label)	Yes	Yes	Yes	NA	No	No	Yes	No	Yes	Fair
Toyoda 2018 ¹⁹⁹ CERTAIN-2	Single	Yes	NA	NA	NA	Yes	Yes	NA	No	No	Yes	No	Yes	Fair
Waked 2016 ¹⁶²	Sinale	Yes	NA	NA	NA	Yes	Yes	NA	No	No	Yes	No	Yes	Good
Wei 2018 ¹⁶³	Single	Unclear	NA	NA	NA	Yes	Yes	NA	No	No	Yes	No	Yes	Fair
Wei 2019a ¹⁶⁴ C-CORAL	Multi	NA	Yes	Yes	Yes	Yes	Yes	Yes	Yes	Yes	Yes	No	Yes	Good
Wei 2019b ¹⁶⁵	Single	Unclear	NA	NA	NA	Yes	Yes	NA	No	No	Yes	No	Yes	Fair
Zeuzem 2015 ¹⁶⁶	Multi	NA	Yes	Yes	Yes	Yes	Yes	NA	Yes	Yes	Yes	No	Yes	Good
Zeuzem 2018 ¹⁶⁷ ENDURANCE- 1	Multi	NA	Yes	No	Yes	Yes	Yes	NA	No	No	Yes	No	Yes	Fair

		Non-												
		randomized												
		studies:												
		Enrolled all												
		(or a												
		random					Primary					Loss to	Analyze	
	Single-	sample of)	Randomized	Randomized			outcome					followup:	people in	
	or	patients	studies:	studies:			pre-				Attrition	differential	the groups	
	multi-	meeting	Random-	Allocation	Groups	Eligibility	specified	Outcome	Care		and	(>10%)/	in which	
	arm	inclusion	ization	concealment	similar at	criteria	and	assessors	provider	Patient	withdrawals	high	they were	
Author year	study?	criteria?	adequate?	adequate?	baseline?	specified?	reported?	masked?	masked?	masked?	reported?	(>20%)?	assigned?	Quality
Zeuzem	Single	Yes	NA	NA	NA	Yes	Yes	NA	No	No	Yes	No	Yes	Fair
2018 ¹⁶⁷														
ENDURANCE-														
3														

Abbreviation: NA = not applicable. Study names are not acronyms.

Appendix B Table 14. Key Question 9: Association of Sustained Virologic Response and Clinical Outcomes – Study Characteristics

		Country	
		Dates of enrollment	
Author year		Number of centers	
Quality	Study type	(location)	Inclusion criteria
Arase 2007 ²⁰⁴	Cohort*	Japan	≥60 years of age: ALT elevation greater than double upper limits within
Fair	•••••	1989 to 2004	6 months: no corticosteroids or antiviral agents in last 6 months: no
' un		Single Center	HBV surface antigen, antinuclear antihodies, or antimitochondrial
		(Toranomon Hospital)	antibodies: leukocytes >3000/mm ³ platelet count >80 000/mm ³ and
		(Toranomorriospital)	bilirubin <2.0 mg/ml : IEN thoropy >4 works
			billubili <2.0 fig/file, iFN therapy >4 weeks
			Evaluded: History of cleaned abuse or advanced simbosis
			Excluded. Filsioly of alcohol abuse of advanced cirrinosis,
Apphing 2010217	Cabart	lanan	encephalopathy, bleeding esophageal valices, of ascites
Asanina 2010-11	Conort		HCV infection with histologically proven chronic nepatitis or cirmosis
Fair		1992 to 2008	
		Single center (Musashino	
D 1 004460	0 1 (†	Red Cross Hospital)	
Backus 201109	Conort+	U.S. (VA)	HCV genotype 1, 2, or 3; treated with pegylated interferon + ribavirin
Fair		2001 to 2008	
	A 1 1 1	Multicenter (national)	Exclusion: HIV infection, HCC prior to treatment
Butt 2017 ²⁰⁵	Cohort+	U.S. (VA)	HCV infected initiating paritaprevir + ritonavir + ombitasvir + dasabuvir
Fair		Enrollment dates NR	or ledipasvir + sofosbuvir
		Multicenter (national)	
Carrat 2019 ¹⁶⁸	Cohort	France	Patients with chronic HCV infection recruited from 32 hepatology
French National	(prospective)	2012 to 2015	centers in France.
Agency for		32 centers	Excluded: HBV, HIV coinfection, previous HCC diagnosis, history of
Research on AIDS			decompensated cirrhosis, liver transplant recipient
CO22 Hepather			
Cohort			
Fair			
Cozen 2013 ²⁰⁶	Cohort [‡]	U.S.	>18 years of age, HCV infection, underwent liver biopsy and follow-up
San Francisco VA		1992 to 2007	liver imaging study, biopsy, or clinic visit
Cohort		Two centers (San	
Fair		Francisco VA and	
		University of California at	
		San Francisco)	
Cozen 2013 ²⁰⁶	Cohort [‡]	U.S.	>18 years of age, HCV infection, underwent liver biopsy and follow-up
University of		1992 to 2007	liver imaging study, biopsy, or clinic visit
California at San		Two centers (San	
Francisco Cohort		Francisco VA and	
Fair		University of California at	
		San Francisco)	
Dieperink 2014 ²⁰⁷	Cohort [‡]	U.S. (VA)	Chronic HCV infection, initiated antiviral therapy
Fair		1997 to 2009	
		Single center	
		(Minneapolis VA)	
Dohmen 2013 ²¹⁸	Cohort	Japan	Chronic HCV infection with viral load ≥5 log IU/mL; HBV negative
Fair	(prospective)	2004 to 2010	Excluded: history of HCC or HCC developed in the first 6 months
		Multicenter (10 centers,	
		primarily in Fukuoka)	
El-Serag 2014 ²¹⁵	Cohort [‡]	U.S. (VA)	HCV infection, ≥1 year followup in VA
Fair		1999 to 2010	
		Multicenter (national)	
Ikeda 1999 ²¹⁹	Cohort*	Japan 1974-1995	Included: age 15 to 86
		Single center (Toronoman	Excluded: HBV, HCC, cirrhosis
Fair		Hospital)	
Imai 1998 ²²⁰	Cohort	Japan	Included: adults with HCV. Childs A cirrhosis
Fair		1992 to 1993	Excluded: HCC
		Multicenter (8 centers	
		primarily in Osaka	
		Japan)	

Appendix B Table 14. Key Question 9: Association of Sustained Virologic Response and Clinical Outcomes – Study Characteristics

		Country	
		Dates of enrollment	
Author year	Study type	Number of centers	Inclusion criteria
Imazeki 2003 ²⁰⁸	Cohort [§]	Japan	Chronic HCV infection underwent liver biopsy
Fair	Conort	1986 to 1998	
		Single center (Chiba University Hospital)	Excluded: HCC detected within six months of liver biopsy
Innes 2011 ²⁰⁹ Fair	Cohort	U.K. 1996 to 2007	HCV infection, treatment naive
		Multicenter (throughout Scotland)	Excluded: Nonsustained SVR (presence of viremia subsequent to meeting definition for SVR), liver transplant, HIV-positive, unknown treatment response
Ioannou 2018 ²²¹ <i>Fair</i>	Cohort	U.S. (VA) 1999 to 2015 Multicenter (national)	Initiation of antiviral regimen within VA from January 1999 to December 2015
Izumi 2005 ²²² Fair	Cohort [†]	Japan 1994 to 2001 Single center (Musashino Red Cross Hospital)	Chronic HCV infection, underwent interferon monotherapy
Kasahara 1998 ²²³ <i>Fair</i>	Cohort [¶]	Japan 1989 to 1995 10 centers (primarily in Osaka)	Included: adults with HCV Excluded: HCC, cirrhosis
Kasahara 2004 ²¹⁰ Fair	Cohort [¶]	Japan Enrollment dates NR Multicenter (number and location of centers unclear)	Histological diagnosis of chronic hepatitis or cirrhosis; no clinical complications of cirrhosis; no evidence of HCC on ultrasonography and/or computed tomography Excluded: HBV; HIV; co-existing liver diseases such as autoimmune hepatitis or primary biliary cirrhosis; excessive alcohol consumption (>80 g/day)
Kurokawa 2009 ²²⁴ <i>Fair</i>	Cohort [¶] (prospective)	Japan 2002 to 2005 Multicenter (number of centers unclear, primarily in Osaka)	All patients treated with interferon alfa-2a + ribavirin during study period Excluded: HBV, HIV positive; liver disease including history of HCC or HCC within 6 months after treatment cessation
Lee 2017 ²²⁵ Fair	Cohort	South Korea 2004 to 2013 Single center (Inha University Hospital)	HCV positive treated during study period Excluded: HBV positive; liver disease
Maruoka 2012 ²¹¹ <i>Fair</i>	Cohort [§]	Japan1986 to 2005Single center (Chiba University Hospital)	HCV positive, underwent liver biopsy Excluded: Other causes of chronic liver disease, HIV-positive, detection of HCC within 1 year of antiviral therapy, dropout within 1 year
Okanoue 2002 ²²⁶ Fair	Cohort	Japan 1995 to 1998 Multicenter (15 centers)	HCV infection, 18 to 68 years of age Excluded: HBV infection, HIV infection, daily alcohol intake >60 g of ethanol for more than 5 years, ALT <30 IU/L
Osaki 2012 ²²⁷ Fair	Cohort	Japan 2002 to 2010 Single center (Osaka Red Cross Hospital)	HCV infection, elevated liver enzymes, and ultrasound image demonstrating chronic liver damage Exclusion: neutrophil count <750 cells/uL, platelet count <50,000 cells/uL, hemoglobin level ≤9.0 g/dL, and renal insufficiency (serum creatinine levels >2 mg/dL), follow-up <24 weeks after the termination of the interferon therapy, previously treated for HCC, or occurrence of HCC during or within 24 weeks after treatment
Singal 2013 ²¹² Fair	Cohort	U.S. 2001 to 2006 Single center (Parkland Health and Hospital System)	HCV infection, life expectancy >5 years, platelet count >50,000/uL

Appendix B Table 14. Key Question 9: Association of Sustained Virologic Response and Clinical Outcomes – Study Characteristics

		Country	
Authorizon		Dates of enrollment	
Author year	Study type	(location)	Inclusion critoria
Sinn 2009231	Cohort	South Koroa	
Epir	Conon	1994 to 2004	
		Single center	
		(Sungkyunkwan	
		University School of	
		Medicine)	
Tanaka 2000 ²²⁸	Cohort	Japan	Chronic HCV infection with liver biopsy
Fair		1980 to 1996	
		Multicenter (6 hospitals in	Excluded: HBV infection, HCC or other liver disease such as alcoholic
		Osaka)	liver disease, autoimmune hepatitis, or primary biliary cirrhosis
Tateyama 2011 ²²⁹	Cohort	Japan,	Chronic HCV infection
Fair		1992 to 2003	
		Single center (National	
		Nagasaki Medical Center)	
Tseng 2016 ²¹⁶	Cohort	Taiwan	Age ≥65 years, chronic HCV infection, treated with pegylated
Fair		2005 to 2011	interferon; elevated ALT
		Single center (Dalin Izu	Excluded: Decompensated cirrhosis; malignant neoplasms;
		Chi General Hospital)	autoimmune diseases; HIV infection, neutropenia; thrombocytopenia;
Vachida 1000230	Cobort#	lance 1096 to 1009	anemia; poony controlled psychiatric diseases
Foshida 1999-00	Conon	Multicenter (8 centers	Excluded: HCC or other liver diseases (chronic HBV/ alcoholic liver
raii		throughout Japan	disease, autoimmune benetitis, or primary biliary cirrhosis)
		Inhibition of	disease, addiminitine nepatitis, or primary billary cirriosis
		Hepatocarcinogenesis by	
		Interferon Therapy Study	
		Group])	
Yoshida 2002 ²¹³	Cohort#	Japan	HCV positive, underwent liver biopsy
Fair		1986 to 1998	
		Multicenter (8 centers	Exclusion: HBV co-infection, alcoholic liver disease, autoimmune
		throughout Japan	hepatitis, or primary biliary cirrhosis
		[Inhibition of	
		Hepatocarcinogenesis by	
		Interferon Therapy Study	
Vu 2006 ²¹⁴	Cohort	Groupj)	Dianov proven abronia UCV infection, with an without simbosi-
1 u 2006 ² '7	Conort		Biopsy-proven chronic HUV intection, with or without cirrhosis
i all		Multicenter (1 centers in	Excluded: HBV or HIV, autoimmune benatitis, alcohol abuse (>80 g
		Taiwan)	ethanol per day) HCC at treatment initiation or within 6 months
		i aiwaii)	emanor per day, noo at treatment initiation of within 0 months

* Study populations overlap.

† Study populations overlap.‡ Study population appears to overlap with Ioannou 2018.

§ Study populations overlap.

Study population appears to overlap with Backus 2011, Butt 2017, Cozen 2013, Dieperink 2014, and El-Serag 2014.

¶ Study populations likely overlap.

Study populations appear to overlap.

Abbreviations: ALT = alanine aminotransferase; HBV = hepatitis B virus; HCC = hepatocellular carcinoma; HCV = hepatitis C virus; IFN = interferon; NR = not reported; SVR = sustained virologic response; U.K. = United Kingdom; U.S. = United States of America; VA = Veterans Affairs.

Author year	Treatment duration	Intervention (c)	Denulation	Variables accounted	Outcomes	Funding course
	Followup		Population	for in analyses		Funding source
Arase 2007 ²⁰⁴	I reatment duration:	SVR VS. NO SVR	Antiviral treatment: n=500	Liver fibrosis, sex, age,		Okinaka Memorial
Fair	(range 28 to 720)	SVR=Undetectable HCV RNA 6	SVR: n=140	HCV genotype, AST,	SVR: 0.19 (95% CI, 0.08 to 0.45)	Institute for Medical Research
	(Tallye 20 to 730)	torm IEN thorapy	Moon and (voars): 64	ALT, HCV VIIal IOau,	0.00 (0 0.45)	and Jananasa
	Followup: Moon 7.4	септисти спетару	Econolo: 50%	liver histology (activity)	NO SVR. Relefence	Ministry of Hoalth
	vears	IEN-2a or IEN-2b monotherapy:	Race: NR		Mortality aHR	Labor and Welfare
	years	94% IFN plus ribavirin	Genotype 1b: 60%		SVR: 0.39 (95% CI	
		combination therapy: 6%	Genotype 2: 34%		0.16 to 0.93	
		combination therapy. 070	Other genotype: 8.0%		No SVR: Reference	
			F1: 36%			
			F2: 31%		Liver-related mortality.	
			F3: 7.0%		aHR	
			F4: 14%		SVR: 0.13 (95% CI,	
					0.03 to 0.59)	
					No SVR: Reference	
Asahina 2010 ²¹⁷	Treatment:	SVR vs. no SVR	Antiviral treatment: n=2166	Age, sex, BMI, fibrosis	HCC, aHR, annual	Japanese Ministry
Fair [†]	24 or 48 weeks	SVR=Undetectable HCV RNA 6	SVR: n=686	stage, degree of	incidence	of Education,
	up to 2 to 5 years	months after completion of	No-SVR: n=1356	steatosis,	SVR: 0.38 (95% CI,	Culture, Sports,
		antiviral therapy	Prolonged therapy: n=59	esophagogastric	0.18 to 0.83), 0.4%	Science, and
	Followup:		Undetermined response: n=65	varices, genotype,	No SVR: Reference,	Technology
	Mean 7.5 years	IFN-alpha or beta monotherapy	Mean Age: 55.4 (SD±3.1)	albumin, ALI, ASI,	20.2%, 1.4%	
	(range 0.5 to 17	(n=1062)	Female: 50%	GGT, alkaline		Japanese Ministry
	years)			pnosphatase, total		of weifare, Health
		Combination therapy IFIN-alpha		Dillrubin, total		and Labor
			F1. 40%	facting blood sugar		
		Pegylated IEN-alpha	F2. 34%	white blood cell red		
		monotherany (n=386)	F4: 5%	blood cell platelet count		
			Genotype 1a: 0.3%	AFP (baseline and post		
		Combination pegylated IFN-	Genotype 1b: 70%	treatment) viral load		
		alpha and ribavirin (n=412)	Genotype 2a: 18%	IFN regimen		
			Genotype 2b: 10%			

Author year	Treatment duration			Variables accounted		
Quality	Followup	Intervention(s)	Population	for in analyses	Outcomes	Funding source
Backus 2011 ⁶⁹	Treatment duration:	SVR vs. no SVR	Antiviral treatment: n=16,864	Age, sex, albumin, AST,	All-cause mortality,	VA, Veterans
Fair [‡]	48 weeks for	SVR=Undetectable HCV RNA 6	SVR: n=7434	AST/ALT ratio,	aHR, 5-year mortality	Health
	genotype 1, 24	months after completion of	No SVR: n=9430	creatinine clearance,	rate	Administration,
	weeks for genotypes	antiviral therapy	Mean age (years): 52	platelets, sodium,	Genotype 1	Office of Public
	2 and 3	PEG-IFN (alfa-2a or alfa-2b) plus	Female: 4%	cirrhosis, chronic	SVR: 0.71 (0.60 to	Health and
	Followup: Median 3.8	ribavirin	Non-White: 43%	obstructive pulmonary	0.86), 6.7%	Environmental
	years (IQR 2.6 to 5.2)		Genotype 1: 72%	disease, diabetes,	No SVR: Reference,	Hazards
			Genotype 2: 17%	hypertension, tobacco	14%	
			Genotype 3: 11%	use, treatment duration	Genotype 2	
			Fibrosis stage: NR	<60% recommended,	SVR: 0.62 (0.44 to	
			Cirrhosis: 13%	bilirubin, BMI, HBV co-	0.87), 7.3%	
				infection, HCV viral load,	No SVR: Reference,	
				hemoglobin, coronary	16%	
				artery disease, cancer,	Genotype 3	
				congestive heart failure,	SVR: 0.51 (0.35 to	
				cerebrovascular	0.75), 8.0%	
				disease, schizophrenia,	No SVR: Reference,	
				recent alcohol abuse	24%	
				diagnosis, anxiety	SVR vs. no SVR	
				disorder, depression,	(calculated): 0.66 (0.57	
				hard drug use, post-	to 0.76)	
				traumatic stress		
				disorder, socioeconomic		
				status instability,		
				multiple treatment		
				courses, erythropolesis		
				stimulating agent use,		
				granulocyte colony		
				stimulating factor use,		
				year of treatment start		

Author year	Treatment duration			Variables accounted		
Quality	Followup	Intervention(s)	Population	for in analyses	Outcomes	Funding source
Butt 2017 ²⁰⁵	Treatment duration:	SVR vs. no SVR	Antiviral treatment: n=6,970	Age, sex, race/ethnicity,	Mortality, aHR	VA, Pittsburgh
Fair‡	NR	SVR not defined	SVR: n=6,371	BMI, FIB-4 score >3.5;	SVR: 0.57 (95% CI,	
			No SVR: n=599	diabetes, chronic kidney	0.33 to 0.99)	
	Followup: 1.5 years	Paritaprevir + ritonavir +		disease stage 3-5;	No SVR: Reference	
		ombitasvir + dasabuvir (n=1,473)	Paritaprevir + ritonavir +	alcohol		
		Ledipasvir + sofosbuvir	ombitasvir + dasabuvir vs.	use/dependence; drug		
		(n=5,497)	ledipasvir + sofosbuvir	abuse/dependence;		
			Median age (years): 61 to 62	HCV RNA, genotype,		
			Female: 3% vs. 4%	anemia		
			White: 47% vs. 55%			
			Black: 32% vs. 26%			
			Hispanic: 2% vs. 2%			
			Genotype 1a: 61% vs. 64%			
			Genotype 1b: 38% vs. 17%			
			Child-Turcotte-Pugh class A:			
			94% vs. 90%			
			Class B: 6% vs. 10%			
			Class C: 0.1% vs. 0.5%			
			FIB-4 score >3.5 (cirrhosis):			
			13% vs. 15%			

Author year	Treatment duration			Variables accounted		
Quality	Followup	Intervention(s)	Population	for in analyses	Outcomes	Funding source
Carrat 2019 ¹⁶⁸ French National Agency for Research on AIDS CO22 Hepather Cohort <i>Fair</i>	Treatment duration: NR Followup: Median 33.4 months (IQR: 24.0 to 40.7)	SVR vs. no SVR SVR not defined DAA regimen (sofosbuvir + simeprevir +/- ribavirin; sofosbuvir + daclatasvir +/- ribavirin; sofosbuvir + ledipasvir +/- ribavirin; sofosbuvir + velpatasvir +/- voxilaprevir; paritaprevir + ritonavir + ombitasvir +/- dasabuvir +/- ribavirin; elbasvir + grazoprevir +/- ribavirin) (n=4,521, non- cirrhosis only)	Antiviral treatment: 4,521 SVR: n=3,286 No SVR: n=146 Unknown SVR: n=1,089 No treatment: 2,329 Total study population (including 3,045 patients with cirrhosis) Treatment vs. no treatment Mean age: 57 vs. 54 Female: 44% vs. 54% Race NR Fibrosis stage: F0, F1, or F2: 41% vs. 84% F3: 17% vs. 6% F4: 42% vs. 10% Genotype 1: 67% vs. 64% Genotype 2: 6% vs. 10% Genotype 3: 13% vs. 9% Genotype 4: 13% vs. 14% Genotypes 5 to 7: 2% vs. 3%	Age, sex, BMI, geographical origin, infection route, fibrosis score, treatment history, genotype, alcohol consumption, diabetes, arterial hypertension, biological variables, time-dependent covariates of treatment response	All-cause mortality, <u>aHR, rate SVR</u> : 0.64 (95% CI, 0.33 to 1.23), 21/4,422 person-years No SVR: 0.47 (95% CI, 0.06 to 4.04), 1/239 person-years No treatment: Reference, 48/11,131 person-years <u>HCC, aHR, rate SVR</u> : 0.75 (95% CI, 0.23 to 2.40), 9/4,400 person- years No SVR: 3.46 (95% CI, 0.61 to 19.7), 3/234 person-years No treatment: Reference, 14/11,120 person-years No SVR: NR, 0/239 person-years No SVR: NR, 0/239 person-years No treatment: Reference, 6/11,131 person-years No SVR: NR, 0/236 person-years No SVR: NR, 0/236 person-years No SVR: NR, 0/236 person-years No treatment: Reference, 4/11,131 person-years No treatment: Reference, 4/11,131 person-years	French National Agency for Aids and Viral Hepatitis Research; French National Agency of Research; French Ministry of Social Affairs and Health; Merck Sharp & Dohme; Janssen; AbbVie; Bristol- Myers Squibb; Roche

Author year	Treatment duration			Variables accounted		
Quality	Followup	Intervention(s)		tor in analyses	Outcomes	Funding source
Cozen 2013 ²⁰⁰	I reatment duration:	SVR vs. nonresponder vs.	San Francisco VA Conort	FIDROSIS Stage, age,	Cirrnosis, aHR, rate	National Institutes
San Francisco VA		RVB-Undetectable HCV/ RNA 6		race/ethnicity, HCV	SVR: 0.68 (95 % CI	or Health, VA merit
Conort	(SD ZZ.SZ)	SVR=Undelectable HCV RINA o	Neprospondor: p. 40	genolype, alconol use,	(7/60)	awaru
rall†	rollowup. Mean To	antiviral therapy	Rolopaor: n=22	subsidiice use,	(7/09) Noprospondor: 2.25	
	years	Relansor-Undetectable viral	Forly treatment	psychiatric	(05%) CL 1 18 to 1 60)	
		load during treatment with	discontinuation/unknown: n=10	stability	(95% CI, 1.10104.09),	
		detectable virus at 6 month	Mean Age 50.98 (SD 6.68)	Stability	Relanser: 1 00 (95%	
		followup	Female: 1 1%		CL 0 28 to 3 56) 22%	
		IFN alpha +/- ribavirin	African-American: 20.2%		(4/22)	
			Latino: 8.7%		Never treated:	
			Asian: 5%		Reference 14%	
			Genotype 1: 68.7%		(28/199)	
			Genotype 2: 14.5% Genotype		SVR vs. no SVR	
			3: 8.4% Genotype 4: 1.7%		(calculated): 0.35 (95%	
			Mixed genotype: 0.6%		CI, 0.11 to 1.10)	
			F0: 31%			
			F1: 24%		Mortality, aHR, rate	
			F2: 26%		SVR: 0.23 (95% CI,	
			F3: 8.4%		0.07 to 0.75), 8.7%	
			F4: 1.7%		(6/69)	
					Nonresponder: 0.56	
					(95% CI, 0.24, to 1.32),	
					29% (14/49)	
					Relapser 0.11 (95% CI,	
					(4/22)	
					Never treated:	
					Reference 24%	
					(47/199)	
					SVR vs. no SVR	
					(calculated): 0.50 (95%	
					CI, 0.12 to 2.10)	
					, ,	

Author year	Treatment duration		_	Variables accounted		
Quality	Followup	Intervention(s)	Population	for in analyses	Outcomes	Funding source
Cozen 2013 ²⁰⁶ University of California, San Francisco Cohort <i>Fair</i> [‡]	Treatment duration: mean 40.45 weeks (SD 22.32) Followup: Mean 10 years	SVR vs. nonresponder vs. relapser SVR=Undetectable HCV RNA 6 months after completion of antiviral therapy Relapser=Undetectable viral load during treatment with detectable virus at 6 month followup IFN alpha +/- ribavirin	University of California, San Francisco Cohort Antiviral treatment: n=131 SVR: n=43 Nonresponder: n=42 Relapser: n=21 Early treatment discontinuation/unknown: n=25 Mean age: 48.42 (SD 8.39) Female: 38.9% African-American: 9.9% Latino: 4.6% Asian: 13.0% Genotype 1: 63.3% Genotype 2: 18.3% Genotype 3: 12.2% Genotype 4: 0% Genotype 6: 1.5% F0: 11.5% F1: 23.7% F2: 30.5% F3: 19.1% F4: 15.3%	Fibrosis stage, age, race/ethnicity, HCV genotype, alcohol use, substance use, psychiatric comorbidities, social stability	Cirrhosis, aHR, rate SVR: 1.12 (0.12 to 10.33), $5.1%$ (2/43) Nonresponder: 5.90 (1.50 to 23.24), 36% ($11/42$) Relapser: 0.23 (0.02 to 2.27), $5.3%$ ($1/21$) Never treated: Reference, 7.8% ($10/134$) SVR vs. no SVR (calculated): 0.43 (95% CI, 0.03 to 5.35) Death or liver transplant University of California, San Francisco cohort, aHR, rate SVR: 0.24 (0.05 to 1.10), $7.0%$ ($3/43$) Nonresponder: 0.43 (0.13 to 1.38), 26% ($11/42$) Relapser: 0.80 (0.21 to 3.04), $19%$ ($4/21$) Never treated: Reference, 11% ($15/134$)	National Institutes of Health, VA merit award

Author year	Treatment duration			Variables accounted		
Quality	Followup	Intervention(s)	Population	for in analyses	Outcomes	Funding source
Dieperink 2014 ²⁰⁷ Fair [‡]	Followup: Median 7.5 years (IQR 4.9 to 9.8)	SVR vs. no SVR SVR not defined PEG-IFN-alpha plus ribavirin (68%) IFN-alpha plus ribavirin (26%) IFN-alpha (3.0%) Consensus IFN and ribavirin (3.0%)	Antiviral Treatment: n=536 SVR: n=222 Non-SVR: n=314 Median age (years): 52 (range 36 to 72) Female: 2% Black: 10% White: 81% Hispanic: 0.4% Asian: 0.4% Native American: 1.5% Unknown/other race: 7.3% Genotype 1: 70% Genotype 2: 15% Genotype 3: 12% Genotype 4: 0.2 Unknown genotype: 2.6% Clinical cirrhosis: 7.1% F0: 2.6% F1: 12% F2: 22% F3: 22% F4: 21% No biopsy: 21%	SVR, integrated care, genotype, fibrosis stage, diabetes, thrombocytopenia, age, depression Not significant in univariate analyses (excluded from model): alcohol use diagnoses, substance use diagnoses, psychosis, number of antiviral treatments, cardiac disease	SVR vs. no SVR All-cause mortality, aHR, rate SVR: 0.47 (95% CI, 0.26 to 0.85), 9% (19/222) No SVR: Reference, 26% (81/314) Liver related mortality, rate SVR: 3% (6/222) No SVR: 18% (56/314) Liver transplant, rate SVR: <1% (2/222) No SVR: 4% (13/314) HCC, aHR, rate SVR: 0.41 (95% CI, 0.18 to 0.96), 4% (9/222) No SVR: Reference, 9% (29/314)	Supported by VA Research Service
Dohmen 2013 ²¹⁸ Fair	Treatment duration: Range 24-72 weeks Followup: median 4.75 years (range 1 to 6.25 years)	SVR vs. no SVR SVR=Undetectable HCV RNA by PCR at 24 weeks after completion of antiviral therapy Oral ribavirin plus subcutaneous PEG-IFN-α-2a or subcutaneous PEG-IFN-α-2b	Antiviral treatment: n=474 SVR: n=285 No SVR: n=189 Mean age: 55 years Female: 52% Race: NR Genotype 1: 67% Genotype 2: 33% Fibrosis stage: NR	Age, sex, genotype, hemoglobin, platelet count, albumin, ALT, viral load, alpha- fetoprotein level	HCC, aHR, rate SVR: 0.39 (calculated 95% CI, 0.24 to 0.64, p=0.0002), 2% (6/285) No SVR: Reference, 9% (17/189)	NR

Author year	Treatment duration			Variables accounted		
Quality	Followup	Intervention(s)	Population	for in analyses	Outcomes	Funding source
El-Serag 2014 ²¹⁵ Fair [‡]	Treatment duration: NR Followup: Mean: 5.2 years	SVR vs. no SVR vs. undeterminable vs. no treatment SVR=Undetectable HCV RNA 12 weeks after completion of antiviral therapy Treatment NR	Demographics reported for all patients Antiviral treatment: n=16344 SVR: n=7577 No SVR: n=8767 Undeterminable: n=7188 No treatment: n=125875 Age: 52.5% Female: 2.9% White: 56% African American: 36% Hispanic: 6.0% Genotype 1: 55% Genotype 2: 8% Genotype 3: 5% Genotype 3: 5% Genotype 4: 1% Genotype 5/6: <1% Unknown genotype: 31% Fibrosis stage: NR	Age, sex, service period, HCV diagnosis year, genotype, diabetes, alcohol abuse, BMI, HIV coinfection, HBV coinfection	Cirrhosis, aHR SVR: 0.75 (95% CI, 0.69 to 0.82) No SVR: 2.07 (95% CI, 1.97 to 2.18) Undeterminable: 1.55 (95% CI, 1.45 to 1.66) No treatment: Reference SVR vs. no SVR (calculated): 0.36 (95% CI, 0.33 to 0.40) HCC, aHR SVR: 0.40 (95% CI, 0.32 to 0.50) No SVR: 1.34 (95% CI, 1.19 to 1.50) Undeterminable: 0.96 (95% CI, 0.82 to 1.12) No treatment: Reference SVR vs. no SVR (calculated): 0.30 (0.23 to 0.38)	National Institutes of Health grant - National Cancer Institute R01 116845 Houston VA Health Services Research & Development Center for Innovations in Quality, Effectiveness and Safety Texas Digestive Disease Center National Institutes of Health DK58338
Ikeda 1999 ²¹⁹ <i>Fair*</i>	Treatment duration:14 to 24 weeks Followup: Median 5.4 years (range 0.1 to 22.8)	Responder vs. nonresponder Complete response=Persistent undetectable HCV RNA 6 months after completion of antiviral therapy Incomplete responder=normal ALT values without elimination of HCV RNA for ≥6 months after treatment IFN alpha, beta or both	Antiviral treatment: n=1191 Responders: n=606 (461 complete responders and 145 incomplete [biochemical] responders) Nonresponders: n=585 No treatment: n=452 Median age (years): 50 (range15-86) Female 33% (389/1191) Race: NR Genotype 1a, 1b: 67% Genotype 2a, 2b: 28% Unknown genotype: 5% F1: 67% F2 and F3: 33% F4: 0%	Age, sex, alcohol intake, family history of HCC, history of blood transfusion, fibrosis stage, AST, ALT, albumin, bilirubin, globulin, gamma- glutamyl transferase, platelet count, indocyanine green retention rate at 15 minutes, HCV genotype, HCV viral load	HCC, aHR, rate Responder: 0.32 (95% Cl, 0.13 to 0.78), 1.2% (7/606) Nonresponder: 0.96 (95% Cl, 0.55 to 1.70), 3.6% (21/585) No treatment: Reference, rate NR SVR vs. no SVR (calculated): 0.33 (95% Cl, 0.12 to 0.96)	NR

Author year	Treatment duration			Variables accounted		
Quality	Followup	Intervention(s)	Population	for in analyses	Outcomes	Funding source
Imai 1998 ²²⁰	Treatment duration:	SVR vs. relapse vs.	Antiviral treatment: n=419	Age, sex, ALT, AFP,	HCC, aHR, rate	NR
Fair	24 weeks	nonresponder	SVR: n=151	platelet count, fibrosis	SVR: 0.06 (95% CI,	
	Follow-up: 47.6	SVR=Persistent normalization of	Relapse: n=120	stage, Histologic Activity	0.01 to 0.46), 0.7%	
	months (range 3.3 to	ALT levels during treatment and	Nonresponder: n=148	Index	(1/151)	
	65.2 months)	followup	No treatment (historical		Relapse: 0.51 (95% CI,	
		Relapse=Normal ALT at end of	control): 144		0.20 to 1.27), 6.1%,	
		treatment, but abnormally	Age <60: 71%		5.8% (7/120)	
		elevated levels after treatment	Female 33%		Nonresponder: 0.95	
			Race: NR		(95% CI, 0.48 to 1.84),	
		Human lymphoblastoid IFN,	Genotype: NR		13% (20/148)	
		recombinant IFN alpha 2a,	F1: 30%		No treatment:	
		recombinant IFN alpha 2b	F2: 33%		Reference, 13%	
			F3: 29%		(19/144)	
			F4: 8%		SVR vs. no SVR	
					(calculated): 0.06 (95%	
					Cl, 0.01 to 0.48)	

Author vear	Treatment duration			Variables accounted		
Quality	Followup	Intervention(s)	Population	for in analyses	Outcomes	Funding source
Imazeki 2003 ²⁰⁸ Fair [§]	Treatment duration: Mean 167 (range 6 to 560) days Followup: Mean 8.2 years (range 7 to 183 months)	SVR vs. no SVR SVR=Undetectable HCV RNA 6 months after completion of antiviral therapy IFN-2a: 84% IFN-2b: 12% Both: 4%	Antiviral treatment: n=355 SVR: n=116 No SVR: 239 Mean age (years): 49 Female: 36% Race: NR Genotype 1: 74% F0/F1: 56% F2: 17% F3: 14% F4: 13%	Age, sex, fibrosis stage, AST, ALT, albumin, platelet count, alcohol consumption, duration of HCV infection	Liver-related mortality, aHR, rate SVR: 0.030 (95% CI, 0.003 to 0.27), 0.9% (1/116) No SVR: 0.26 (95% CI, 0.11 to 0.61), 7.5% (18/239) No treatment: Reference, 12% (12/104) SVR vs. no SVR (calculated): 0.12 (95% CI, 0.01 to 1.28) All-cause mortality, aHR, rate SVR: 0.22 (95% CI, 0.068 to 0.71), 3.4% (4/116) No SVR: 0.63 (95% CI, 0.32 to 1.26), 12% (29/239) No treatment: Reference, 14% (15/104) SVR vs. no SVR (calculated): 0.35 (95% CI, 0.09 to 1.36)	NR
Innes 2011 ²⁰⁹ Fair	Treatment duration: Not specified Followup: Mean 5.3 years (range 27 days to 12.4 years)	SVR vs. no SVR SVR=Undetectable HCV RNA >6 months after completion of antiviral therapy PEG-IFN plus ribavirin: 61% PEG-IFN monotherapy: 1% IFN plus ribavirin: 21% IFN monotherapy: 18%	Antiviral treatment: n=1215 SVR: n=560 No SVR: n=655 Mean age (years): 42 Female: 31% Non-White: 7.8% Genotype 1: 36% Non-genotype 1: 55% Unknown genotype: 9.2% Fibrosis stage: NR Cirrhosis: 14%	Sex, age, race, IVDU, genotype, cirrhosis, alcohol-related hospitalization, elevated ALT	Liver-related mortality, aHR, rate SVR: 0.22 (95% CI, 0.09 to 0.58), 0.9% (5/560) No SVR: Reference, 7.6% (50/655) Liver-related hospital episode, aHR SVR: 0.22 (95% CI, 0.15 to 0.34) No SVR: Reference	Scottish Government

Author year	Treatment duration			Variables accounted		
Quality	Followup	Intervention(s)	Population	for in analyses	Outcomes	Funding source
Ioannou 2018 ²²¹	Treatment duration:	SVR vs. no SVR	Antiviral treatment=50,886	Cirrhosis,	HCC, aHR, rate	National Institutes
Fair	NR	SVR=HCV RNA <lower limit="" of<="" th=""><th>(excluding persons with</th><th>decompensated</th><th>All regimens (excludes</th><th>of Health/National</th></lower>	(excluding persons with	decompensated	All regimens (excludes	of Health/National
		detection 12 weeks after	cirrhosis)	cirrhosis, age, sex,	cirrhotics)	Cancer Institute
	Followup duration:	completion of antiviral therapy	SVR: 28,655	race/ethnicity, BMI, HCV	SVR: 0.32 (95% CI,	grant
	mean 6.1 years		No SVR: 23,231	genotype, HCV viral	0.28 to 0.37), 1.1%	R01CA196692
		IFN or pegylated IFN: 58%		load, HIV co-infection,	(316/28,655)	
		DAA + IFN: 7.3%	All patients (included persons	HBV co-infection, type 2	No SVR: Reference,	VA Clinical
		DAA only: 35%	with cirrhosis)	diabetes mellitus,	7.7% (1,778/23,231)	Science Research
			Mean age: 55.8 (SD ±7.6)	alcohol use disorders,	All regimens (includes	& Development
			years	substance abuse	<u>cirrhotics)</u>	grant
			Female: 3.4%	disorders, liver	SVR: 0.39 (95% CI,	101CX001156
			White: 55.6%	transplantation, platelet	0.35 to 0.43 , $1.9%$	
			Black: 26.3%	count, AST/ALT ratio,	(642/34,660)	
			Alspanic: 6.0%	International normalized	NO SVR: Reference,	
			Other: 1.6%	ratio, nemoglobin	9.5% (2629/27,694)	
			Construct 1: 77%		cirrbotice)	
			Constyne 2: 14%			
			Genotype 2: 14%		0.32 (95% C),	
			Genotype 3: 0.3%		(303/11 988)	
			Fibrosis stage: NR		No SVR: Reference	
			Cirrhosis: 16.8%		9.8% (2348/23.883)	
			(decompensated 4 7%)		DAA + IFN (includes	
					cirrhotics)	
					SVR: aHR 0.48 (95%	
					Cl. 0.32 to 0.73), 2.1%	
					(59/2763)	
					No SVR: 6.5%	
					(116/1772)	
					DAA only (includes	
					cirrhotics)	
					SVR: HR 0.29 (95% CI,	
					0.23 to 0.37), 1.4%	
					(280/19,909)	
					No SVR: Reference,	
					8.1% (165/2039)	

Author year Quality	Treatment duration	Intervention(s)	Population	Variables accounted for in analyses	Outcomes	Funding source
Izumi 2005 ²²² Fair [†]	Treatment duration: 24 weeks Followup: Duration NR	SVR vs. no SVR SVR=Undetectable HCV RNA 6 months after completion of antiviral therapy IFN monotherapy	Antiviral therapy: n=495 SVR: n=155 No SVR: n=340 Mean age (years): 52 Female: 43% Race: NR Genotype 1b: 50% Genotype 2a: 13% Genotype 2b: 7.9% F1: 27% F2: 37% F3: 25% F4: 0.7%	Age, sex, and fibrosis stage reported as statistically significant predictors of outcomes in multivariate model, otherwise unclear	HCC, aHR, rate SVR: 0.36 (95% CI, 0.04 to 0.83), 1.9% (3/155) No SVR: Reference, 8.2% (28/340)	Japanese Ministry of Health Labor and Welfare
Kasahara 1998 ²²³ <i>Fair</i> [¶]	Treatment duration: 14 to 52 weeks Follow up, mean: 37.4 months (range 13 to 97 months)	SVR vs. relapse vs. nonresponder SVR=Normalized ALT levels 24 weeks after completion of antiviral therapy Relapse=normalized ALT during therapy, abnormal ALT levels 24 weeks after therapy IFN alpha 2a, IFN alpha 2b, IFN beta, natural IFN alpha	Antiviral treatment: n=1022 SVR: n=313 Relapse: n=304 Non-responder: n=405 Mean age (years): 53 Female: 33% Race: NR Genotype 1: 58% Genotype 2: 18% Mixed or unclassified: 1.5% Genotype not tested: 23% METAVIR stage (mean): 1.9 to 2.3 Cirrhosis: Excluded	Age, gender, total histological score, Knodell's scores (periportal necrosis, intralobular or portal inflammation, and fibrosis), HCV genotype, HCV viral load, IFN dose, number of courses of IFN treatment, period of observation, ALT response	HCC, aHR, rate SVR: 0.13 (95% CI, 0.03 to 0.57), 1.6% (5/313) Non-responder: Reference, 7.9% (32/405) HCC, aHR, rate SVR: 0.32 (95% CI, 0.06 to 1.69), 1.6% (5/313) Relapse: Reference, 3.0% (9/304) HCC SVR vs. no SVR (calculated): 0.19 (95% CI, 0.06 to 0.58)	NR

Author vear	Treatment duration			Variables accounted		
Quality	Followup	Intervention(s)	Population	for in analyses	Outcomes	Funding source
Kasahara 2004 ²¹⁰ Fair [¶]	Treatment duration: 4 to12 months Followup: Mean 5.7 (SD± 2.0) years vs. 5.8 (SD±1.9)	SVR vs. No SVR SVR=Normalized ALT levels 24 weeks after completion of antiviral therapy IFN	Antiviral Treatment: n=2698 SVR: n=738 No SVR: n=1930 No treatment: n=256 Median age (years): 53 (range 20 to 76) Female: 36% Race: NR Genotype: NR F0: 0.7% vs. 0.6% F1: 35% vs. 25% F2: 36% vs. 32% F3: 26% vs. 38% F4: 3% vs. 5%	Age, gender, fibrosis stage, liver biopsy date	All-cause mortality, aHR, rate SVR: 0.14 (95% CI, 0.06 to 0.35), 0.9% (7/738) No SVR: 0.59 (95% CI, 0.33 to 1.06), 4.9% (94/1930) No treatment: Reference, 20% (52/256) SVR vs. no SVR (calculated): 0.24 (95% CI, 0.08 to 0.68) Liver-related mortality SVR: 0.04 (95% CI, 0.005 to 0.30), 0.1% (1/738) No SVR: 0.76 (95% CI, 0.40 to 1.42), 3.5% (68/1930) No treatment: Reference, 16% (42/256) SVR vs. no SVR (calculated): 0.05 (95% CI, 0.01 to 0.45)	NR
Kurokawa 2009 ²²⁴ <i>Fair</i> [¶]	Treatment duration: NR Followup: median 3 years (range 6 months to 5 years)	SVR vs. no SVR SVR=Undetectable HCV-RNA 24 weeks after completion of antiviral therapy Subcutaneous IFN-α-2b + oral ribavirin	Antiviral treatment: n=403 SVR: n=139 No SVR: n=264 Mean age (years): 55.8 (SD 10.9) Female: 36% Race: NR Genotype 1: 89% F0: 4% F1: 37% F2: 14% F3: 23% F4: 2%	Sex, age, fibrosis	HCC, aHR, rate SVR: 0.28 (95% CI, 0.08 to 0.96), 2.9% (4/139) No SVR: Reference, 8.0% (21/264)	NR

Author year	Treatment duration			Variables accounted		
Quality	Followup	Intervention(s)	Population	for in analyses	Outcomes	Funding source
Lee 2017 ²²⁵ <i>Fair</i>	Treatment duration: NR Followup: Median 2.6 years (range 6 months to 12 years)	SVR vs. no SVR SVR=Undetectable HCV RNA 24 weeks after completion of antiviral therapy PEG-IFN + ribavirin: 93% IFN followed by PEG-IFN + ribavirin: 7%	Antiviral Treatment: n=489 SVR: n=306 No SVR: n=183 Median age (years): 46 Female: 36% Race: NR Genotype 1: 51% Genotype 2: 40% Mixed genotype 1 and 2: 0.2% Mixed genotype 3 or 4: 0.2% Fibrosis stage: NR Cirrhosis: 13%	Age, sex, BMI, cirrhosis, ALT, HCV RNA, HCV genotype	HCC, aHR, rate SVR: 0.09 (95% CI, 0.02 to 0.40), 1.1% (n/N unclear) No SVR: Reference, 9.8% (18/183)	Inha University Hospital
Maruoka 2012 ²¹¹ <i>Fair</i> §	Treatment duration: Median 25 (range 1- 267) weeks Followup: Mean 9.9±5.3 years	SVR vs. no SVR SVR=Undetectable HCV RNA >6 months after completion of antiviral therapy IFN-alfa or -beta monotherapy: 83% IFN-alfa or -beta sequential therapy: 3.3% IFN-alfa plus ribavirin combination therapy: 14%	Antiviral treatment: n=577 SVR: n=221 No SVR: n=356 No treatment: n=144 Mean age (years): 50 Female: 36% Race: NR Genotype 1: 73% Genotype 2: 27% F0 or F1: 53% F2: 23% F3: 14% F4: 10%	Sex, age, fibrosis stage, inflammatory grade, genotype, high viral load, genotype 1 and high viral load, ALT, platelets, albumin	All-cause mortality, aHR, rate SVR: 0.17 (95% CI, 0.075 to 0.40), 4.5% (10/221) No SVR: 0.84 (95% CI, 0.50 to 1.42), 21% (74/356) No treatment: Reference, 26% (37/144) SVR vs. no SVR (calculated): 0.20 (0.08 to 0.54) HCC, aHR, rate SVR: 0.14 (95% CI, 0.046 to 0.42), 2.3% (5/221) No SVR: 1.18 (95% CI, 0.69 to 2.01), 22% (80/356) No treatment: Reference, 24% (35/144) SVR vs. no SVR (calculated): 0.12 (95% CI, 0.03 to 0.41)	NR

Author year	Treatment duration			Variables accounted		
Quality	Followup	Intervention(s)	Population	for in analyses	Outcomes	Funding source
Okanoue 2002 ²²⁶ Fair	Treatment duration: 16 to 26 weeks Followup: Mean 5.6 years	SVR vs. relapse vs. nonresponder SVR=Normalized ALT levels 6 months after completion of antiviral therapy Relapse=Normalized ALT during treatment, elevated levels 6 months after treatment Natural IFN Recombinant IFN2a Recombinant IFN2b Natural IFNB	Antiviral Treatment: n=1,370 SVR: n=426 Relapse: n=358 Nonresponder: n=586 Mean age 50.4 (SD±11.5) Female: 37% Race: NR Genotype: NR F1: 17% F2: 52% F3: 28% F4: 4%	Sex, age, fibrosis stage, serum ALT level, platelet count	HCC, aHR, rate SVR: 0.10 (95% CI, 0.04 to 0.28), 0.2% (1/426) Relapse: 0.55 (95% CI, 0.34 to 0.89), 2% (8/358) Non-responder: Reference, 7.5% (44/586) SVR vs. no SVR (calculated): 0.13 (95% CI, 0.06 to 0.27) All-cause mortality, rate SVR: 1% (2/426) Relapse: 3% (10/358) Non-responder: 6% (37/637)	Ministry of Education of Japan and Health and Welfare of Japan
Osaki 2012 ²²⁷ Fair	Treatment: 48 to 72 weeks for HCV genotype 1 and serum HCV RNA >5 log IU/mL, 24 weeks otherwise Followup: Median 4.1 (range 0.1 to 8.4) years	SVR vs. no SVR SVR=Undetectable HCV RNA 6 months after completion of antiviral therapy IFN + ribavirin (n=69) Or PEG-IFN + ribavirin (n=313)	Antiviral Treatment: n=382 SVR: n=185 No SVR: n=197 Median age (years): 59 (range 18-81) Female: 50% Race: NR Genotype 1b: 60% (genotype otherwise NR) Fibrosis stage: NR Cirrhosis: Excluded	Age, sex, HCV genotype, virological response, biochemical response, ALT, AFT, platelet count	HCC, aHR, rate SVR: 0.12 (95% CI, 0.01 to 0.94), 1% (1/185) No SVR: Reference, 11% (22/197)	Ministry of Education, Culture, Sports, Science and Technology, and the Ministry of Health, Labor and Welfare of Japan

Author year	Treatment duration			Variables accounted		
Quality	Followup	Intervention(s)	Population	for in analyses	Outcomes	Funding source
Singal 2013 ²¹² Fair	Treatment Duration: 48 weeks for genotypes 1,4, 6 and 24 weeks for genotypes 2 and 3 Followup: Median 72 months in SVR patients, 36-65 months in nonresponders	SVR vs. no SVR SVR=Undetectable HCV RNA 24 weeks after completion of antiviral therapy PEG-IFN α-2b and ribavirin	Antiviral treatment: n=242 SVR: n=83 No SVR: n=159 Median age: 48 (IQR 43-54) Female: 49% Caucasian: 47% African-American: 31% Hispanic: 14% Genotype 1: 68% Genotype: 2 or 3: 27% Other genotype: 5% Fibrosis stage: NR Clinical cirrhosis: 17%	Genotype, age, gender, race, comorbidities, cirrhosis, albumin level, white blood cell level, platelet count, SVR	Mortality, aHR, rate SVR: 0.11 (95% CI, 0.03 to 0.47), 2% (2/83) No SVR: Reference, 27% (43/159)	Grants: KL2 RR024983-04 and Adjusted Clinical Group Junior Faculty Development Award
Sinn 2008 ²³¹ Fair	Treatment duration: NR Followup: Median 4.6 years	SVR vs. no SVR SVR not defined IFN monotherapy or combination therapy with pegylated IFN or IFN and ribavirin	Antiviral treatment: n=490 SVR: n=296 No SVR: n=194 Mean age: 48.4 (SD±10.8) Female: 58% (286/490) Race: NR Genotype (n=240) Genotype 1b: 44% Genotype 1, non-1b: 2% Genotype 2: 52% Genotype 3 and 6: 2% Fibrosis stage (n=122) F0 and 1: 52% F3 and 4: 48%	Age, gender, diabetes, alcohol intake, body weight, HCV duration, platelet level, ALT, AST, AST:platelet ratio, AFP, genotype, fibrosis stage	Disease progression (increase in Child-Pugh score of ≥2 points, HCC, spontaneous bacterial peritonitis, bleeding gastric or esophageal varices, hepatic encephalopathy, or liver death), aHR SVR: 0.32 (95% CI, 0.11 to 0.91) No SVR: Reference	NR

Author year	Treatment duration			Variables accounted		
Quality	Followup	Intervention(s)	Population	for in analyses	Outcomes	Funding source
Tanaka 2000 ²²⁸ Fair	Treatment: 6 months Followup: Mean 55 to 68 months	SVR vs. relapse vs. nonresponders vs. no treatment SVR=Normalized ALT levels 24 weeks after completion of antiviral therapy Relapse=normalized ALT levels during treatment, elevated after 24 weeks of treatment IFN alpha 2a, recombinant IFN alpha 2b	Antiviral Treatment: n=594 SVR: n=175 Relapse: n=165 Nonresponders: n=254 No treatment: n=144 Mean age (years): 52 Female: 31% Race: NR Genotype 1: 75% Genotype 2: 25% F0: 2.4% F1: 54% F3: 40% F4: 2.9%	Age, sex, ALT, platelet count, fibrosis stage, HCV genotype, HCV viral load	HCC, aHR, rate SVR: 0.16 (95% CI, 0.04 to 0.62), 2% (3/175) Relapse: 0.27 (95% CI, 0.09 to 0.79), 3% (5/165) Non-responder: 0.74 (95% CI, 0.37 to 1.48),10% (25/254) No treatment: Reference, 12% (17/144) SVR vs. no SVR (calculated): 0.29 (95% CI, 0.07 to 1.28) SVR vs. relapse vs. non-responder All-cause mortality: 1.1% (2/175) vs. 0.6% (1/165) vs. 5.9% (15/254)	Osaka Prefectural Government and New Ten-Year Strategy for Center Control, Prevention of Cancer, from the Ministry of Health and Welfare of Japan
Tateyama 2011 ²²⁹ <i>Fair</i>	Treatment duration:NR Followup: <u>Mean:</u> 8.2 (SD±4.4) years	SVR vs. no SVR SVR=Undetectable HCV RNA 6 months after completion of antiviral therapy IFN monotherapy PEG-IFN monotherapy IFN and ribavirin combination PEG-IFN with ribavirin	Antiviral Treatment: n=373 SVR: n=139 No SVR: n=234 No treatment: n=334 (patient characteristics include untreated patients) Mean age (years): 57 Female: 50% Race: NR Genotype 1b: 72% Genotype 1b: 72% Genotype 2: 28% Other genotype: 0.3% F0 or F1: 39% F2: 27% F3: 17% F4: 17%	Age, sex, alcohol consumption, fibrosis stage, platelet count, albumin, AST, ALT, AFP, HCV genotype	HCC, aHR, 10-year cumulative incidence SVR: 0.099 (95% CI, 0.03 to 0.33), 3.1% No SVR: 0.70 (95% CI, 0.45 to 1.09), 14.6% No treatment: Reference, 29.5% SVR vs. no SVR (calculated): 0.14 (95% CI, 0.04 to 0.52)	Ministry of health, Labor and Welfare of Japan

Author year	Treatment duration			Variables accounted		
Quality	Followup	Intervention(s)	Population	for in analyses	Outcomes	Funding source
Tseng 2016 ²¹⁶ Fair	Treatment duration: 6 months Followup: mean 5.5 years (SD 2.5)	SVR vs. no SVR SVR=Undetectable HCV RNA 24 weeks after completion of antiviral therapy Subcutaneous PEG-IFN-α-2a or PEG-IFN-α-2b + oral ribavirin	Antiviral Treatment: n=145 SVR: n=95 No SVR: n=50 Mean age: 69 (SD±3.3) years Female: 60% Race: NR Genotype 1: 61% Fibrosis stage: NR	Sex, diabetes, HBV co- infection, alcoholism, fatty liver, HCV genotype	Cirrhosis, aHR, rate SVR: 0.29 (95% CI, 0.10 to 0.76), 15% (14/95) No SVR: Reference, 26% (13/50)	Dalin Tzu Chi General Hospital
Yoshida 1999 ²³⁰ <i>Faii</i> #	Treatment: NR Followup: mean 4.3 years	SVR vs. no SVR SVR=Undetectable HCV RNA 6 months after completion of antiviral therapy IFN	Cirrhosis: NR Antiviral Treatment: n=2357 SVR: n=789 No SVR: n=1568 No antiviral treatment: n=490 Mean age, years: 49.5 (SD±11.3) Female: 36% F0: 2% F1: 28% F2: 37% F3: 24% F4: 10% Genotype 1: 70% Genotype 2: 30%	Age, sex, fibrosis stage	HCC, aHR, rate SVR: 0.20 (95% CI, 0.099 to 0.39), 0% (10/789) No SVR: 0.63 (0.43 to 0.92), 1% (76/1568) No treatment: Reference, 12.0% (59/490) SVR vs. no SVR (calculated): 0.32 (95% CI, 0.14 to 0.70)	The Japan Ministry of Health and Welfare

Author year	Treatment duration			Variables accounted		
Quality	Followup	Intervention(s)	Population	for in analyses	Outcomes	Funding source
Yoshida 2002 ²¹³	Treatment duration:	SVR vs. no SVR	Antiviral treatment: n=2,430	Age, sex	Mortality, aHR, rate	Ministry of Health,
Fair#	Mean 137 days	SVR=Undetectable HCV RNA 6	SVR: n=817		SVR: 0.15 (95% CI,	Labor, and Welfare
		months after completion of	No SVR: n=1613		0.064 to 0.34), 0.9%	of Japan and
	Followup: Mean	antiviral therapy	No treatment: n=459		(7/817)	Ministry of
	5.4±2.4 years		Mean age (years): 50		No SVR: 0.47 (95% CI,	Education, Culture,
		IFN alfa: 84%	Female: 37%		0.29 to 0.76), 3.0%	Sports, Science,
		IFN beta: 14%	Race: NR		(49/1613)	and Lechnology of
		Both: 2%	Genotype: NR		No treatment:	Japan
			FU OF F1: 30%		Reference, 6.5%	
			F2: 37%		(30/459) SV(D vo no SV(D	
			F3. 23%		SVR VS. 110 SVR	
			F4. 9.3%		(Calculated). 0.32 (95%)	
					CI, 0.12 (0 0.00)	
					l iver-related mortality	
					aHR rate	
					SVR: 0.050 (95% CL	
					0.012 to 0.22). 0.2%	
					(2/817)	
					No SVR: 0.39 (95% CI.	
					0.22 to 0.68), 2.0%	
					(33/1613)	
					No treatment:	
					Reference, 5.0%	
					(23/459)	
					SVR vs. no SVR	
					(calculated): 0.13 (95%	
					CI, 0.03 to 0.61)	

Author year	Treatment duration		Des latio	Variables accounted		
	Followup	Intervention(s)	Population	for in analyses	Outcomes	Funding source
Yu 2006 ²¹⁴	I reatment duration:	SVR vs. no SVR	Antiviral Treatment: n=1057	Age, sex, ALI, fibrosis	HCC, aHR, rate	Department of
Fair	20-48 weeks	SVR=Undetectable HCV RNA 6	SVR: n=/15	stage, HCV genotype	SVR: HR 0.24 (95% CI,	Health, Taiwan
		months after completion of	No SVR: n=342		(0.13 to 0.46), 0.4%	and the Talwan
	Followup: Mean 5.18	antiviral therapy	No treatment: n=562			Liver Research
	years		Mean age (years): 46.9		NO SVR: 0.99 (95% CI,	Foundation
		IFIN alpha, combination	$(5D\pm11.49)$		$(0.04 \ (0 \ 1.51), 2.0\%)$	
			Page NP		(9/342)	
			Genotype 1: 46%		(6/562)	
			Other Genotypes: 54%		SVR vs. no SVR	
			Fibrosis stage: NR		(calculated): 0.24 (95%)	
			Cirrhosis: 16%		CL = 0.11 to 0.52	
					0., 0	
					Mortality, aHR, rate	
					SVR: 0.37 (95% CI,	
					0.14 to 0.99), 0.6%	
					(4/715)	
					No SVR: 1.32 (95% CI,	
					0.56 to 3.06), 3.5%	
					(12/342)	
					No treatment:	
					Reference, 1.8%	
					(10/562)	
					SVR vs. No SVR	
					(calculated): 0.28 (95%	
					CI, 0.08 to 1.02)	
					Liver-related mortality,	
					$ \Im VK: U.4\% (3/715)$	
					NU SVK: 3.2% (11/342)	
					(10/562)	

* Study populations overlap.

† Study populations overlap.

‡ Study population appears to overlap with Ioannou 2018.

§ Study populations overlap.

Study population appears to overlap with Backus, 2011, Butt, 2017, Cozen, 2013, Dieperink, 2014, and El-Serag, 2014.

¶ Study populations likely overlap.

Study populations appear to overlap.

Abbreviations: AFP = alpha fetoprotein; aHR = adjusted hazard ratio; ALT = alanine aminotransferase; AST = aspartate amino transferase; BMI = body mass index; CI = confidence interval; DAA = direct acting antiviral; FIB-4 = Fibrosis 4; HBV = hepatitis B virus; HCC = hepatocellular carcinoma; HCV = hepatitis C virus; HR = hazard ratio; IFN = interferon; IQR = interquartile range; IVDU

= injection drug use; PCR = polymerase chain reaction; PEG-IFN = pegylated interferon; NR = not reported; RNA = ribonucleic acid; SD = standard deviation; SVR = sustained virologic response; VA = Veterans Affairs.

Appendix B Table 16. Key Question 9: Quality Assessment of Studies of the Association Between Sustained Virologic Response After Antiviral Therapy and Clinical Outcomes

	Did the study attempt	Were the groups	Did the study			Did the study			
	to enroll all (or a	comparable at	use accurate	Were outcome		perform	Is there	Were outcomes	
	random sample of)	baseline on key	methods for	assessors and/or		appropriate	important	pre-specified and	
	patients meeting	prognostic	ascertaining	data analysts	Did the	statistical	differential loss	defined, and	
	inclusion criteria, or a	factors (e.g., by	exposures and	blinded to the	article	analyses on	to follow-up or	ascertained	
	random sample	restriction or	potential	exposure being	report	potential	overall high loss	using accurate	Quality
Author year	(inception cohort)?	matching)?	confounders?	studied?	attrition?	confounders?	to follow-up?	methods?	rating
Arase 2007 ²⁰⁴	Yes	No	Yes	Unclear	No	Yes	Unclear	Yes	Fair
Asahina 2010 ²¹⁷	Yes	Unclear	Yes	Unclear	No	Yes	Unclear	Yes	Fair
Backus 2011 ⁶⁹	Yes	No	Yes	No	Yes	Yes	Yes	Yes	Fair
Butt 2017 ²⁰⁵	Yes	Yes	Yes	Unclear	No	Yes	Unclear	Yes	Fair
Carrat 2019 ¹⁶⁸	Yes	Unclear	Yes	Unclear	Yes	Yes	No	Yes	Fair
Cozen 2013 ²⁰⁶	Yes	No	Yes	Unclear	No	Yes	Unclear	Yes	Fair
Dieperink 2014 ²⁰⁷	Yes	No	Yes	No	Yes	Yes	Unclear	Yes	Fair
Dohmen 2013 ²¹⁸	Yes	No	Yes	Unclear	No	Yes	Unclear	Yes	Fair
EI-Serag 2014 ²¹⁵	Unclear	Unclear	Yes	No	Yes	Yes	Unclear	Yes	Fair
Ikeda 1999 ²¹⁹	Unclear	Unclear	Unclear	Unclear	Yes	Yes	No	Yes	Fair
Imai 1998 ²²⁰	Yes	No	Yes	Yes	Yes	Yes	No	Yes	Fair
Imazeki 2003 ²⁰⁸	Yes	No	Yes	Unclear	Yes	Yes	No	Yes	Fair
Innes 2011 ²⁰⁹	Yes	No	Yes	Unclear	Yes	Yes	No	Yes	Fair
Ioannou 2018 ²²¹	Yes	No	Yes	No	No	Yes	Unclear	Yes	Fair
Izumi 2005 ²²²	Yes	Unclear	Yes	Unclear	No	No	Unclear	Yes	Fair
Kasahara 1998 ²²³	Unclear	No	Yes	Unclear	No	Yes	Unclear	Yes	Fair
Kasahara 2004 ²¹⁰	Yes	Yes	Yes	Unclear	No	No	Unclear	Yes	Fair
Kurokawa 2009 ²²⁴	Yes	Unclear	Yes	Unclear	No	No	Unclear	Yes	Fair
Lee 2017 ²²⁵	Yes	No	Yes	Unclear	No	Yes	Unclear	Yes	Fair
Maruoka 2012 ²¹¹	Yes	Unclear	Yes	Unclear	Yes	Yes	Yes	Yes	Fair
Okanoue 2002 ²²⁶	Yes	No	Yes	No	Yes	Yes	No	Yes	Fair
Osaki 2012 ²²⁷	Yes	No	Yes	Unclear	No	Yes	Unclear	Yes	Fair
Singal 2013 ²¹²	Yes	Unclear	Yes	Unclear	Yes	Yes	Yes	Yes	Fair
Sinn 2008 ²³¹	Yes	No	Yes	Unclear	Yes	Yes	No	Yes	Fair
Tanaka 2000 ²²⁸	Yes	No	Yes	No	Yes	Yes	Unclear	Yes	Fair
Tateyama 2011 ²²⁹	Unclear	Unclear	Yes	Unclear	No	Yes	Unclear	Yes	Fair
Tseng 2016 ²¹⁶	Yes	Unclear	Yes	Unclear	Yes	Yes	No	Yes	Fair
Yoshida 1999 ²³⁰	Yes	No	Yes	No	Yes	Yes	No	Yes	Fair
Yoshida 2002 ²¹³	Yes	No	Yes	No	Yes	No	No	Yes	Fair
Yu 2006 ²¹⁴	Yes	No	Yes	No	No	Yes	Unclear	Yes	Fair