Comparative Effectiveness Review Number 76

Treatment for Hepatitis C Virus Infection in Adults



Number 76

Treatment for Hepatitis C Virus Infection in Adults

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Preface

The Agency for Healthcare Research and Quality (AHRQ) conducts the Effective Health Care Program as part of its mission to organize knowledge and make it available to inform decisions about health care. As part of the Medicare Prescription Drug, Improvement, and Modernization Act of 2003, Congress directed AHRQ to conduct and support research on the comparative outcomes, clinical effectiveness, and appropriateness of pharmaceuticals, devices, and health care services to meet the needs of Medicare, Medicaid, and the Children's Health Insurance Program (CHIP).

AHRQ has an established network of Evidence-based Practice Centers (EPCs) that produce Evidence Reports/Technology Assessments to assist public- and private-sector organizations in their efforts to improve the quality of health care. The EPCs now lend their expertise to the Effective Health Care Program by conducting comparative effectiveness reviews (CERs) of medications, devices, and other relevant interventions, including strategies for how these items and services can best be organized, managed, and delivered.

Systematic reviews are the building blocks underlying evidence-based practice; they focus attention on the strength and limits of evidence from research studies about the effectiveness and safety of a clinical intervention. In the context of developing recommendations for practice, systematic reviews are useful because they define the strengths and limits of the evidence, clarifying whether assertions about the value of the intervention are based on strong evidence from clinical studies. For more information about systematic reviews, see www.effectivehealthcare.ahrq.gov/reference/purpose.cfm.

AHRQ expects that CERs will be helpful to health plans, providers, purchasers, government programs, and the health care system as a whole. In addition, AHRQ is committed to presenting information in different formats so that consumers who make decisions about their own and their family's health can benefit from the evidence.

Transparency and stakeholder input from are essential to the Effective Health Care Program. Please visit the Web site (www.effectivehealthcare.ahrq.gov) to see draft research questions and reports or to join an email list to learn about new program products and opportunities for input. Comparative Effectiveness Reviews will be updated regularly.

We welcome comments on this CER. They may be sent by mail to the Task Order Officer named below at: Agency for Healthcare Research and Quality, 540 Gaither Road, Rockville, MD 20850, or by email to epc@ahrq.hhs.gov.

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Treatment for Hepatitis C Virus Infection in Adults

Structured Abstract

Objectives. This report systematically reviews the comparative benefits and harms of current antiviral treatment regimens for chronic hepatitis C virus (HCV) infection in treatment-naïve adults.

Data sources. MEDLINE[®] (1947 to August 2012), the Cochrane Central Register of Controlled Trials (through 3rd quarter 2012), clinical trial registries, and reference lists.

Review methods. We used predefined criteria to determine study eligibility. We selected randomized trials of dual therapy with pegylated interferon (alfa-2a or alfa-2b) plus ribavirin or triple therapy with pegylated interferon (alfa-2a or alfa-2b), ribavirin, and either boceprevir or telaprevir that reported clinical outcomes, sustained virologic response (SVR), or harms. We also selected randomized trials or cohort studies that compared clinical outcomes in patients who experienced an SVR after antiviral therapy with patients who did not experience an SVR.

Results. We included 90 randomized trials and observational studies. No study evaluated the comparative effectiveness of current antiviral regimens on long-term clinical outcomes. In trials of treatment-naïve patients, the likelihood of achieving an SVR was slightly lower for dual therapy with pegylated interferon alfa-2b plus ribavirin than for dual therapy with pegylated interferon alfa-2a plus ribavirin, with a difference in absolute SVR rates of about 8 percentage points. There were no clear differences in estimates of relative effectiveness in patient subgroups defined by demographic or clinical characteristics, although absolute response rates were lower in older patients, Black patients, patients with high viral load, patients with more advanced fibrosis or cirrhosis, and patients with genotype 1 infection. Differences in harms were relatively small, with no difference in withdrawals due to adverse events, although dual therapy with pegylated interferon alfa-2b plus ribavirin was associated with a lower risk of serious adverse events than dual therapy with pegylated interferon alfa-2a plus ribavirin. In patients with genotype 2 or 3 infection, trials found dual therapy with pegylated interferon for 12 to 16 weeks associated with a lower likelihood of achieving SVR as compared with 24 weeks of therapy. Lower doses of pegylated interferon alfa-2b were less effective than standard doses, and limited evidence showed no clear differential effects of ribavirin dosing.

Five trials found triple therapy with pegylated interferon (alfa-2a or alfa-2b), ribavirin, and either boceprevir or telaprevir associated with higher likelihood of SVR (66–80 percent) than dual therapy with pegylated interferon plus ribavirin for genotype 1 infection, with an absolute increase in SVR rate of 22–31 percentage points. Triple therapy with boceprevir was associated with increased risk of hematological adverse events, and triple therapy with telaprevir was associated with increased risk of anemia and rash, including severe rash, versus dual therapy.

A large cohort study that controlled well for confounders found that patients with an SVR after antiviral therapy had a lower risk of all-cause mortality than patients with no SVR, with adjusted hazard ratio estimates ranging from 0.51 to 0.71, depending on genotype. Other, smaller cohort studies also found that SVR was associated with reduced risk of all-cause mortality and long-term complications of HCV infection, but had more methodological shortcomings.

Conclusions. Although there is no direct evidence on the comparative effects of current antiviral regimens on long-term clinical outcomes, SVR rates are substantially higher in patients with HCV genotype 1 infection who receive triple therapy with pegylated interferon (alfa-2a or alfa-2b), ribavirin, and boceprevir or telaprevir compared with dual therapy with pegylated interferon plus ribavirin. Achieving an SVR following antiviral therapy appears to be associated with decreased risk of all-cause mortality compared with no SVR, although estimates are susceptible to residual confounding.

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Executive Summary

Background

Hepatitis C virus (HCV) is the most common chronic bloodborne pathogen in the United States. HCV is primarily acquired by large or repeated percutaneous exposures to blood, with injection drug use being the strongest risk factor. Based on a national survey of households, approximately 1.6 percent of U.S. adults over 20 years of age have antibodies to HCV, indicating prior acute HCV infection. About 78 percent of patients with acute HCV infection develop chronic HCV infection, defined by the presence of persistent viremia.

Chronic HCV infection has a variable course, but it is a leading cause of complications from chronic liver disease, including cirrhosis, liver failure, and hepatocellular carcinoma (HCC). Chronic HCV infection is associated with an estimated 15,000 deaths each year in the United States, and it is the most common indication for liver transplantation among American adults, accounting for more than 30 percent of cases. The prevalence of chronic HCV infection is thought to have peaked in 2001 at 3.6 million people, and the yearly incidence has declined from more than 200,000 cases per year in the 1980s to around 16,000 cases in 2009. However, complications related to chronic HCV infection, which frequently occur only after decades of infection, are expected to rise for another 10 to 13 years.

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, but it is extremely difficult to design and carry out clinical trials long and large enough to provide direct evidence related to these outcomes. The sustained virologic response (SVR) rate, typically defined as the proportion of patients who experience a decline in HCV-RNA (hepatitis C virus ribonucleic acid) to undetectable levels 24 weeks following completion of antiviral treatment, is the standard marker of successful treatment in clinical trials because an SVR is strongly associated with the long-term absence of viremia. Recent studies have evaluated the association between achieving an SVR and reductions in mortality, liver failure, and cancer. Under the complex complex complex contents are contents as a successful treatment in clinical trials because an SVR is strongly associated with the long-term absence of viremia.

In the early 2000s, the combination of "pegylated" interferon plus ribavirin became the standard antiviral treatment for HCV infection. Pegylation refers to the cross-linking of polyethylene glycol molecules to the interferon molecule, which delays renal clearance and thereby permits less frequent dosing (once weekly vs. three times a week with standard interferon). Dual therapy with pegylated interferon plus ribavirin is associated with higher SVR rates (about 55–60 percent overall) than either standard interferon plus ribavirin or pegylated interferon monotherapy. Currently, two pegylated interferons are available: pegylated interferon alfa-2a and pegylated interferon alfa-2b. Although previous reviews found insufficient evidence to determine whether combination therapy with pegylated interferon alfa-2a or pegylated interferon alfa-2b plus ribavirin is more effective, more head-to-head trials directly comparing these two regimens are now available.

A number of factors affect response to antiviral treatment. The two major pretreatment predictors of SVR are the viral genotype and the pretreatment viral load. In the United States, genotype 1 infection is found in around three-quarters of HCV-infected patients. HCV genotype 1 infection is associated with a substantially lower response to antiviral treatment than infection with genotypes 2 and 3, which are present in about 20 percent of HCV-infected patients. A pretreatment viral load of <600,000 international units per milliliter (IU/mL) is

associated with higher likelihood of achieving an SVR. ¹¹ Other factors less consistently or less strongly associated with an increased likelihood of achieving an SVR include female sex, age less than 40 years, non-Black race, lower body weight (≤75 kg), absence of insulin resistance, elevated alanine aminotransferase levels, and absence of bridging fibrosis or cirrhosis on liver biopsy. ¹¹ Effects of race on the likelihood of achieving an SVR may be due in part to polymorphisms in the interleukin-28B (IL28B) gene. ^{21, 22}

An issue complicating antiviral treatment is the high rate of adverse effects observed with interferon-based therapy, including flulike symptoms, fatigue, and neuropsychiatric and hematologic adverse effects. Such adverse effects can be difficult to tolerate and can lead to premature discontinuation of therapy.

In 2011, the U.S. Food and Drug Administration (FDA) approved the first direct acting antiviral agents, boceprevir (trade name VictrelisTM) and telaprevir (trade name Incivek[®]), for treatment of chronic HCV genotype 1 infection.^{24, 25} Both drugs are classified as nonstructural 3/4A protease inhibitors, with a potential advantage of shorter duration of therapy (24 to 28 weeks) compared with standard dual therapy with pegylated interferon (alfa-2a or 2b) plus ribavirin for genotype 1 infection (48 weeks).²⁶⁻²⁸ Either drug is administered in combination with pegylated interferon (alfa-2a or 2b) plus ribavirin.

Understanding the comparative benefits and harms of the various antiviral regimens is critical for making informed treatment decisions in patients with chronic HCV infection, particularly given the availability of new treatment options. This review assesses the comparative effectiveness of antiviral treatments in adults with chronic HCV infection who have not received previous antiviral drug treatment. In addition to assessing the comparative effectiveness of different drug regimens, the review evaluates the effects of different medication doses, durations of therapy, and dosing strategies (such as weight-based or response-guided vs. fixed treatment). To help with individualized clinical decisionmaking regarding antiviral therapy for chronic HCV infection, the review also evaluates how comparative effectiveness varies depending on HCV genotype, viral load, and other demographic and clinical characteristics. Given the need to understand the effects of treatment in people with HCV infection identified by screening in order to assess the potential benefits and harms of screening, this review will be used, together with a separate review on HCV screening, ²⁹ by the U.S. Preventive Services Task Force to update its HCV screening recommendations.

Objectives

The following Key Questions are the focus of our report:

Key Question 1

- a. What is the comparative effectiveness of antiviral treatment in improving health outcomes in patients with HCV infection?
- b. How does the comparative effectiveness of antiviral treatment for health outcomes vary according to patient subgroup characteristics, including but not limited to HCV genotype, age, race, sex, stage of disease, or genetic markers?

Key Question 2

a. What is the comparative effectiveness of antiviral treatments on intermediate outcomes, such as the rate of SVR or histologic changes in the liver?

b. How does the comparative effectiveness of antiviral treatment for intermediate outcomes vary according to patient subgroup characteristics, including but not limited to HCV genotype, age, race, sex, stage of disease, or genetic markers?

Key Question 3

- a. What are the comparative harms associated with antiviral treatments?
- b. Do these harms differ according to patient subgroup characteristics, including HCV genotype, age, race, sex, stage of disease, or genetic markers?

Key Question 4

KQ = Key Question

Have improvements in intermediate outcomes (SVR, histologic changes) been shown to reduce the risk or rates of adverse health outcomes from HCV infection?

Analytic Framework

The analytic framework that guided this report is shown in Figure A. The numbers in the analytic framework indicate the Key Questions listed above. The population was patients with chronic HCV infection who were receiving antiviral therapy. The interventions were dual therapy with pegylated interferon (alfa-2a or alfa-2b) plus ribavirin, or triple therapy with pegylated interferon (alfa-2a or alfa-2b) plus ribavirin plus a protease inhibitor approved by the FDA (either boceprevir or telaprevir). Comparisons were between different regimens, as well as between regimens including the same drugs administered at different doses or for different durations. Intermediate outcomes were sustained virologic response and hepatic histological improvement. Final outcomes were morbidity and mortality from HCV infection (including hepatic cirrhosis, HCC, and liver transplantation rates) and quality of life, as well as harms of antiviral therapies (including flulike symptoms, hematologic effects, rash, and psychiatric effects).

KO. 1 Final Clinincal Outcomes Intermediate Antiviral Outcomes Treatment Patients with KQ. 4 KO. 2 Mortality Sustained chronic Morbidity virologic Hepatitis C Virus · Quality of life response infection Transmission Histological of Hepatitis C improvements Virus KQ. 3 Harms

Figure A. Analytic framework for treatment of hepatitis C infection in adults

Methods

Input From Stakeholders

The topic of treatment for HCV infection was nominated for a comparative effectiveness review (CER) in a public process. The Key Questions were proposed in the public nomination process and developed by investigators from the Evidence-based Practice Center (EPC) with contributions from expert Key Informants (KI), who helped refine Key Questions, identify important methodological and clinical issues, and define parameters for the review of evidence. The revised Key Questions were then posted to a public Web site for comment. The Agency for Healthcare Research and Quality (AHRQ) and the EPC agreed on the final Key Questions after reviewing the public comments and receiving additional advice from a Technical Expert Panel (TEP) convened for this report. We then drafted a protocol for this CER, which the TEP reviewed. Access it from the AHRQ Web site, where it was posted in November 2011: (www.effectivehealthcare.ahrq.gov/index.cfm/search-for-guides-reviews-and-reports/?pageaction=displayproduct&productid=855).

A multidisciplinary group of clinicians, researchers, and patient advocates with expertise in hepatitis C treatment and research were selected to serve as the TEP members to provide high-level content and methodological expertise throughout the development of the review. Prior to participation in this report, the TEP members disclosed all financial or other conflicts of interest. The AHRQ Task Order Officer and the authors reviewed all of these disclosures and determined the panel members had no significant conflicts of interest that precluded participation. KIs and TEP members had expertise in hepatology, epidemiology, screening, and primary care. TEP members and other experts were invited to provide external peer review of the draft report.

Search Strategy and Study Selection

To identify articles relevant to each Key Question, a research librarian searched the Cochrane Database of Systematic Reviews, the Cochrane Central Register of Controlled Trials, and Ovid MEDLINE® from 1947 to April 2011 (see Appendix A for the search strategies), and a final updated search was conducted in August 2012. The search strategies were peer reviewed by another research librarian and revised prior to finalization. Unpublished trials were sought by searching clinical trial registries (ClinicalTrials.gov, Current Controlled Trials, Clinical Trial Results, WHO Trial Registries) and grants databases (NIHRePORTER, HSRProj, and AHRQ GOLD). Scientific Information Packets on unpublished and published trials were solicited from manufacturers of included antiviral drugs through the Scientific Resource Center. We also hand-searched the reference lists of relevant studies. Searches were updated before the report was finalized to identify relevant new publications.

Studies were selected according to criteria developed for inclusion and exclusion. The selection criteria were based on the Key Questions and the populations, interventions, comparators, outcomes, timing, and setting (PICOTS) approach. Papers were selected for full review if they were about chronic HCV infection, were relevant to Key Questions in the analytic framework, and met the predefined inclusion criteria. To evaluate the potential effects of publication bias, we included trials published only as conference abstracts of sensitivity analyses. We restricted inclusion to English language articles. Studies of nonhuman subjects were also excluded, and studies had to include original data.

Abstracts and full-text articles were dual reviewed for inclusion and exclusion for each Key Question. Full-text articles were obtained for all studies identified as potentially meeting inclusion criteria. Two investigators independently reviewed all full-text articles for final inclusion or exclusion, and discrepancies were resolved through discussion and consensus, with a third investigator making the final decision if necessary.

Data Extraction and Quality Assessment

We assessed the quality of each study based on predefined criteria (Appendix E). We adapted criteria from methods proposed by Downs and Black (observational studies),³⁰ the USPSTF,³¹ and the Quality Assessment of Diagnostic Accuracy Studies-2 Group.³² The criteria used are consistent with the approach recommended by AHRQ in the Methods Guide for Effectiveness and Comparative Effectiveness Reviews (Methods Guide).³³ We used the term "quality" rather than the alternate term "risk of bias." Although both refer to internal validity, "quality" may be more familiar to most users and has potential advantages in terms of readability.

We rated the quality of each randomized trial based on the methods used for randomization, allocation concealment, and blinding; the similarity of compared groups at baseline; maintenance of comparable groups; adequate reporting of dropouts, attrition, crossover, adherence, and contamination; loss to followup; the use of intent-to-treat analysis; and ascertainment of outcomes.³¹

We rated the quality of each cohort study based on whether it used nonbiased selection methods to create an inception cohort; whether it evaluated comparable groups; whether rates of loss to followup were reported and acceptable; whether it used accurate methods for ascertaining exposures, potential confounders, and outcomes; and whether it performed appropriate statistical analyses of potential confounders. ³¹

Following assessment of individual quality criteria, individual studies were rated good, fair, or poor quality, as defined below.³³

Good-quality studies are considered likely to be valid. Good-quality studies clearly describe the population, setting, interventions, and comparison groups; use a valid method for allocation of patients to interventions; clearly report dropouts and have low dropout rates; use appropriate methods for preventing bias; and appropriately measure outcomes and fully report results.

Fair-quality studies have some methodological deficiencies but no flaw or combination of flaws judged likely to cause major bias. The study may be missing information, making it difficult to assess its methods or assess limitations and potential problems. The fair-quality category is broad, and studies with this rating vary in their strengths and weaknesses—the results of some fair-quality studies are likely to be valid, while others are only probably valid.

Poor-quality studies have significant flaws that may invalidate the results. They have a serious or fatal flaw in design, analysis, or reporting; large amounts of missing information; or discrepancies in reporting. The results of these studies are judged to be at least as likely to reflect flaws in the study design as true effects of the interventions under investigation. We did not exclude studies rated poor quality a priori, but they were considered to be the least reliable studies when synthesizing the evidence, particularly when discrepancies between studies were present.

We recorded factors important for understanding the applicability of studies, such as whether the publication adequately described the study population, how similar patients were to populations likely to be targeted by screening, whether differences in outcomes were clinically (as well as statistically) significant, and whether the interventions and tests evaluated were reasonably representative of standard practice.³⁴ We also recorded the funding source and role of the sponsor. We did not assign a rating of applicability (such as high or low) because applicability may differ based on the user of this report.

Data Synthesis and Rating the Strength of the Body of Evidence

We performed meta-analysis of trials that evaluated similar populations, interventions, comparisons, and outcomes to estimate pooled relative risks. When present, statistical heterogeneity was explored through subgroup and sensitivity analyses, as well as qualitatively. Subgroup analyses were performed in groups stratified by HCV genotype as well as by race, age, body weight, viral load, stage/severity of disease, and IL-28b status when these data were available. We performed sensitivity analysis by excluding poor-quality studies and outlier trials, and by including results from studies published only as abstracts to evaluate the stability of estimates and conclusions. We did not perform meta-analyses for Key Question 4 because all studies were observational and had important methodologic shortcomings. These studies were synthesized qualitatively.

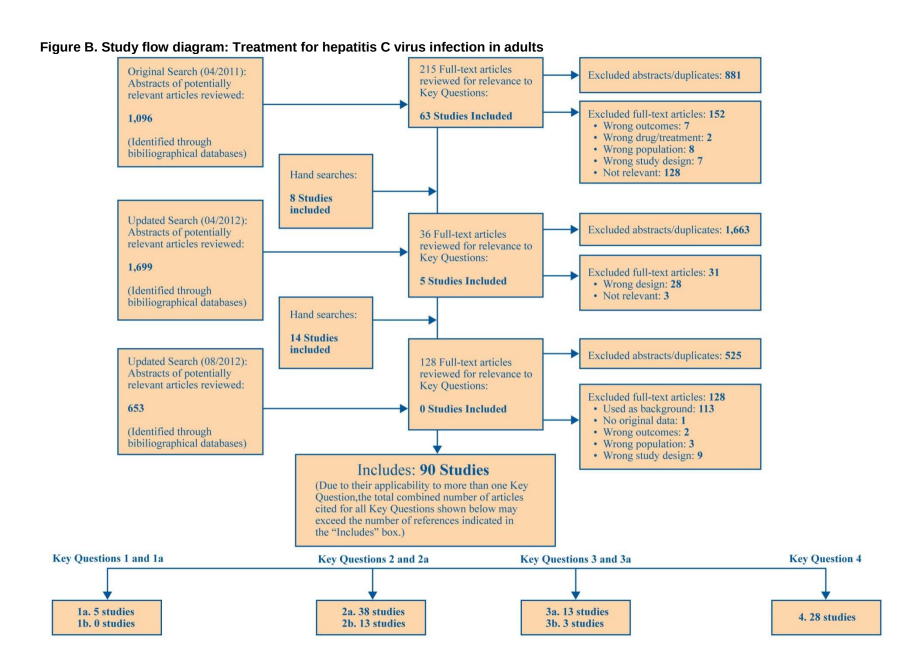
We rated the strength of evidence for each Key Question using the four categories recommended in the AHRQ Methods Guide. We synthesized the overall quality of each body of evidence based on the type and quality of studies (graded good, fair, or poor); the precision of the estimate of effect based on the number and size of studies and confidence intervals for the estimates (graded high, moderate, or low); the consistency of results between studies (graded high, moderate, or low); and the directness of the evidence linking the intervention and health outcomes (graded direct or indirect). We did not downgrade a body of evidence for directness that evaluated an intermediate outcome if the intermediate outcome was the specific focus of the Key Question. We were not able to formally assess for publication bias due to small numbers of studies, methodological shortcomings, or differences across studies in designs, measured outcomes, and other factors.

We graded the strength of evidence for each comparison and outcome by using the four categories recommended in the AHRQ Methods Guide: ³³ A "high" grade indicates high confidence that the evidence reflects the true effect and that further research is very unlikely to change our confidence in the estimate of effect and will not change the estimate. A "moderate" grade indicates moderate confidence that the evidence reflects the true effect and that further research may change our confidence in the estimate of effect and may change the estimate. A "low" grade indicates low confidence that the evidence reflects the true effect and that further research is likely to change the confidence in the estimate of effect and is likely to change the estimate. An "insufficient" grade indicates evidence either is unavailable or is too limited to permit any conclusion.

Results

The search and selection of articles are summarized in the study flow diagram (Figure B). Of the 1,096 citations identified at the title and abstract level in the original search, 215 articles met inclusion criteria and were selected for further review of the full text. From updated searches and peer reviewer suggested citations, an additional 2,352 citations were identified, and 164 of these met inclusion criteria and were selected for full-text review. Of the 379 articles reviewed at the full-text level, a total of 90 studies met inclusion criteria.

No study evaluated comparative effectiveness of current antiviral regimens on long-term clinical outcomes such as mortality, complications of chronic HCV infection, or quality of life.



ES-7

Dual Therapy Regimens with Pegylated Interferon Plus Ribavirin

In trials of treatment-naïve patients, dual therapy with pegylated interferon alfa-2b plus ribavirin was associated with a slightly lower likelihood of achieving an SVR than dual therapy with pegylated interferon alfa-2a plus ribavirin, with a difference in absolute SVR rates of about 8 percentage points. ^{16-19, 36-38} In patients with genotype 2 or 3 infection, dual therapy for 12 to 16 weeks appears to be associated with a lower likelihood of SVR, compared with dual therapy for 24 weeks, with no differences between 24 weeks and longer courses of therapy. ³⁹⁻⁴⁴ In trials comparing different doses of dual therapy with pegylated interferon plus ribavirin, lower doses of pegylated interferon alfa-2b were less effective than standard doses, ^{41, 45-49} and limited evidence found no clear differential effects of ribavirin dosing. ^{39, 50}

There were no clear differences in estimates of relative effectiveness between dual therapy with pegylated interferon alfa-2a plus ribavirin versus dual therapy with pegylated interferon alfa-2b plus ribavirin in patient subgroups defined by demographic or clinical characteristics, although absolute response rates were lower in older patients, Black patients, patients with high viral load, patients with more advanced fibrosis or cirrhosis, and patients with genotype 1 infection. ^{16, 17, 19, 51}

Differences in harms between dual therapy with pegylated interferon alfa-2a plus ribavirin versus pegylated interferon alfa-2b plus ribavirin were relatively small, with no differences in withdrawals due to adverse events, although dual therapy with pegylated interferon alfa-2b was associated with a lower risk of serious adverse events. ^{16-19, 38, 52}

Triple Therapy Regimens With Pegylated Interferon, Ribavirin, and Either Boceprevir or Telaprevir

Trials of antiviral regimens including either boceprevir or telaprevir have been primarily conducted in patients with genotype 1 infection. Triple antiviral regimens (pegylated interferon alfa-2a or alfa-2b, ribavirin, and boceprevir or telaprevir) were associated with a substantially increased likelihood of achieving an SVR than dual therapy with pegylated interferon alfa-2a or alfa-2b plus ribavirin). ^{26-28, 53-57}

Two trials found triple therapy with boceprevir for 48 weeks (dual therapy with pegylated interferon alfa-2b plus ribavirin for 4 weeks followed by 44 weeks of triple therapy with the addition of boceprevir) was associated with a higher likelihood of SVR than dual therapy with pegylated interferon alfa-2b plus ribavirin for 48 weeks (pooled relative risk [RR] 1.81, 95% confidence interval [CI] 1.58 to 2.06, I^2 =0.0%) with an absolute increase in SVR rate of 31 percentage points (95% CI 23 to 39).

Three trials found triple therapy with telaprevir for 24 weeks (pegylated interferon alfa-2a, ribavirin, and telaprevir triple therapy for 12 weeks followed by 12 weeks of pegylated interferon alfa-2a plus ribavirin without telaprevir) was associated with a higher likelihood of SVR than dual therapy with pegylated interferon alfa-2a plus ribavirin for 48 weeks (pooled RR 1.48, 95% CI 1.26 to 1.75, I²=0.0%), with an absolute increase in SVR rate of 22 percentage points (95% CI 13 to 31). One trial found response-guided telaprevir triple therapy (8 or 12 weeks of pegylated interferon alfa-2a, ribavirin, and telaprevir followed by 12 or 36 weeks of response-guided dual therapy with pegylated interferon alfa-2a plus ribavirin) was associated with a higher likelihood of SVR than dual therapy with pegylated interferon alfa-2a plus ribavirin for 48 weeks (RR 1.6, 95% CI 1.4 to 1.9), with an absolute increase in SVR rate of 25–31 percentage points. States of 25–31 percentage points.

Relative estimates of the effects of triple therapy with either boceprevir or telaprevir, compared with dual therapy, were similar across subgroups, except in patients with low viral load, in whom triple therapy was no more effective than dual therapy in achieving an SVR. Triple therapy with boceprevir was associated with increased risk of hematological adverse events and triple therapy with telaprevir with increased risk of anemia and rash (including severe rash) than dual therapy; adverse events were generally self-limited with discontinuation of therapy. All antiviral regimens were associated with a high incidence of flulike symptoms, with small or no clear differences in risk.

Sustained Virologic Response After Antiviral Therapy and Clinical Outcomes

A large cohort study that was well controlled for confounders found that patients with an SVR after antiviral therapy had a lower risk of all-cause mortality than patients with no SVR (adjusted hazard ratio estimates 0.51 to 0.71). Eighteen other cohort studies also found SVR associated with reduced risk of all-cause mortality, liver-related mortality, and other hepatic complications rather than no SVR, but had more methodological shortcomings. 9, 58-74 Ten of the studies were conducted in Asian countries and might not be directly applicable to U.S. populations.

Discussion

Key Findings and Strength of Evidence

The evidence reviewed in this study is summarized in Table A. The specific domain scores used to determine the overall strength of evidence for each body of evidence are shown in Appendix G. We identified no studies that evaluated comparative effectiveness of current antiviral regimens on long-term clinical outcomes such as mortality, complications of chronic HCV infection, or quality of life. Such trials would be difficult to design and carry out due to the long time required for complications of chronic HCV infection to develop in most patients.

Dual Therapy Regimens With Pegylated Interferon and Ribavirin

In lieu of direct evidence on long-term clinical outcomes, SVR rates are the primary outcome to assess comparative benefits of different antiviral regimens. In trials of treatment-naïve patients, the likelihood of achieving an SVR was slightly lower with dual therapy with pegylated interferon alfa-2b plus ribavirin compared with dual therapy with pegylated interferon alfa-2a plus ribavirin (pooled RR 0.87, 95% CI 0.80 to 0.95; I²=27.4%), with a difference in absolute SVR rates of about 8 percentage points. Although the largest study, the Individualized Dosing Efficacy vs. Flat Dosing to Assess Optimal Pegylated Interferon Therapy (IDEAL) trial, found no difference in SVR rates for dual therapy with pegylated interferon alfa-2a plus ribavirin compared with dual therapy with pegylated interferon alfa-2b plus ribavirin, excluding the IDEAL trial from pooled analyses, resulted in similar effect estimates. Although there was no difference between types of dual therapy regimens in risk of withdrawals due to adverse events, dual therapy with pegylated interferon alfa-2b plus ribavirin was associated with a lower risk of serious adverse events than dual therapy with pegylated interferon alfa-2a plus ribavirin (pooled RR 0.76, 95% CI 0.71 to 0.88, I²=0.0%), suggesting a potential tradeoff between greater benefits and greater harms. However, serious adverse events were only reported in two trials, ^{18, 19} and the

rate of serious adverse events was relatively low (about 4 percent overall in IDEAL), with an absolute difference of about 1 percent, and adverse events with antiviral treatments generally resolve following discontinuation of therapy. Trials found no clear difference in estimates of relative effectiveness of dual therapy with pegylated interferon alfa-2a plus ribavirin compared with dual therapy with pegylated interferon alfa-2b plus ribavirin in patient subgroups stratified by age, sex, race, viral load, fibrosis stage, and genotype, although absolute response rates were lower in older patients, Black patients, patients with high viral load, patients with more advanced fibrosis or cirrhosis, and patients with genotype 1 infection. SVR rates ranged from 24 to 42 percent lower in patients with genotype 1 infection compared with patients with genotype 2 or 3.

In patients with genotype 2 or 3 infection, dual therapy for 12 to 16 weeks appears to be associated with a lower likelihood of SVR compared with dual therapy for 24 weeks, with no differences between 24 weeks and longer courses of therapy. Standard doses of pegylated interferon alfa-2b were more effective than lower doses (no trials compared different doses of pegylated interferon alfa-2a). Although trials comparing different ribavirin doses found no clear differences, they evaluated different dose comparisons, precluding firm conclusions. Although trials comparisons, precluding firm conclusions.

Triple Therapy Regimens With Pegylated Interferon, Ribavirin, and Either Boceprevir or Telaprevir

Trials of triple therapy regimens with the protease inhibitors boceprevir or telaprevir (both approved by the FDA in 2011) in treatment-naïve patients with genotype 1 infection found each associated with substantially higher SVR rates than standard dual therapy without a protease inhibitor. SVR rates with triple therapy were similar to the 70–80 percent observed with dual therapy in patients with genotype 2 or 3 infection. ^{23, 26-28, 53-57, 77} Trials that evaluated the telaprevir regimen recommended by the FDA (12 weeks of triple therapy with telaprevir followed by response-guided duration of 12 or 36 weeks of dual therapy) reported SVR rates of 75–80 percent. ^{54, 56} Trials that evaluated the boceprevir regimen recommended by the FDA for antiviral-naïve patients with cirrhosis (4 weeks of dual therapy lead-in followed by 44 weeks of triple therapy with boceprevir) reported SVR rates of 66–75 percent. ^{26, 28} Trials that evaluated other regimens in antiviral naïve patients, including fixed duration telaprevir regimens, shorter fixed duration triple therapy boceprevir therapy, and boceprevir without dual therapy lead-in, reported similar or lower SVR rates.

As with the head-to-head trials of dual therapy with pegylated interferon alfa-2a plus ribavirin compared with pegylated interferon alfa-2b plus ribavirin, RR estimates for triple, compared with dual, therapy were similar (or there were no clear differences) in patient subgroups based on age, sex, or race, although absolute SVR rates were lower in older patients and Black patients. In two trials, triple therapy with boceprevir was no more effective than dual therapy in the subgroup of patients with lower HCV-RNA viral load (<600,000 or <800,000 IU/mL), ^{26, 28} but two trials of triple therapy with telaprevir were inconsistent in showing differential effects depending on baseline viral load. ^{54, 55} There was insufficient evidence to evaluate relative effectiveness of triple, compared with dual, therapy based on fibrosis stage.

In addition to a higher likelihood of SVR, another advantage of triple therapy regimens in patients with genotype 1 infection is the potential for a shorter duration of treatment (24 or 28 weeks in patients with early virologic response, compared with the standard 48 weeks of dual therapy with pegylated interferon plus ribavirin). Shorter courses of treatment would probably be

appealing to patients, given the frequency of bothersome flulike symptoms associated with interferon-based therapy. On the other hand, triple therapy regimens were associated with increased risk of certain harms, in particular hematological adverse events (neutropenia, anemia, and thrombocytopenia) with boceprevir, and anemia and rash (including severe rash in up to about 10 percent of patients, which could result in treatment discontinuation) with telaprevir. However, there was no clear increase in risk of serious adverse events or overall withdrawal due to adverse events with use of protease inhibitors, and the adverse events appear to be self-limited following drug discontinuation.

Sustained Virologic Response After Antiviral Therapy, and Clinical Outcomes

The strongest evidence on the association between an SVR after antiviral therapy and improved clinical outcomes is a large U.S. Department of Veterans Affairs (VA) cohort study (n=16,864) that adjusted for many confounders and found decreased risk of all-cause mortality compared with no SVR across patient groups stratified by genotype (adjusted hazard ratio [HR] 0.71 [0.60–0.86], 0.62 [0.44–0.87] and 0.51 [0.35–0.75] for genotypes 1, 2, and 3, respectively).8 Despite controlling for important confounders, the possibility of residual confounding is suggested by the very rapid separation of mortality curves for people with an SVR versus those without an SVR, which was observed at 3 months after assessment for SVR. This is more rapid than expected given the typically prolonged natural history of HCV infection. Therefore, estimates of effects of SVR on clinical outcomes from this study may be exaggerated, although it is not possible to determine to what degree. Eighteen other cohort studies also found an SVR after antiviral therapy associated with decreased risk of all-cause mortality and complications of chronic HCV infection, including studies specifically of patients with baseline cirrhosis, but had more methodological shortcomings. In addition, 10 of the 19 studies were conducted in Asia, where the incidence of HCC in patients with chronic HCV infection is higher than in the United States, ⁷⁸ potentially limiting their generalizability. Other studies found an SVR after antiviral therapy associated with better scores on measures of quality of life than with no SVR, but those studies focused on short-term outcomes and typically did not adjust for confounders or blind patients to SVR status when assessing outcomes.

Key Question	Outcome	Summary of Evidence	Strength of Evidence
Koy Question 1a	Long-term clinical outcomes	No evidence.	Insufficient
Key Question 1a What is the comparative effectiveness of antiviral	Short-term mortality	Three trials that compared current antiviral regimens ^a found no differences in risk of short-term mortality, but reported very few (20 total) events.	Low
treatment in improving health outcomes in patients with HCV infection?	Short-term quality of life	One open-label randomized trial of patients with genotype 4 infection found dual therapy with pegylated interferon alfa-2a plus ribavirin associated with statistically significant, slightly better short-term scores on some quality of life assessments compared with dual therapy with pegylated interferon alfa-2b plus ribavirin.	Low

for hepatitis C (co	Outcome	Summary of Evidence	Strength of Evidence
Key Question 1b How does the comparative effectiveness of antiviral treatment for health outcomes vary according to patient subgroup characteristics?	Any clinical outcome	No evidence.	Insufficient
		egylated Interferon Alfa-2b Plus Ribavirin vs. Du Pegylated Interferon Alfa-2a Plus Ribavirin	ıal Therapy
	Sustained virologic response	Seven trials found dual therapy with standard doses of pegylated interferon alfa-2b plus ribavirin associated with lower likelihood of achieving an SVR than pegylated interferon alfa-2a plus ribavirin (pooled RR 0.87, 95% CI 0.80 to 0.95; I ² =27.4%), with an absolute difference in SVR rates of 8 percentage points (95% CI 3 to 14).	Moderate
	Dual Therapy With Pegylated Interferon Alfa-2a or Alfa-2b Plus Ribavirin: Duration Effects		
Key Question 2a What is the comparative effectiveness of antiviral	Sustained virologic response	Two trials of patients with genotype 2 or 3 infection found no difference in likelihood of achieving an SVR between 48 vs. 24 weeks of dual therapy with pegylated interferon alfa-2a plus ribavirin (pooled RR 0.97, 95% CI 0.84 to 1.1; I ² =43%).	Moderate
treatments on intermediate outcomes?	Sustained virologic response	Four trials of patients with genotype 2 or 3 infection found 24 weeks of dual therapy with pegylated interferon (alfa-2a or alfa-2b) more effective than 12-16 weeks for achieving an SVR (pooled RR 1.15, 95% CI 1.02 to 1.29; I ² =79.5%). Relative risk estimates ranged from 1.01 to 1.33 in the four trials and may have varied in part due to differences across studies in ribavirin dosing.	Moderate
	Sustained virologic response	Three trials of patients with genotype 2 or 3 infection with a rapid virologic response (undetectable HCV-RNA by week 4) found no differences between 24 vs. 12-16 weeks of dual therapy with pegylated interferon (alfa-2a or alfa-2b) plus ribavirin (pooled RR 0.99, 95% CI 0.86 to 1.14; I ² =66.7%). Relative risk estimates ranged from 0.89 to 1.12.	Moderate

for hepatitis C (co	Outcome	Summary of Evidence	Strength of Evidence
	Dual Therapy With Pegylated Interferon Alfa-2a or Alfa-2b Plus Ribavirin: Dose Effects		
	Sustained virologic response	Six trials of patients with genotype 2 or 3 infection found lower doses of pegylated interferon alfa-2b (0.75-1.0 mcg/kg or 50 mcg) associated with lower likelihood of achieving an SVR than higher doses (1.5 mcg/kg or 100-150 mcg) (pooled RR 0.90; 95% CI 0.81 to 0.99; I ² =20.2%).	Moderate
	Sustained virologic response	Three trials of patients with genotype 2 or 3 infection who did not specifically have advanced fibrosis or cirrhosis found no clear difference in likelihood of SVR between lower doses of ribavirin (400 or 800 mg flat dose or 600 to 800 mg weight-based dose) vs. higher doses (800 or 1,200 mg flat dose or 800 to 1400 mg weight-based dose).	Moderate
Key Question 2a What is the comparative effectiveness of antiviral treatments on	Sustained virologic response	One small trial of patients with genotype 2 or 3 infection (N=60) and advanced fibrosis or cirrhosis (Ishak stage 4-6) found 600 to 800 mg daily of ribavirin associated with lower likelihood of SVR than 1000 to 1200 mg daily (45 vs. 72 percent, RR 0.62, 95% C I 0.40 to 0.98).	Low
intermediate outcomes?	Triple Therapy With Pegylated Interferon Alfa-2b, Ribavirin, and Boceprevir vs. Dual Therapy With Pegylated Interferon Alfa-2b Plus Ribavirin		
(continued)	Sustained virologic response	Two trials of patients with genotype 1 infection found triple therapy with boceprevir (pegylated interferon alfa-2b plus ribavirin for 4 weeks, followed by the addition of boceprevir for 44 weeks) associated with higher likelihood of SVR than dual therapy with pegylated interferon alfa-2b plus ribavirin therapy for 48 weeks (pooled RR 1.81; 95% CI 1.58 to 2.06; I ² =0.0%), with an absolute increase in SVR rate of 31% (95% CI 23 to 39).	Moderate
	Sustained virologic response	One trial of patients with genotype 1 infection found 48 weeks of triple therapy with boceprevir using a low dose of ribavirin (400-1000 mg daily) associated with a non–statistically significant trend toward lower likelihood of SVR compared with 48 weeks of triple therapy with a standard ribavirin dose (800-1400 mg daily) (36% vs. 50%, RR 0.71, 95% CI 0.39 to 1.3).	Low

for hepatitis C (co	Outcome	Summary of Evidence	Strength of Evidence
	Triple Therapy With Pegylated Interferon Alfa-2a or Alfa-2b, Ribavirin, and Telaprevir vs. Dual Therapy With Pegylated Interferon Alfa-2a or Alfa-2b Plus Ribavirin		
	Sustained virologic response	Three trials of patients with genotype 1 infection found triple therapy with telaprevir for 24 weeks (12 weeks of pegylated interferon alfa-2a, ribavirin, and telaprevir followed by 12 weeks of pegylated interferon alfa-2a plus ribavirin) associated with a higher likelihood of SVR than dual therapy with pegylated interferon alfa-2a plus ribavirin for 48 weeks (pooled RR 1.48, 95% CI 1.26 to 1.75; I ² =0.0%), with an absolute increase in SVR rate of 22% (95% CI 13 to 31).	Moderate
Key Question 2a What is the	Sustained virologic response	One trial of patients with genotype 1 infection found no difference in likelihood of SVR between triple therapy with pegylated interferon, ribavirin, and telaprevir for 12 weeks vs. dual therapy with pegylated interferon alfa-2a plus ribavirin for 48 weeks.	Moderate
comparative effectiveness of antiviral treatments on intermediate outcomes? (continued)	Sustained virologic response	One trial of patients with genotype 1 infection found response-guided triple therapy with telaprevir (pegylated interferon alfa-2a, ribavirin, and telaprevir for 8 or 12 weeks followed by a response-guided dual therapy with pegylated interferon alfa-2a plus ribavirin for an additional 12 or 36 weeks) associated with a higher likelihood of SVR than dual therapy with pegylated interferon alfa-2a plus ribavirin for 48 weeks (RR 1.6, 95% CI 1.4 to 1.9), with an absolute increase in SVR rate ranging from 25% to 31%. The regimen with 8 weeks of telaprevir was associated with a slightly lower SVR rate than the 12 week telaprevir regimen (69% vs. 75%).	Low
	Sustained virologic response	One trial of patients with genotype 1 infection found no difference in likelihood of SVR between triple therapy with telaprevir for 48 weeks (12 weeks of triple therapy with pegylated interferon alfa-2a, ribavirin, and telaprevir followed by 36 weeks of dual therapy with pegylated interferon alfa-2a plus ribavirin) vs. triple therapy with telaprevir for 24 weeks (12 weeks of triple therapy followed by 12 weeks of dual therapy).	Low

for hepatitis C (co			Strength of
Key Question	Outcome	Summary of Evidence	Evidence
	Triple Therapy With Pegylated Interferon Alfa-2a, Ribavirin, and Telaprevir: Dose Effects of Pegylated Interferon Alfa-2a vs. Alfa-2b and Duration Effects		
Key Question 2a What is the comparative effectiveness of antiviral treatments on	Sustained virologic response	One trial of response-guided triple therapy with telaprevir (24 or 48 weeks, based on absence or presence of HCV-RNA from weeks 4 through 20) found similar SVR rates (81–85%) for regimens that varied on telaprevir dose (750 mg tid vs. 1125 mg bid) and type of pegylated interferon (alfa-2a or alfa-2b).	Low
intermediate outcomes? (continued)	Sustained virologic response	One trial of patients with an extended rapid virologic response to initial triple therapy with telaprevir reported similar, high (92% and 88%) SVR rates in patients randomized to a total of 24 or 48 weeks of therapy.	Low
		Pegylated Interferon Alfa-2b Plus Ribavirin vs. Du	ial Therapy
	Sustained virologic response	Pegylated Interferon Alfa-2a Plus Ribavirin The largest randomized trial (n=3070) of dual therapy with pegylated interferon alfa-2a plus ribavirin vs. dual therapy with pegylated interferon alfa-2b plus ribavirin found no clear differences in relative risk estimates for SVR in genotype 1 patients stratified by race, sex, age, baseline fibrosis stage, or baseline viral load. Characteristics associated with lower absolute SVR rates across dual therapy regimens were older age, Black race, advanced fibrosis or cirrhosis, and high baseline viral load.	Low
Key Question 2b How does the comparative effectiveness of antiviral treatment for intermediate		Four randomized trials of dual therapy with pegylated interferon alfa-2a plus ribavirin vs. dual therapy with pegylated interferon alfa-2b plus ribavirin found no clear differences in relative risk estimates for SVR in patients stratified by genotype. Genotype 1 infection was associated with a lower absolute SVR rate than genotypes 2 or 3.	
outcomes vary according to patient subgroup characteristics?	Sustained virologic response	With Pegylated Interferon Alfa-2b Plus Ribavirin Two trials of triple therapy with boceprevir for 48 weeks (4 weeks of dual therapy lead-in with pegylated interferon plus ribavirin followed by 44 weeks of triple therapy with pegylated interferon, ribavirin, and boceprevir) found no difference in relative risk estimates for SVR in men vs. women, and no clear difference in relative risk estimates for Black vs. non-Black patients. Black race was associated with a lower absolute SVR rate than non-Black race.	Moderate
	Sustained virologic response	Two trials found triple therapy with pegylated interferon alfa-2b, ribavirin, and boceprevir associated with higher likelihood of achieving SVR than dual therapy with pegylated interferon alfa-2b plus ribavirin in patients with high baseline HCV-RNA viral load (>600,000 or ≥800,000 IU/mL), but found no difference in likelihood of SVR in patients with lower viral load.	Moderate

for hepatitis C (co	Outcome	Summary of Evidence	Strength of Evidence
	Triple Therapy With Pegylated Interferon Alfa-2a or Alfa-2b, Ribavirin, and Telaprevir vs. Dual Therapy With Pegylated Interferon Alfa-2a or Alfa-2b Plus Ribavirin		
Key Question 2b How does the comparative effectiveness of antiviral treatment for intermediate outcomes vary according to patient subgroup characteristics? (continued)	Sustained virologic response	One trial of response-guided triple therapy with telaprevir (12 weeks of pegylated interferon alfa-2a, ribavirin, and telaprevir followed by response-guided dual therapy with pegylated interferon alfa-2a and ribavirin) vs. dual therapy with pegylated interferon plus ribavirin for 48 weeks found no clear differences in relative risk estimates in patients stratified by age, sex, race, baseline fibrosis status, or body mass index. Characteristics associated with lower absolute rates of SVR were older age, Black race, advanced fibrosis or cirrhosis, and higher body mass index. One other trial of 24-week fixed duration triple therapy with telaprevir, pegylated interferon alfa-2b, and ribavirin vs. 48 weeks of dual therapy found no differences in estimates of effect in patients stratified by sex or age.	Moderate (for age and sex) Low (for other factors)
	Sustained virologic response	Two trials of triple therapy with pegylated interferon (alfa-2a or alfa-2b), ribavirin, and telaprevir vs. dual therapy depending reported inconsistent findings for differential relative risk estimates according baseline viral load.	Insufficient
	Dual Therapy With Pegylated Interferon Alfa-2b Plus Ribavirin vs. Dual Therapy With Pegylated Interferon Alfa-2a Plus Ribavirin		
Key Question 3a What are the comparative harms associated with antiviral treatments?	Harms	Dual therapy with pegylated interferon alfa-2b was associated with slightly greater risk of headache (three trials, pooled RR 1.1, 95% CI 1.1 to 1.2, I ² =0%), and a lower risk of serious adverse events (two trials, pooled RR 0.76; 95% CI 0.71 to 0.88; I ² =0%), lower risk of neutropenia (five trials, pooled RR 0.61, 95% CI 0.46 to 0.83, I ² =38%), and lower risk of rash (two trials, pooled RR 0.79, 95% CI 0.71 to 0.88, I ² =0.0%) than dual therapy with pegylated interferon alfa-2a plus ribavirin, with no differences in withdrawals due to adverse events.	Moderate

Key Question	Outcome	Summary of Evidence	Strength of Evidence
	Triple Therapy With Pegylated Interferon Alfa-2b, Ribavirin, and Boceprevir vs. Dual Therapy With Pegylated Interferon Alfa-2b Plus Ribavirin		
	Harms	Triple therapy with boceprevir for 48 weeks (pegylated interferon alfa-2b plus ribavirin for 4 weeks followed by addition of boceprevir for 44 weeks) was associated with increased risk of neutropenia (two trials, pooled RR 1.8, 95% CI 1.5 to 2.3, I²=0.0%), dysgeusia (two trials, pooled RR 2.5, 95% CI 2.0 to 3.2, I²=0.0%), anemia (two trials, pooled RR 2.0, 95% CI 1.4 to 2.8, I²=0.0%), and thrombocytopenia (two trials, pooled RR 3.2, 95% CI 1.2 to 8.2; I²=0.0%) than dual therapy with pegylated interferon alfa-2b plus ribavirin. The incidence of anemia was about 25% with triple therapy and the incidence of neutropenia about 33%, with severe anemia in 4–5% and severe neutropenia in 8–15%.	Moderate
Key Question 3a	Triple Therapy With Pegylated Interferon Alfa-2a or Alfa-2b, Ribavirin, and Telaprevir vs. Dual Therapy With Pegylated Interferon Alfa-2a or Alfa-2b Plus Ribavirin		
What are the comparative harms associated with antiviral treatments? (continued)	Harms	In two trials, there were no statistically significant differences between a 12-week regimen of triple therapy with pegylated interferon alfa-2a, ribavirin, and telaprevir vs. dual therapy with pegylated interferon alfa-2a plus ribavirin in risk of any assessed adverse event.	Moderate
	Harms	In three trials, a 24-week regimen of triple therapy with telaprevir (pegylated interferon alfa-2a or alfa-2b, ribavirin, and telaprevir for 12 weeks followed by pegylated interferon alfa-2a plus ribavirin for 12 weeks) was associated with increased risk of anemia (three trials, pooled RR 1.3, 95% CI 1.1 to 1.5, I²=0%) and rash (three trials, pooled RR 1.4, 95% CI 1.1 to 1.7; I²=0.0%) vs. dual therapy with pegylated interferon alfa-2a plus ribavirin for 48 weeks. Among patients randomized to the 24-week telaprevir regimen, one to two-thirds experienced a rash (7–10% experienced severe rash) and 27–91% experienced anemia (4–11% experienced severe anemia). There was no difference in risk of withdrawal due to adverse events.	Moderate

Key Question	Outcome	Summary of Evidence	Strength of Evidence
		With Pegylated Interferon Alfa-2a or Alfa-2b, Rib	
	and Telaprevir vs.	Dual Therapy With Pegylated Interferon Alfa-2a	or Alfa-2b
		Plus Ribavirin (continued)	T
Key Question 3a What are the comparative harms associated with antiviral treatments? (continued)	Harms	In one trial, response-guided triple therapy with telaprevir (pegylated interferon alfa-2a, ribavirin, and telaprevir for 8 or 12 weeks followed by response-guided duration pegylated interferon alfa-2a and ribavirin) was associated with increased risk of withdrawal due to adverse events (27% vs. 7.2%, RR 3.8, 95% CI 2.6 to 5.7), anemia (38% vs. 19%, RR 2.0, 95% CI 1.6 to 2.5), any rash (36% vs. 24%, RR 1.5, 95% CI 1.2 to 1.8), and severe rash (5% vs. 1%, RR 4.6, 95% CI 1.6 to 13) vs. therapy with pegylated interferon alfa-2a plus ribavirin for 48 weeks.	Low
	Dual Therapy With Pegylated Interferon Alfa-2b Plus Ribavirin vs. Dual Therapy With Pegylated Interferon Alfa-2a Plus Ribavirin		
Key Question 3b Do these harms differ according to patient subgroup	Harms	No trial of dual therapy with pegylated interferon alfa-2b plus ribavirin vs. dual therapy with pegylated interferon alfa-2a plus ribavirin reported harms in patients stratified by factors such as HCV genotype, age, race, sex, stage of disease, or genetic markers. Three trials that restricted enrollment to patients with genotype 1 infection reported risk estimates for risk of harms that were similar to the risk estimates based on all trials.	Insufficient
characteristics?	Triple Therapy With Pegylated Interferon Alfa-2a or Alfa-2b, Ribavirin,		
	and Telaprevir or	Boceprevir vs. Dual Therapy With Pegylated Int	terteron
	Harms	Alfa-2a or Alfa-2b Plus Ribavirin No trial evaluated harms associated with triple therapy with pegylated interferon, ribavirin, and boceprevir or telaprevir vs. dual therapy with pegylated interferon plus ribavirin in patient subgroups. All trials evaluated patients with genotype 1 infection.	Insufficient

Key Question	Outcome	Summary of Evidence	Strength of Evidence
Key Question 4 Have improvements in intermediate outcomes been shown to reduce the risk or rates of adverse health outcomes from HCV infection?	Mortality and long-term hepatic complications	A large VA hospital study that controlled well for potential confounders found an SVR after antiviral therapy associated with lower risk of all-cause mortality vs. no SVR (adjusted HR 0.71 [0.60-0.86], 0.62 [0.44-0.87] and 0.51 [0.35-0.75] for genotypes 1, 2, and 3, respectively). Eighteen other cohort studies found an SVR associated with decreased risk of all-cause mortality, liver-related mortality, HCC, and other complications of ESLD compared with no SVR, with stronger effect estimates than the VA study (adjusted HRs generally ranged from around 0.10 to 0.33). However, the studies had methodological shortcomings, including inadequate handling of confounders, and 10 were conducted in Asia.	Moderate
	Short-term quality of life	Nine studies found an SVR associated with greater improvement in measures related to quality of life (generic or disease-specific) 24 weeks after the end of antiviral treatment vs. no SVR, with differences averaging less than 5 to 10 points on various SF-36 domains. All studies were poor-quality and were characterized by failure to adjust for confounders, high loss to followup, and failure to blind patients to SVR status.	Low

bid = twice daily; CI = confidence interval; ESLD = end-stage liver disease; HCC = hepatocellular carcinoma; HCV = hepatitis C virus; HCV-RNA = hepatitis C virus ribonucleic acid; HR = hazard ratio; IU = international units; kg = kilograms; mcg = micrograms; mL = milliliters; RR = relative risk; SF-36=Short Form (36) Health Survey; SVR = sustained virologic response; tid = three times daily; VA = U.S. Department of Veterans Affairs

Findings in Relationship to What Is Already Known

Our findings regarding the comparative effectiveness of dual therapy with pegylated interferon alfa-2b plus ribavirin compared with dual therapy with pegylated interferon alfa-2a plus ribavirin are consistent with recent systematic reviews that also found the former associated with a lower likelihood of SVR. ^{14,79} Our findings of no clear difference in comparative effectiveness between 12 to 16 weeks compared with 24 weeks of response-guided dual therapy with pegylated interferon plus ribavirin in hepatitis C genotype 2 or 3 infection with rapid virologic response are discordant with a recent systematic review, which found a shorter duration of treatment associated with a lower likelihood of achieving an SVR. ⁸⁰ The discrepancy may be explained by the inclusion in the other systematic review of a study that we excluded because it evaluated a nonstandard dose of pegylated interferon, ⁸¹ as well as its inclusion of subgroup analyses from trials of patients randomized to different fixed durations of therapy prior to assessment of rapid virologic response, ^{40, 42, 43} which we considered separately because they did not represent randomized comparisons of response-guided treatment.

Because telaprevir and boceprevir are so new, we are unaware of other published systematic reviews on the comparative benefits and harms of regimens including these drugs, compared with standard dual therapy. Our findings on the association between achieving an SVR and

^a "Current antiviral treatment regimen" refers to dual therapy with pegylated interferon (alfa-2a or alfa-2b) plus ribavirin, or triple therapy with pegylated interferon (alfa-2a or alfa-2b) plus ribavirin and boceprevir or telaprevir.

reduced risk of mortality or complications associated with chronic HCV infection are consistent with a recent review that used some systematic methods.⁸²

Applicability

The trials included in this review generally met criteria for efficacy studies based on the exclusion of patients with common comorbidities (such as serious psychiatric conditions or recent or ongoing substance abuse). In addition, the trials may have overestimated efficacy compared with what would be seen in typical practice due to improved adherence as a result of closer followup, effects of trial participation, selection of patients, or other factors. A separate review funded by AHRQ will be focusing on issues related to the screening for HCV infection in adults. ²⁹

The severity of baseline liver disease in the patients enrolled in the trials suggests a broad range of patients were enrolled. In trials of triple therapy with boceprevir or telaprevir, the proportion of patients with cirrhosis at enrollment ranged from <1 to 11 percent. ^{26-28, 53, 54, 56, 57} Trials that reported the proportion of patients with minimal or no fibrosis reported rates of 27–39 percent. ^{27, 53, 54, 56, 57}

Evidence to evaluate potential differences in comparative benefits or harms in patient subgroups based on age, sex, race, and other clinical factors was relatively limited, precluding strong conclusions in these specific subgroups. The strongest evidence on the association between an SVR versus no SVR after antiviral therapy and reduced mortality comes from a study performed in a VA population, which might limit generalizability to other settings. As described above, studies conducted in Asia on the association between an SVR after antiviral therapy and risk of clinical outcomes may be of limited applicability to U.S. populations because of a higher incidence of HCC in Asian patients with chronic HCV infection. However, the incidence of HCC is increasing in the United States in HCV-infected people, which may attenuate such concerns regarding applicability.

The results of this CER are not applicable to populations excluded from the review, including patients previously treated with antiviral therapies and excluded populations such as patients with Human immunodeficiency virus (HIV) coinfection, post-transplant patients, or hemodialysis patients. Antiviral therapy is not recommended in patients following kidney transplant, and ribavirin is not recommended in those with more severe (stage 3 to 5) kidney disease since it is cleared via renal function and associated with increased risk of hemolytic anemia in this setting. Such patients were typically excluded from randomized trials of antiviral treatment.

Implications for Clinical and Policy Decisionmaking

Our review has potential implications for clinical and policy decisionmaking. For patients with genotype 1 infection, triple therapy regimens with pegylated interferon alfa-2a or alfa-2b, ribavirin, and telaprevir or boceprevir may be considered an alternative to dual therapy with pegylated interferon alfa-2a or alfa-2b plus ribavirin as standard treatment due to substantially superior efficacy for achieving SVR compared with dual therapy with pegylated interferon alfa-2a or alfa-2b, as well as a shorter duration of treatment. Factors that may affect decisions to use regimens with boceprevir or telaprevir include cost and specific harms associated with use of these drugs (such as hematologic adverse events with boceprevir and anemia and rash with telaprevir). Dual therapy with pegylated interferon alfa-2a plus ribavirin appears to be associated with a higher likelihood of achieving SVR compared with dual therapy with pegylated interferon

alfa-2b plus ribavirin, but absolute differences were relatively small. Therefore, decisions about which pegylated interferon to use may be affected by other considerations, such as cost, patient preferences, or other factors. For genotype 2 or 3 infection, standard doses and duration (24 weeks) of pegylated interferon as part of dual therapy are more effective than shorter regimens or lower doses, lending support to dosing guidance from the FDA and clinical practice guidelines. ^{11, 85, 86} Evidence on differential effects of ribavirin dose are too limited to draw strong conclusions about optimal dosing of this component of antiviral regimens, although differences appeared relatively small.

The findings that absolute SVR rates are lower in certain subgroups (such as older patients, Black patients, patients with worse baseline fibrosis, and patients with high viral load) can be used to guide individualized decisionmaking. Patients who are less likely to achieve an SVR may make different informed decisions about therapy compared with those more likely to achieve an SVR, given the adverse effects associated with treatment.

The findings of the review are also relevant to screening recommendations, which are based in part on the effectiveness of treatments in people found through screening to have HCV infection. Important new evidence that may affect assessments regarding potential benefits of screening include stronger evidence on the link between achieving an SVR and improvement in clinical outcomes, as well as evidence showing substantially higher SVR rates with newer triple therapy regimens with boceprevir or telaprevir in patients with genotype 1 infection, the predominant type of HCV infection in the United States.

Limitations of the Comparative Effectiveness Review Process

Our review had some potential limitations. We excluded non–English-language articles, which could result in language bias, although a recent systematic review found little empirical evidence that exclusion of non–English-language articles leads to biased estimates for noncomplementary or alternative medicine interventions.⁸⁷

We did not formally assess for publication bias with funnel plots due to small numbers (<10) of studies for all comparisons. Small numbers of studies can make interpretation of funnel plots unreliable, and experts suggest 10 studies as the minimum number of studies to perform them. 88 We included some studies that were published only as abstracts and found their inclusion or exclusion from analyses did not change conclusions. In addition, we searched trial registries and solicited drug manufacturers for additional unpublished trials and identified none.

Another potential limitation is that we included cohort studies to evaluate the association between SVR and either mortality or hepatic complications associated with chronic HCV infection. Such studies are susceptible to confounding if factors associated with SVR (such as age, race, viral load, or fibrosis stage) are also associated with these outcomes. Therefore, we only included studies that reported adjusted risk estimates, and we evaluated how well studies addressed key potential confounders as part of our quality assessment. Nonetheless, residual confounding is a possibility, even in cohort studies that adjust for potential confounding.

Limitations of the Evidence Base

We identified several important limitations of the evidence base. First, studies assessing important long-term clinical outcomes associated with current antiviral treatments for chronic HCV infection are not available. In the case of antiviral regimens involving newly approved antiviral drugs, such studies are not possible yet because of the extended followup required to adequately evaluate effects on clinical outcomes. Second, no trials directly compared regimens

with boceprevir with regimens with telaprevir. Given the increased efficacy of these regimens for genotype 1 infection, trials directly comparing their effects would be helpful for guiding health care providers' treatment choices between these drugs. Third, few trials have evaluated the regimens approved specifically by the FDA for these drugs, limiting confidence in conclusions regarding estimates of benefits and harms for the regimens likely to be used in clinical practice. Fourth, few methodologically rigorous studies conducted in settings applicable to U.S. populations evaluated the association between achieving an SVR and improvements in clinical outcomes. Such studies would be very helpful for confirming the results of the recent large, well-conducted VA cohort study showing an association between achieving an SVR and reduced mortality risk.⁸

Future Research

Evaluating the comparative effectiveness of current antiviral regimens on clinical outcomes in randomized trials or cohort studies is a challenge due to the long lead time and large sample sizes necessary to adequately assess these outcomes. This might be more feasible if the studies were to focus on populations at higher risk for complications from chronic HCV infection (e.g., patients with baseline cirrhosis, high viral load, or other risk factors for progression).

For all trials of antiviral treatments, studies that enroll broader populations with medical and psychological comorbidities, as frequently encountered in clinical practice, are needed to better understand comparative effectiveness, rather than just comparative efficacy. Studies designed using an effectiveness paradigm would also be helpful for understanding real-world outcomes of antiviral regimens, including effects related to the poorer treatment adherence than expected from efficacy trials.

Trials directly comparing triple therapy with telaprevir compared with triple therapy with boceprevir would be very helpful for understanding comparative effectiveness of these two protease inhibitors. In addition, trials evaluating the boceprevir regimen recommended by the FDA in antiviral-naïve patients without baseline cirrhosis are needed to verify that results from studies of previously treated patients were appropriately generalized. Prolonged followup of patients exposed to telaprevir and boceprevir is needed to understand the long-term harms associated with these medications. A number of other protease inhibitors and other newer drugs for treatment of hepatitis C virus infection are currently in active development, and further studies with new drugs and drug regimens are expected, including regimens without interferon. ⁸⁹

It is critical that future studies that evaluate clinical outcomes in patients with an SVR versus no SVR after antiviral therapy adequately control for other factors that influence clinical outcomes in chronic HCV infection. Studies on effects of achieving an SVR on long-term quality of life would be very helpful for understanding other potential clinical benefits of antiviral therapy, but a significant challenge is whether it is possible to ethically blind patients to virologic status, which may have an important effect on assessments of quality of life.

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Introduction

Hepatitis C virus (HCV) is the most common chronic bloodborne pathogen in the United States. HCV is primarily acquired by large or repeated percutaneous exposures to blood, with injection drug use the strongest risk factor. Based on a national survey of households, approximately 1.6 percent of U.S. adults over 20 years of age have antibodies to HCV, indicating prior acute HCV infection. About 78 percent of patients with acute HCV infection develop chronic HCV infection, defined by the presence of persistent viremia.

Chronic HCV infection has a variable course, but it is a leading cause of complications from chronic liver disease, including cirrhosis, liver failure, and hepatocellular carcinoma (HCC). Chronic HCV infection was associated with an estimated 15,000 deaths in the United States in 2007,² and it is the most common indication for liver transplantation among American adults, accounting for more than 30 percent of cases.³ The prevalence of chronic HCV infection is thought to have peaked in 2001 at 3.6 million people, and the yearly incidence has declined from more than 200,000 cases per year in the 1980s to around 16,000 cases in 2009.^{4,5} However, complications related to chronic HCV infection, which frequently occur only after decades of infection, are expected to rise for another 10 to 13 years.⁴

The goals of antiviral treatment for chronic HCV infection are to prevent the long-term health complications associated with HCV infection, such as cirrhosis, hepatic decompensation, and liver cancer, but it is a challenge to design and carry out clinical trials long and large enough to provide direct evidence related to these outcomes. The sustained virologic response (SVR) rate, typically defined as a decline in HCV-RNA (Hepatitis C virus ribonucleic acid) to undetectable levels 24 weeks following completion of antiviral treatment, is the standard marker for successful treatment in clinical trials because it is strongly associated with long-term absence of viremia. Recent studies have evaluated the association between achieving an SVR and reductions in mortality, liver failure, and cancer. Recent studies have evaluated the association between achieving an SVR and

The treatment of HCV infection has evolved dramatically over the past several decades. Recombinant type I interferons were introduced as monotherapy in the mid-1980s, but were only modestly successful at achieving SVR (overall <20 percent). Subsequent trials found dual therapy with interferon and the synthetic nucleoside analogue ribavirin more effective than monotherapy with interferon, although the SVR rates remained under 50 percent. $^{10-13}$

In the early 2000s, the combination of "pegylated" interferon plus ribavirin became the standard antiviral treatment for HCV infection. He first pegylated interferon was approved by the FDA in 2001. Pegylation refers to the cross-linking of polyethylene glycol molecules to the interferon molecule, which delays renal clearance and thereby permits less frequent dosing (once weekly vs. three times a week with nonpegylated interferon). Currently, two pegylated interferons are available: pegylated interferon alfa-2a and pegylated interferon alfa-2b. Both are Type I alfa interferons, but differ in the size and structure of the interferon and polyethylene glycol molecules, as well as in their pharmacokinetic properties (Table 1). One pegylated interferon consists of 31-kilodalton (kDa) interferon alfa-2b conjugated to 12-kDa polyethylene glycol (brand name PEG-intron). The other consists of recombinant 20-kDa interferon alfa-2a linked to 40-kDa polyethylene glycol (trade name Pegasys). The dosing schedule is fixed for pegylated interferon alfa-2a and is based on weight for pegylated interferon alfa-2b. Each pegylated interferon is approved for dual therapy with ribavirin. Although each pegylated interferon is approved for combination therapy with a specific brand of ribavirin manufactured by the respective manufacturer (Copegus) for pegylated interferon alfa-2a and Rebetol for alfa-2b), the ribavirin is pharmacologically identical. The FDA-recommended doses of ribavirin are

800 to 1200 mg/day for pegylated interferon alfa-2a, depending on weight and genotype, and 800 to 1400 mg/day for pegylated interferon alfa-2b, depending on weight.

Dual therapy with pegylated interferon (alfa-2a or alfa-2b) plus ribavirin is associated with higher SVR rates (about 55–60 percent overall) than either nonpegylated interferon plus ribavirin or pegylated interferon (alfa-2a or alfa-2b) monotherapy. Although previous reviews found insufficient evidence to determine whether dual therapy with pegylated interferon alfa-2a or pegylated interferon alfa-2b is more effective, ^{18, 19} more head-to-head trials directly comparing these two regimens are now available. ²⁰⁻²³

A number of factors affect response to antiviral treatment. The two major pretreatment predictors of SVR are the viral genotype and the pretreatment viral load. In the United States, genotype 1 infection is found in around three-quarters of HCV-infected patients. HCV genotype 1 infection is associated with a substantially lower response to antiviral treatment than infection with genotypes 2 and 3, which are present in about 20 percent of HCV-infected patients. A pretreatment viral load of <600,000 international units per milliliter (IU/mL) is associated with higher likelihood of achieving an SVR. Other factors less consistently or less strongly associated with increased likelihood of SVR include female sex, age less than 40 years, non-Black race, lower body weight (\leq 75 kg), absence of insulin resistance, elevated alanine aminotransferase (ALT) levels, and absence of bridging fibrosis or cirrhosis on liver biopsy. Effects of race on the likelihood of SVR may be due in part to polymorphisms in the interleukin-28B (IL28B) gene. Strong pretreatment viral load. The United States, and absence of bridging fibrosis or cirrhosis on liver biopsy.

An issue complicating antiviral treatment is the high rate of adverse effects observed with interferon-based therapy, including flulike symptoms, fatigue, and neuropsychiatric and hematologic adverse effects.²⁷ Such adverse effects can be difficult to tolerate and can lead to premature discontinuation of therapy.

In 2011, the U.S. Food and Drug Administration (FDA) approved the first direct acting antiviral agents, boceprevir (trade name Victrelis®) and telaprevir (trade name Incivek®), for treatment of chronic HCV genotype 1 infection (Table 1). Both drugs are classified as nonstructural (NS) 3/4A protease inhibitors, with a potential advantage of shorter duration of therapy (24 to 28 weeks) when used in combination with pegylated interferon (alfa-2a or alfa-2b) compared with standard dual therapy with pegylated interferon (alfa-2a plus -2b) plus ribavirin for genotype 1 infection (48 weeks) (Table 1). October 10.

Table 1. Pharmacokinetics, indications, and dosing of included drugs^{28, 29, 33, 34}

Drug	Indications Labeled by the U.S. Food and	Dosing Recommended by the U.S. Food and
Trade Name	Drug Administration	Drug Administration
Pegylated interferon alfa-2a Pegasys [®]	Patients 5 years of age and older with chronic HCV infection with compensated liver disease not previously treated with interferon alfa	180 mcg once weekly in combination with ribavirin for 24 weeks with ribavirin for genotypes 2 or 3, or 48 weeks for genotype 1or 4 infection
Pegylated interferon alfa-2b PEG-Intron®	Patients 5 years of age and older with chronic HCV infection with compensated liver disease	1.5 mcg/kg weekly in combination with ribavirin for 24 weeks with ribavirin for genotypes 2 or 3, or 48 weeks for genotype 1 infection
Boceprevir Victrelis [®]	Adults with chronic HCV genotype 2 infection with compensated liver disease, including cirrhosis, who are previously untreated or who have been previously treated with interferon and ribavirin therapy	Four weeks of treatment with pegylated interferon (alfa-2a or 2b) plus ribavirin, then the addition of boceprevir 800 mg 3 times daily as follows: ^a In treatment-naïve patients without cirrhosis: - If HCV-RNA undetectable from treatment week 8 through week 24, complete triple therapy at treatment week 28 - If HCV-RNA detectable at treatment week 8 and undetectable at treatment week 24, continue triple therapy through treatment week 36 and continue pegylated interferon (alfa-2a or 2b) with ribavirin through treatment week 48
		In treatment-naïve patients with cirrhosis: - 44 weeks of triple therapy
Telaprevir Incivek [®]	Adults with chronic HCV genotype 1 infection with compensated liver disease, including cirrhosis, who are previously untreated or who have been previously treated with interferon and ribavirin therapy	750 mg 3 times a day with pegylated interferon (alfa-2a or 2b) and ribavirin for all patients for 12 weeks, followed by response-guided regimen of pegylated interferon and ribavirin ^a In treatment-naïve patients without cirrhosis:
		If HCV-RNA is undetectable at weeks 4 and 12, then continue dual therapy for 12 more weeks (total treatment 24 weeks) If HCV-RNA is detectable at week 4 and/or week 12, then continue dual therapy for 36 more weeks (total treatment 48 weeks)
	C virus: HCV PNA – hanatitis C virus ribonuclaic a	In treatment-naïve with cirrhosis: - Continue dual therapy for 36 more weeks (total treatment 48 weeks)

HCV = hepatitis C virus; HCV-RNA = hepatitis C virus ribonucleic acid

Understanding the comparative benefits and harms of the various antiviral regimens is critical for making informed treatment decisions in patients with chronic HCV infection, particularly given the availability of new treatment options. This review will assess the comparative effectiveness of antiviral treatments in adults with chronic HCV infection who have not received previous antiviral drug treatment. In addition to assessing the comparative effectiveness of different drug regimens, the review will evaluate effects of different medication doses, durations of therapy, and dosing strategies (such as weight-based or response-guided vs. fixed treatment). To help with individualized clinical decisionmaking regarding antiviral therapy for chronic HCV infection, it will also evaluate how comparative effectiveness varies depending

^a The manufacturer packaging and dosage information does not specify a particular pegylated interferon (alfa-2a or alfa-2b) for either drug, though in trials conducted to obtain FDA approval, boceprevir was tested with pegylated interferon alfa-2b and telaprevir with pegylated interferon alfa-2a.

on HCV genotype, viral load, and other demographic and clinical characteristics. Because estimating potential benefits and harms of HCV screening requires an understanding of the effects of treatment in people with HCV infection, this review will be used, together with a separate review on HCV screening,³⁵ by the U.S. Preventive Services Task Force to update its HCV screening recommendations.

Scope and Key Questions

The analytic framework and Key Questions used to guide this report are shown below (Figure 1). The analytic framework shows the target populations, interventions, and intermediate and health outcome measures we examined.

The following Key Questions are the focus of our report:

Key Question 1

- a. What is the comparative effectiveness of antiviral treatment in improving health outcomes in patients with HCV infection?
- b. How does the comparative effectiveness of antiviral treatment for health outcomes vary according to patient subgroup characteristics, including but not limited to HCV genotype, age, race, sex, stage of disease, or genetic markers?

Key Question 2

- a. What is the comparative effectiveness of antiviral treatments on intermediate outcomes, such as the rate of SVR or histologic changes in the liver?
- b. How does the comparative effectiveness of antiviral treatment for intermediate outcomes vary according to patient subgroup characteristics, including but not limited to HCV genotype, age, race, sex, stage of disease, or genetic markers?

Key Question 3

- a. What are the comparative harms associated with antiviral treatments?
- b. Do these harms differ according to patient subgroup characteristics, including HCV genotype, age, race, sex, stage of disease, or genetic markers?

Key Question 4

Have improvements in intermediate outcomes (SVR, histologic changes) been shown to reduce the risk or rates of adverse health outcomes from HCV infection?

Key Question 1 focuses on direct evidence on the comparative effectiveness of antiviral treatments for chronic HCV infection on health outcomes (such as death, cirrhosis, hepatic decompensation, HCC, need for transplantation, or quality of life). Because of the long duration (typically decades) necessary develop major hepatic complications related to chronic HCV infection, it is difficult to assess for such outcomes in clinical trials. In addition, dual therapy with pegylated interferon (alfa-2a or alfa-2b) plus ribavirin has only been available since 2001, and protease inhibitors only became approved by the FDA in 2011, which might not be enough time to adequately evaluate some long-term clinical outcomes. Therefore, Key Question 2 focuses on evidence on the comparative effectiveness of antiviral treatments for chronic HCV infection on intermediate outcomes (SVR and histological improvements). Key Question 4 assesses the link between intermediate and clinical outcomes, in order to facilitate interpretation

of results obtained for Key Question 2. Key Question 3 focuses on the comparative harms of different antiviral treatments.

KQ. 1 Final Clinincal Outcomes Intermediate Antiviral Outcomes Treatment Patients with KQ. 4 KQ. 2 Mortality Sustained chronic Morbidity virologic Hepatitis C Virus • Quality of life response infection Transmission • Histological of Hepatitis C improvements Virus KQ. 3 Harms

Figure 1. Analytic framework for treatment of hepatitis C infection in adults

KQ = Key Question

Methods

Input From Stakeholders

The topic of hepatitis C virus (HCV) treatment was nominated for a comparative effectiveness review (CER) in a public process. The Key Questions were proposed in the public nomination process and developed by investigators from the Evidence-based Practice Center (EPC) with input from expert Key Informants (KI), who helped to refine Key Questions, identify important methodological and clinical issues, and define parameters for the review of evidence. The revised Key Questions were then posted to a public Web site for comment. The Agency for Healthcare Research and Quality (AHRQ) and the EPC agreed upon the final Key Questions after reviewing the public comments and receiving additional input from a Technical Expert Panel (TEP) convened for this report. We then drafted a protocol for this CER, which was reviewed by the TEP and is available on the AHRQ Web site where it was posted in November 2011: www.effectivehealthcare.ahrq.gov/index.cfm/search-for-guides-reviews-and-reports/?pageaction=displayproduct&productid=855.

A multidisciplinary group of clinicians, researchers, and patient advocates with expertise in hepatitis C treatment and research were selected to serve as the TEP members to provide high-level content and methodological expertise throughout the development of the review. Prior to participation in this report, the TEP members disclosed all financial or other conflicts of interest. The AHRQ Task Order Officer and the authors reviewed all of these disclosures and determined the panel members had no significant conflicts of interest that precluded participation. KIs and TEP members had expertise in the areas of hepatology, epidemiology, screening, and primary care. TEP members and other experts were invited to provide external peer review of the draft report.

Search Strategy

To identify articles relevant to each Key Question, a research librarian searched the Cochrane Database of Systematic Reviews, the Cochrane Central Register of Controlled Trials, and Ovid MEDLINE® (Appendix A) from 1947 to April 2011 (see Appendix A for the search strategies and a final updated search was conducted in August 2012 following the receipt of peer reviewer comments. The search strategies were peer reviewed by another research librarian and revised prior to finalization. Unpublished trials were sought by searching clinical trial registries (ClinicalTrials.gov, Current Controlled Trials, Clinical Trial Results, WHO Trial Registries) and grants databases (NIHRePORTER, HSRProj, and AHRQ GOLD). Scientific Information Packets on unpublished and published trials were solicited from manufacturers of included antiviral drugs through the Scientific Resource Center. We also hand-searched the reference lists of relevant studies. Searches were updated prior to finalization of the report to identify relevant new publications.

Study Selection

We developed criteria for inclusion and exclusion of studies based on the Key Questions and the populations, interventions, comparators, outcomes, timing, and setting (PICOTS) approach. Inclusion and exclusion criteria, summarized below, are described in more detail by Key Question in Appendix B. Papers were selected for full review if they were about chronic HCV infection, were relevant to Key Questions in the analytic framework, and met the predefined

inclusion criteria. To evaluate potential effects of publication bias, we included trials published only as conference abstracts as sensitivity analyses. We restricted inclusion to English language articles. Studies of nonhuman subjects were also excluded, and studies had to include original data.

Abstracts and full-text articles were dual reviewed for inclusion and exclusion for each Key Question (Appendix B). Full-text articles were obtained for all studies that either investigator identified as potentially meeting inclusion criteria. Two investigators independently reviewed all full-text articles for final inclusion or exclusion (Appendix C). A list of excluded studies with primary reasons for exclusion can be found in Appendix D. Discrepancies were resolved through discussion and consensus, with a third investigator making the final decision if necessary.

Population and Conditions of Interest

The target population for Key Questions 1 through 3 was nonpregnant adults with chronic HCV infection who have not had previous antiviral drug treatment. Pregnant women were excluded as no antiviral treatment for HCV infection is currently recommended during pregnancy due to potential teratogenic effects. We also evaluated comparative benefits and harms in patient subgroups defined by HCV genotype, race, sex, stage or severity of disease, viral load, weight, genetic markers (i.e., polymorphisms in the IL28B gene), and other factors (such as body weight). For Key Question 4, the target population was adults with chronic HCV infection who had received a course of interferon-based antiviral therapy. We excluded post-transplant patients, HIV patients, and hemodialysis patients, because treatment considerations and response to therapy may differ from what is observed in the general population of patients with chronic HCV infection without these conditions.

Interventions and Comparisons

We included antiviral regimens recommended in current guidelines for treatment of HCV infection, specifically dual therapy with pegylated interferon alfa-2a or alfa-2b plus ribavirin for genotype 2 or 3 infection, ¹⁵ and triple therapy regimens with the recently approved protease inhibitors telaprevir and boceprevir, which are used in combination with pegylated interferon alfa-2a or alfa-2b plus ribavirin, for genotype 1 infection. ³⁷ We included studies of interferon monotherapy and standard interferon plus ribavirin only for Key Question 4, which evaluated the association between intermediate and clinical outcomes. We excluded regimens that involved antiviral drugs that are not approved in the United States for treatment of chronic HCV infection.

For Key Questions 1 through 3, we included studies that compared dual therapy with pegylated interferon alfa-2a plus ribavirin compared with dual therapy with pegylated interferon alfa-2b plus ribavirin, or that compared triple therapy with pegylated interferon (alfa-2a or alfa-2b), ribavirin, and a protease inhibitor (either telaprevir or boceprevir) compared with dual therapy with pegylated interferon (alfa-2a or alfa-2b) plus ribavirin. We also included studies that evaluated different doses or dosing protocols (i.e., weight-based vs. standardized) of the same antiviral drugs, or different durations of therapy or methods (e.g., response-guided therapy vs. fixed-duration therapy) for guiding duration of therapy. We focused on dose and duration comparisons of dual therapy with pegylated interferon (alfa-2a or alfa-2b) plus ribavirin in patients with HCV genotypes 2 and 3. For Key Question 4, we included studies of patients with chronic HCV infection who received antiviral treatment that compared outcomes between those who achieved an SVR (or improved histological findings) after antiviral therapy and those who did not.

Outcomes

Clinical outcomes were mortality, cirrhosis, hepatic decompensation, HCC, need for transplantation, and quality of life. We classified clinical outcomes assessed 1 year or earlier after the end of antiviral treatment as short-term and those assessed after at least 1 year as longterm. Intermediate outcomes were SVR rates and improvements in histological outcomes. We defined a sustained virologic response as the absence of detectable HCV-RNA in the serum six months after the end of a course of therapy. 15 We did not evaluate measures of earlier virologic response (such as undetectable HCV-RNA before or through week 12 of therapy or at the end of therapy). Although such early virologic outcomes predict whether a patient will achieve an SVR and can be used to guide therapy decisions (e.g., whether to continue therapy or duration of therapy), they are less accurate than the SVR for predicting long-term remission. ¹⁵ Histological response has been defined as a 2-point or greater decrease in the inflammatory score or fibrosis score, or a 1-point decrease in the fibrosis score, although relatively few trials evaluate histological response and definitions are less standardized compared with SVR. 15, 38 We did not evaluate improvement in liver function tests as an intermediate outcome (e.g., sustained biochemical response, or normalization of liver transaminases six months after the end of a course of therapy), due to its poor correlation with SVR. 39-42 Harms of treatment included withdrawals due to adverse events, serious adverse events such as neutropenia, anemia, psychological adverse events, flulike symptoms, and dermatologic adverse events.

Timing

We did not apply a minimum threshold for duration of studies. We defined long-term outcomes as those measured one year or more after the completion of antiviral therapy and short-term outcomes as those measured prior to one year after the completion of antiviral therapy.

Setting

Studies conducted in primary care and specialty settings were included.

Types of Studies

We included randomized trials for all Key Questions. For Key Question 4, we included cohort studies that compared clinical outcomes between patients who achieved an SVR compared with those who did not achieve an SVR, or that compared clinical outcome between patients who achieved a histological response compared with those who did not. Many factors (such as age, race, viral load, and fibrosis stage) may be associated with both the likelihood of achieving an SVR as well as the likelihood of hepatic complications. Therefore, we excluded studies on the association between achieving an SVR and mortality or hepatic complications that only reported unadjusted risk estimates, given the strong potential for confounding. Because almost no studies on the association between SVR and quality of life reported adjusted risk estimates, we included studies that reported unadjusted risk estimates for this association.

Data Extraction

We extracted the following data from included studies into Excel spreadsheets: study design, setting, population characteristics, eligibility and exclusion criteria, the antiviral regimen (including duration and dose), and results for each outcome. Data abstraction for each study was

completed by two investigators: the first abstracted the data, and the second reviewed the abstracted data for accuracy and completeness against the original articles.

For Key Question 4, some studies reported adjusted hazard ratios (HRs) for the association between achieving an SVR and clinical outcomes relative to untreated patients, and for no SVR and clinical outcomes relative to untreated patients, but did not report a risk estimate for SVR compared with no SVR. We calculated the HR for SVR compared with no SVR based on the two HRs and their reported confidence intervals, assuming zero correlation between the two reported HRs. Such HRs are usually positively correlated; an assumption of zero correlation results in the most conservative (widest) confidence interval for the HR for SVR compared with no SVR.

Assessing Quality

We assessed quality for each study based on the predefined criteria listed in Appendix E. We adapted criteria from methods proposed by Downs and Black⁴³ and the USPSTF.⁴⁴ The criteria used are consistent with the approach recommended in AHRQ's Methods Guide for Effectiveness and Comparative Effectiveness Reviews (Methods Guide).⁴⁵ We used the term "quality" rather than the alternate term "risk of bias." Although both refer to internal validity, "quality" may be more familiar to most users and has potential advantages in terms of readability.

We rated the quality of each randomized trial based on the methods used for randomization, allocation concealment, and blinding; the similarity of compared groups at baseline; maintenance of comparable groups; adequate reporting of dropouts, attrition, crossover, adherence, and contamination; loss to followup; the use of intent-to-treat analysis; and ascertainment of outcomes.⁴⁴

We rated the quality for each cohort study based on whether it used nonbiased selection methods to create an inception cohort; whether it evaluated comparable groups; whether rates of loss to followup were reported and acceptable; whether it used accurate methods for ascertaining exposures, potential confounders, and outcomes; and whether it performed appropriate statistical analyses of potential confounders. He for Key Question 4, we considered studies to have performed adequate statistical analyses of potential confounders if they adjusted at a minimum for age, sex, genotype, viral load, and hepatic fibrosis stage in a multivariate model including SVR or histological response; evaluated these factors and excluded them from the multivariate model because there was no association in either univariate or step-wise multivariate analyses; or accounted for these factors using other methods such as stratification or restriction.

Following assessment of individual quality criteria, individual studies were rated as good, fair, or poor quality, as defined below. 44, 45

Good quality studies are considered likely to be valid. Good quality studies clearly describe the population, setting, interventions, and comparison groups; use a valid method for allocating patients to interventions; clearly report dropouts and have low dropout rates; use appropriate methods for preventing bias; and appropriately measure outcomes and fully report results.

Fair quality studies have some methodological deficiencies, but no flaw or combination of flaws judged likely to cause major bias. The study may be missing information, making it difficult to assess its methods or assess limitations and potential problems. The fair quality category is broad, and studies with this rating vary in their strengths and weaknesses—the results of some fair quality studies are likely to be valid, while others are only probably valid.

Poor quality studies have significant flaws that may invalidate the results. They have a serious or fatal flaw in design, analysis, or reporting; large amounts of missing information; or

discrepancies in reporting. The results of these studies are judged to be at least as likely to reflect flaws in the study design as true effects of the interventions under investigation. We did not exclude poor quality studies a priori, but they were considered the least reliable studies when synthesizing the evidence, particularly when discrepancies between studies were present.

Assessing Research Applicability

We recorded factors important for understanding the applicability of studies such as whether the publication adequately described the study population, the country in which the study was conducted (studies indicate that the rate of HCC in patients with chronic HCV infection is higher in Japan and other Asian countries compared with the United States), ⁴⁶ how similar patients were to typical populations of those with chronic HCV infection, whether differences in outcomes were clinically (as well as statistically) significant, and whether the antiviral regimens and other aspects of care evaluated were reasonably representative of standard practice. ⁴⁷ We also recorded the funding source and role of the sponsor. We did not assign a rating of applicability (such as high or low) because applicability may differ based on the user of this report.

Data Synthesis

For Key Questions 1 through 3, we performed meta-analysis of trials that evaluated similar populations, interventions, comparisons, and outcomes to estimate pooled relative risks using the DerSimonian-Laird method in a random effects model.⁴⁸ A random effects model results in estimates that are similar to a fixed effects model when there is little or no between-study statistical heterogeneity, but results in more conservative estimates (wider confidence intervals) when statistical heterogeneity is present. Heterogeneity was assessed by calculating the Qstatistic and the percentage of the total variance due to between study variability (1² statistic).⁴⁹ When present, statistical heterogeneity was explored through subgroup and sensitivity analyses, as well as qualitatively. Subgroup analyses were performed in groups stratified by HCV genotype as well as by race, age, body weight, viral load, stage/severity of disease, and IL-28b status when these data were available. We performed sensitivity analysis by excluding poorquality studies, excluding outlier trials and including trials that used nonstandard doses of antiviral drugs, and adding results from trials published only as abstracts to evaluate the stability of estimates and conclusions. We did not formally assess for publication bias with funnel plots due to small numbers (<10) of studies for all comparisons. Small numbers of studies can make interpretation of funnel plots unreliable, and experts suggest 10 studies as the minimum number of studies to perform funnel plots.⁵⁰ All analyses were performed using Stata 11.0 (StataCorp. College Station, TX, 2009).

For Key Question 4, we did not perform meta-analysis, since all studies were cohort studies, and many had methodological shortcomings (including failure to adjust for important confounders) and varied in populations assessed, treatments received, and other factors. Rather, these studies were synthesized qualitatively.

Strength of the Body of Evidence

We assessed the overall strength of evidence for a body of literature about a particular Key Question in accordance with the AHRQ Methods Guide.⁴⁵ The strength of evidence was based on the overall quality of each body of evidence, based on the type and quality of studies (graded good, fair, or poor); the consistency of results within and between study designs (graded high,

moderate, or low); the directness of the evidence linking the intervention and health outcomes (graded direct or indirect); and the precision of the estimate of effect, based on the number and size of studies and confidence intervals for the estimates (graded high, moderate, or low). We did not downgrade a body of evidence for directness that evaluated an intermediate outcome, if the intermediate outcome was the specific focus of the Key Question. We did not grade supplemental domains for cohort studies included in Key Question 4 because they were not relevant (dose-response relationship) or because important methodological shortcomings (in particular failure to adjust for critical confounders) limited their usefulness (magnitude of effect and direction of plausible confounding). We were not able to formally assess for publication bias due to small numbers of studies, methodological shortcomings, or differences across studies in designs, measured outcomes, and other factors.

We graded the strength of evidence for each Key Question using the four key categories recommended in the AHRQ Methods Guide. ⁴⁵ A "high" grade indicates high confidence that the evidence reflects the true effect and that further research is very unlikely to change our confidence in the estimate of effect. A "moderate" grade indicates moderate confidence that the evidence reflects the true effect and further research may change our confidence in the estimate of effect and may change the estimate. A "low" grade indicates low confidence that the evidence reflects the true effect and further research is likely to change the confidence in the estimate of effect and is likely to change the estimate. An "insufficient" grade indicates evidence either is unavailable or is too limited to permit any conclusion.

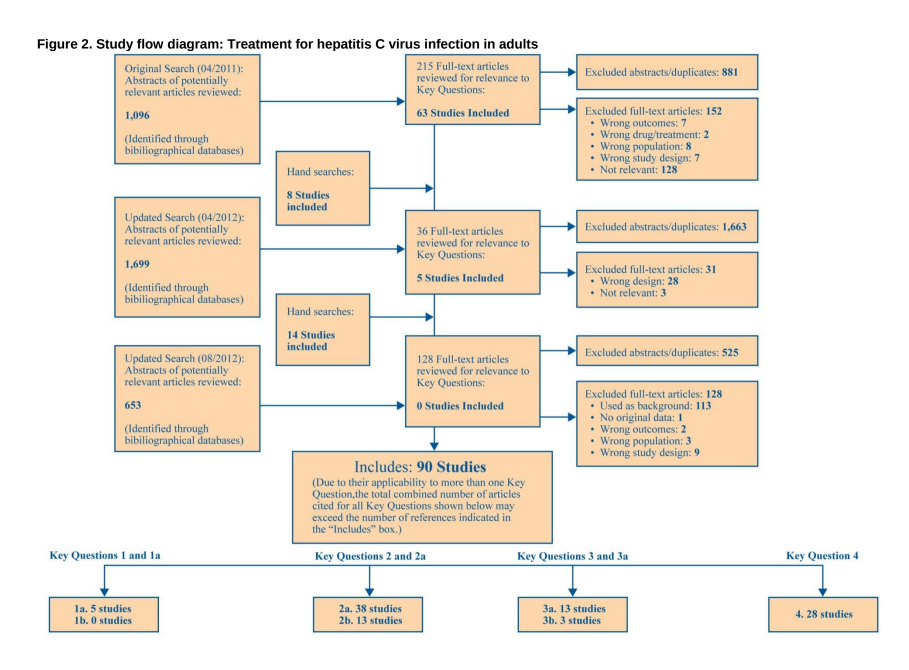
Peer Review and Public Commentary

Experts in gastroenterology, hepatology, primary care, and prevention, and individuals representing stakeholder and user communities were invited to provide external peer review of a draft of this CER; AHRQ and an EPC associate editor also provided comments. The draft report was posted on the AHRQ Web site for 4 weeks to elicit public comment. All comments were reviewed and addressed as documented in a disposition of comments report that will be made available 3 months after the Agency posts the final CER on the AHRQ Web site www.effectivehealthcare.ahrq.gov.

Results

Overview

The search and selection of articles are summarized in the study flow diagram (Figure B). Of the 1,096 citations identified at the title and abstract level in the original search, 215 articles met inclusion criteria and were selected for further review of the full text. From updated searches and peer reviewer suggested citations, an additional 2,352 citations were identified, and 164 of these met inclusion criteria and were selected for full-text review. Of the 379 articles reviewed at the full-text level, a total of 90 studies met inclusion criteria.



Key Question 1a. What is the comparative effectiveness of antiviral treatment in improving health outcomes in patients with HCV infection?

- No randomized trial or observational study evaluated the comparative effectiveness of current antiviral treatment regimens for chronic HCV infection on improving long-term clinical outcomes (strength of evidence: insufficient).
- Three trials that compared current antiviral regimens found no differences in risk of short-term mortality, but reported very few (20 total) events (strength of evidence: low).
- One open-label randomized trial of patients with genotype 4 infection found dual therapy with pegylated interferon alfa-2a plus ribavirin associated with statistically significant, slightly better short-term scores on some generic and liver disease-specific quality of life assessments than dual therapy with pegylated interferon alfa-2b plus ribavirin (strength of evidence: low).

No trial evaluated comparative effects of current antiviral treatment regimens for chronic HCV infection (dual therapy with pegylated interferon alfa-2a or alfa-2b plus ribavirin or triple therapy with pegylated interferon alfa-2a or alfa-2b, ribavirin, and a protease inhibitor) on risk of long-term clinical outcomes.

Three trials reported short-term mortality (through 6 months after the completion of antiviral therapy), but reported few deaths (20 total), resulting in very imprecise estimates (Appendix H: Evidence Table 1). One large trial found no difference between dual therapy with standard dose pegylated interferon alfa-2b (1.5 mcg/kg/week) plus ribavirin versus pegylated interferon alfa-2a plus ribavirin in risk of short-term mortality (risk ratio [RR] 0.85, 95% confidence interval (CI) 0.26 to 2.8), based on 11 deaths in over 2000 subjects. Another trial found no difference between triple therapy with boceprevir and dual therapy in risk of short-term all-cause mortality (RR 0.25, 95% CI 0.03 to 2.2), but only reported 5 deaths in over 700 patients. One trial of response-guided triple therapy with telaprevir versus dual therapy reported four deaths in over 1088 patients, resulting in a very imprecise estimate (RR 1.5, 95% CI 0.16 to 14).

Two trials evaluated comparative effects of current antiviral regimens for chronic HCV infection on short-term quality of life (Appendix H: Evidence Table 11). 52,53 One trial of patients with genotype 4 infection found dual therapy with pegylated interferon alfa-2a plus ribavirin associated with slightly higher (better) scores on some Short-Form 36 (SF-36) health survey subscales than dual therapy with pegylated interferon alfa-2b plus ribavirin 24 weeks after the end of treatment (differences of 3.2 to 5.7 points on the Bodily Pain, Vitality, Social Functioning, and Role Emotional subscales, each on a 0 to 100 scale).⁵³ Dual therapy with pegylated interferon alfa-2a was also associated with slightly higher scores on the Physical Component Summary score (3.2, points, p<0.02), but there was no difference on the Mental Component Summary score, or on five of six domains on the Chronic Liver Disease questionnaire, though dual therapy with pegylated interferon alfa-2a plus ribavirin was associated with a slightly higher overall score (difference 0.4 point on a 1 to 7 scale, p=0.02). The trial was open-label and patients do not appear to have been blinded to virologic response status, which could have affected quality of life assessments. A trial of patients with genotype 1 infection with undetectable HCV-RNA after 24 weeks of pegylated interferon alfa-2a plus ribavirin found continuation of dual therapy for another 24 weeks associated with worse quality of life scores at the end of treatment than pegylated interferon alone for the last 24 weeks, but the clinical relevance of this finding is limited since the shorter regimen was associated with lower

likelihood of achieving an SVR and is not considered the standard of care for genotype 1 infection.⁵²

Key Question 1b. How does the comparative effectiveness of antiviral treatment for health outcomes vary according to patient subgroup characteristics, including but not limited to HCV genotype, age, race, sex, stage of disease or genetic markers?

• No randomized trial or observational study evaluated comparative effects of current antiviral treatment regimens on any clinical outcomes in patients stratified by HCV genotype, age, race, sex, stage of disease, genetic markers, or other factors (strength of evidence: insufficient).

Key Question 2a. What is the comparative effectiveness of antiviral treatments on intermediate outcomes, such as the rate of SVR or histologic changes in the liver?

Dual Therapy With Pegylated Interferon Alfa-2a or Alfa-2b Plus Ribavirin

• Seven trials found dual therapy with standard doses of pegylated interferon alfa-2b plus ribavirin associated with lower likelihood of achieving an SVR than pegylated interferon alfa-2a plus ribavirin (pooled RR 0.87, 95% CI 0.80 to 0.95, I²=27.4%), with an absolute difference in SVR rates of 8 percentage points (95% CI 3 to 14) (strength of evidence: moderate).

Dual Therapy With Pegylated Interferon Alfa-2a or Alfa-2b Plus Ribavirin: Duration Effects

- Two trials of patients with genotype 2 or 3 infection found no difference in likelihood of achieving an SVR between 48 versus 24 weeks of dual therapy with pegylated interferon alfa-2a plus ribavirin (pooled RR 0.97, 95% CI 0.84 to 1.11, I²=42.7%) (strength of evidence: moderate).
- Four trials of patients with genotype 2 or 3 infection found 24 weeks of dual therapy with pegylated interferon (alfa-2a or alfa-2b) more effective than 12-16 weeks for achieving an SVR (pooled RR 1.15, 95% CI 1.02 to 1.29, I²=79.5%). Relative risk estimates ranged from 1.0 to 1.3 (strength of evidence: moderate).
- Three trials of patients with genotype 2 or 3 infection with a rapid virologic response (undetectable HCV-RNA by week 4) found no differences between 24 versus 12-16 weeks of dual therapy with pegylated interferon (alfa-2a or alfa-2b) plus ribavirin (pooled RR 0.99, 95% CI 0.86 to 1.14, I²=66.7%). Relative risk estimates ranged from 0.89 to 1.1 (strength of evidence: moderate).

Dual Therapy With Pegylated Interferon Alfa-2a or Alfa-2b Plus Ribavirin: Dose Effects

- Six trials of patients with genotype 2 or 3 infection found lower doses of pegylated interferon alfa-2b (0.75–1.0 mcg/kg or 50 mcg) associated with lower likelihood of achieving an SVR than higher doses (1.5 mcg/kg or 100–150 mcg) (pooled RR 0.90, 95% CI 0.81 to 0.99, I²=20.2%) (strength of evidence: moderate).
- Three trials of patients with genotype 2 or 3 infection who did not specifically have advanced fibrosis or cirrhosis found no clear difference in likelihood of SVR between lower doses of ribavirin (400 or 800 mg flat dose or 600 to 800 mg weight-based dose) versus higher doses (800 or 1200 mg flat dose or 800 to 1400 mg weight-based dose) (strength of evidence: moderate).
- One small trial of patients with genotype 2 or 3 infection (N=60) and advanced fibrosis or cirrhosis (Ishak stage 4-6) found 600 to 800 mg daily of ribavirin associated with lower likelihood of SVR than 1000 to 1200 mg daily (45 vs. 72 percent, RR 0.62, 95% C I 0.40 to 0.98) (strength of evidence: low).

Trials of Triple Therapy With Pegylated Interferon Alfa-2b, Ribavirin, and Boceprevir

- Two trials of patients with HCV genotype 1 infection found dual therapy lead-in with pegylated interferon alfa-2b plus ribavirin for 4 weeks followed by 44 weeks of triple therapy with boceprevir associated with higher likelihood of SVR than dual therapy for 48 weeks (pooled RR 1.81, 95% CI 1.58 to 2.06, I²=0.0%), with an absolute increase in SVR rate of 31 percentage points (95% CI 23 to 39) (strength of evidence: moderate).
- One trial of patients with genotype 1 infection found 48 weeks of triple therapy with boceprevir using low dose of ribavirin (400–1000 mg daily) associated with a non-statistically significant trend towards lower likelihood of SVR than 48 weeks of triple therapy with a standard ribavirin dose (800–1400 mg daily) (36 vs. 50 percent, RR 0.71, 95% CI 0.39 to 1.3) (strength of evidence: low).

Trials of Triple Therapy With Pegylated Interferon Alfa-2a or Alfa-2b, Ribavirin, and Telaprevir

- Three trials of patients with genotype 1 infection found triple therapy with telaprevir for 24 weeks (12 weeks of pegylated interferon alfa-2a or alfa-2b, ribavirin, and telaprevir followed by 12 weeks of pegylated interferon alfa-2a or alfa-2b plus ribavirin) associated with higher likelihood of SVR than dual therapy with pegylated interferon alfa-2a or alfa-2b plus ribavirin for 48 weeks (pooled RR 1.48, 95% CI 1.26 to 1.75, I²=0.0%), with an absolute increase in SVR rate of 22 percentage points (95% CI 13 to 31) (strength of evidence: moderate).
- One trial of patients with genotype 1 infection found no difference in likelihood of SVR between triple therapy with pegylated interferon alfa-2a, ribavirin, and telaprevir for 12 weeks versus dual therapy with pegylated interferon alfa-2a plus ribavirin for 48 weeks (strength of evidence: moderate).
- One trial of patients with genotype 1 infection found response-guided triple therapy with telaprevir (pegylated interferon alfa-2a, ribavirin, and telaprevir for 8 or 12 weeks

followed by response-guided dual therapy with pegylated interferon alfa-2a plus ribavirin for an additional 12 or 36 weeks) associated with higher likelihood of SVR than dual therapy with pegylated interferon alfa-2a plus ribavirin for 48 weeks (RR 1.6, 95% CI 1.4 to 1.9), with an absolute increase in SVR rate ranging from 25 to 31 percent. The regimen with 8 weeks of telaprevir was associated with a slightly lower SVR rate than the 12 week telaprevir regimen (69 vs. 75 percent) (strength of evidence: low).

- One trial of patients with genotype 1 infection found no difference in likelihood of SVR between triple therapy with telaprevir for 48 weeks (12 weeks of triple therapy with pegylated interferon alfa-2a, ribavirin, and telaprevir followed by 36 weeks of dual therapy with pegylated interferon alfa-2a plus ribavirin) versus triple therapy with telaprevir for 24 weeks (12 weeks of triple therapy followed by 12 weeks of dual therapy) (strength of evidence: low).
- One trial of response-guided triple therapy with telaprevir (24 or 48 weeks, based on absence or presence of HCV-RNA from weeks 4 through 20) found similar SVR rates (81–85 percent) for regimens that varied on telaprevir dose (750 mg three times daily vs. 1125 mg two times daily) and type of pegylated interferon (alfa-2a or alfa-2b) (strength of evidence: low).
- One trial of patients with an extended rapid virologic response to initial triple therapy with telaprevir reported similar high (92 and 88 percent) SVR rates (92 and 88 percent) in patients randomized to a total of 24 or 48 weeks of therapy (strength of evidence: low).

Dual Therapy With Pegylated Interferon Alfa-2a Plus Ribavirin Compared With Dual Therapy With Pegylated Interferon Alfa-2b Plus Ribavirin

Ten trials that directly compared dual therapy with pegylated interferon alfa-2a plus ribavirin to dual therapy with pegylated interferon alfa-2b plus ribavirin in patients with genotypes 1, 2, or 3 infection met inclusion criteria (Table 2, Appendix H: Evidence Table 1). 20-23, 53-58 Two of these trials were published only as abstracts and were included only in sensitivity analyses; we could not adequately assess their quality due to limited information in the abstracts.^{54, 56} One other trial compared pegylated interferon alfa-2a versus pegylated interferon alfa-2b as part of response-guided triple therapy with telaprevir and was also only included in sensitivity analysis.⁵⁹ One trial enrolled a mix of treatment naïve and treatment experienced patients but reported SVR in the treatment-naïve subgroup.⁵⁷ Of the eight trials that could be quality rated, two^{21, 58} were rated poor quality and six were rated fair quality (Appendix H: Evidence Table 2). Frequent methodologic shortcomings were open-label design, ^{20, 21, 23, 53, 55, 57} high or unclear loss to followup, ²⁰⁻²³ and unclear or inadequate methods of allocation concealment. ^{21, 23, 53, 55, 57, 58} Sample sizes ranged from 66 to 3,070. Five trials, including the trial that compared triple therapy regimens, only enrolled patients with genotype 1 HCV infection;^{22, 55, 57-59} the others enrolled either a mix of genotypes or a specific genotype other than genotype 1. The proportion of patients with cirrhosis at baseline ranged from <5–20 percent, ^{20, 23, 59, 60} and the proportion of patients with elevated transaminases ranged from 60–100 percent ^{20, 21, 23, 53, 58, 60} in trials that reported this information. All but two trials 54,57 included a comparison of a standard dose of pegylated interferon alfa-2a (180 mcg/week) with a standard dose of pegylated interferon alfa-2b (1.5 mcg/kg/week). One trial evaluated multiple pegylated interferon alfa-2b doses. ²² Ribavirin dosing varied across studies. All trials used weight-based dosing of ribavirin except for one,

which used an 800 mg daily flat dose (it also enrolled only genotype 3 patients). ⁵⁴ Three trials used different ribavirin doses with pegylated interferon alfa-2a and alfa-2b. ^{22, 23, 59} Nine trials evaluated fixed-duration regimens, with 48 weeks of treatment for genotypes 1 or 4 and 24 weeks for genotypes 2 or 3. 20, 21, 23, 53-58

Table 2. Trials of dual therapy with pegylated interferon alfa-2a plus ribavirin versus dual therapy with pegylated interferon alfa-2b plus ribavirin

with pegylated interferon alfa-2b plus ribavirin								
Trial Country N Quality	Population Characteristics	Genotype Mix	Weekly Pegylated Interferon Dose	Daily Ribavirin Dose	Duration (weeks)	Sustained Virologic Response Rate		
Ascione, 2010 ²⁰ Italy N=320 Quality: Fair	A vs. B Age (mean): 51 vs. 49 years Female: 49% vs. 61% Race: Not reported Cirrhosis: 21% vs. 16% Minimal or no fibrosis: Not reported Elevated transaminases: 100%	~60% genotype 1 or 4	A. Alfa-2a 180 mcg B. Alfa-2b 1.5 mcg/kg	1000-1200 mg	24-48 by genotype	A. 69% B. 54%		
Escudero, 2008 ²¹ Spain N=183 Quality: Poor	A vs. B Age (mean): 44 vs. 44 years Female: 30% vs. 39% Race: Not reported Cirrhosis: Not reported Minimal or no fibrosis: Not reported (100% had at least periportal fibrosis) Elevated transaminases: 100%	~75% genotype 1 or 4	A. Alfa-2a 180 mcg B. Alfa-2b 1.5 mcg/kg	800-1200 mg	24-48 by genotype	A. 66% B. 62%		
Kamal, 2011 ⁵³ Egypt N=217 Quality: Fair	A vs. B Age (mean): 42 vs. 41 years Female: 46% vs. 56% Race: Not reported Cirrhosis: Not reported Minimal or no fibrosis: Not reported Elevated transaminases: 100%	100% genotype 4	A. Alfa-2a 180 mcg B. Alfa-2b 1.5 mcg/kg	1000-1200 mg	48	A. 71% B. 55%		
Khan, 2007 ⁵⁴ Pakistan N=66 Quality: Not assessed ^b	A vs. B Age: Not reported Female: Not reported Race: Not reported Cirrhosis: Not reported Minimal or no fibrosis: Not reported Elevated transaminases: Not reported	100% genotype 3	A. Alfa-2a 180 mcg B. Alfa-2b 1.0 mcg/kg	800 mg	24	A. 79% B. 82%		

Table 2. Trials of dual therapy with pegylated interferon alfa-2a plus ribavirin versus dual therapy with pegylated interferon alfa-2b plus ribavirin (continued)

with pegylated interferon alfa-2b plus ribavirin (continued)							
Trial Country N Quality	Population Characteristics	Genotype Mix	Weekly Pegylated Interferon Dose	Daily Ribavirin Dose	Duration (weeks)	Sustained Virologic Response Rate	
Mach 2011 ⁵⁵ Poland N=260 Quality: Fair	A vs. B Age: 44 vs. 45.2 years Female: 37.7% vs. 42% Race: Not reported (Polish centers) Cirrhosis: Not reported Minimal or no fibrosis: Not reported (78% vs. 73% F0-F2 fibrosis) Elevated transaminases: Not reported	100% genotype 1b	A: Alfa-2a 180 mcg B:Alfa-2b 1.5 mg/kg	1000-1200 mg	48	A. 49% B. 44%	
Magni, 2009 ⁵⁶ Italy N=218 Quality: Not assessed ^b	A vs. B Age: Not reported Female: Not reported Race: Not reported Cirrhosis: Not reported Minimal or no fibrosis: Not reported Elevated transaminases: Not reported	~55% genotype 1 or 4	A. Alfa-2a 180 mcg B. Alfa-2b 1.5 mcg/kg	10.5 mg/kg	24-48 by genotype	A. 68% B. 67%	
Marcellin, 2011 ^{59a} Europe N=161 Quality: Fair	A vs. B vs. C vs. D Age (median): 47 vs. 46 vs. 40 vs. 49 years Female: 50% vs. 52% vs. 48% vs. 51 Non-White race: 10% vs. 10% vs. 10% vs. 8% Cirrhosis: 2.5% vs. 2.4% vs. 0% vs. 5.1% Minimal or no fibrosis: 38% vs. 36% vs. 55% vs. 28% Elevated transaminases: Not reported	100% genotype 1	A. Alfa-2a 180 mcg B. Alfa-2b 1.5 mcg/kg C. Alfa-2a 180 mcg D. Alfa-2b 1.5 mcg/kg	A. 1000- 1200 mg B. 800- 1200 mg C. 1000- 1200 mg D. 800- 1200 mg	24/48	A. 85% B. 81% C. 83% D. 82%	
McHutchison, 2008 (IDEAL) ⁶⁰ U.S. N=3070 Quality: Fair	A vs. B vs. C Age (mean): 48 vs. 48 vs. 48 years Female: 40% vs. 40% vs. 41% Non-White race: 29% vs. 28% vs. 29% Cirrhosis: Not reported (10% vs. 11% vs. 11% severe fibrosis or cirrhosis) Minimal or no fibrosis: Not reported Elevated transaminases: 80% vs. 81% vs. 81%	100% genotype 1	A. Alfa-2a 180 mcg B. Alfa-2b 1.5 mcg/kg C. Alfa-2b 1.0 mcg/kg	A. 1000- 1200 mg B. 800- 1400 mg C. 800- 1400 mg	48	A. 41% B. 40% C. 38%	

Table 2. Trials of dual therapy with pegylated interferon alfa-2a plus ribavirin versus dual therapy with pegylated interferon alfa-2b plus ribavirin (continued)

with pegylated interferon alfa-2b plus ribavirin (continued)							
Trial Country N Quality	Population Characteristics	Genotype Mix	Weekly Pegylated Interferon Dose	Daily Ribavirin Dose	Duration (weeks)	Sustained Virologic Response Rate	
Miyase, 2012 ⁵⁷ Japan N=201 Quality: Fair	A vs. B Age mean: 59.2 vs. 58.9 years Female: 61.4% vs. 60% Nonwhite race: Not reported Cirrhosis: 20% vs. 17% Minimal or no fibrosis: Not reported Elevated transaminases: Not reported	100% genotype 1	A: Alfa-2a 180 mcg B: Alfa-2b 60- 150 mcg/kg (weight- based)	600-1000 mg	48	A. 66% B. 51%	
Rumi, 2010 ²³ Italy N=431 Quality: Fair	A vs. B Age (mean): 52 vs. 53 years Female: 40% vs. 45% Race: Not reported Cirrhosis: 20% vs. 18% Minimal or no fibrosis: Not reported Elevated transaminases (>2 times upper limit of normal): 59% vs. 59%	41% genotype 1 33% genotype 2 15% genotype 3 10% genotype 4	A. Alfa-2a 180 mcg B. Alfa-2b 1.5 mcg/kg	Genotype 1/4: A. 1000- 1200 mg/day for 48 weeks B. 800- 1200 mg/day for 48 weeks Genotype 2/3: A. 800 mg/day for 24 weeks B. 800- 1200 mg/day for 24 weeks	24 -48 by genotype	A: 66% B: 54%	
Yenice, 2006 ⁵⁸ Turkey N=74 Quality: Poor	A vs. B Age (mean): 48 vs. 51 years Female: 35% vs. 27% Race: Not reported Cirrhosis: Not reported Minimal or no fibrosis: Not reported (all patients had at least minimal fibrosis) Elevated transaminases: 70% vs. 76%	100% genotype 1 (1a vs. 1b vs. 1c)	A. Alfa-2a 180 mcg B. Alfa-2b 1.5 mcg/kg	800-1200 mg	24-48 by genotype	A. 49% B. 35%	

Note: Cirrhosis = METAVIR F4, Ishak 5-6, or equivalent. Minimal or no fibrosis=METAVIR F0-F1, Ishak 0-2, or equivalent. ^a All arms included 12 weeks of telaprevir; because this trial compared triple therapy regimens it was excluded from the primary analysis and only included in sensitivity analysis.

^b Published as abstract only; only included in sensitivity analysis.

Dual therapy with a standard dose of pegylated interferon alfa-2b (1.5 mcg/kg/week) plus ribavirin was associated with slightly lower likelihood of achieving an SVR than a standard dose of pegylated interferon alfa-2a (180 mcg/week) plus ribavirin (seven trials, pooled RR 0.87, 95% CI 0.80 to 0.95, I²=27.4%) (Figure 3). 20-23, 53, 55, 58 The pooled absolute reduction in likelihood of SVR was 8 percentage points (95% CI 3 to 14). Results were similar when the meta-analysis included the trial⁵⁹ that evaluated pegylated interferon alfa-2b versus pegylated interferon alfa-2a as part of a triple therapy regimen with telaprevir and ribavirin (eight trials, pooled RR 0.89, 95% CI 0.82 to 0.96, I²=26%) and a trial⁵⁶ only available as a conference abstract (nine trials, pooled RR 0.90, 95% CI 0.83 to 0.97, I²=25%), or excluded two poor-quality trials (five trials, pooled RR 0.86, 95% CI 0.78 to 0.95, I²=47%). Two trials, one published only as an abstract, compared only a standard dose of pegylated interferon alfa-2a (180 mcg weekly) versus nonstandard doses of pegylated interferon alfa-2b (1.0 mcg/kg/week or 60-150 mcg/week). Pooled estimates were similar when these trials were included in the analysis (nine trials, pooled RR 0.88, 95% CI 0.82 to 0.95, I²=22%).

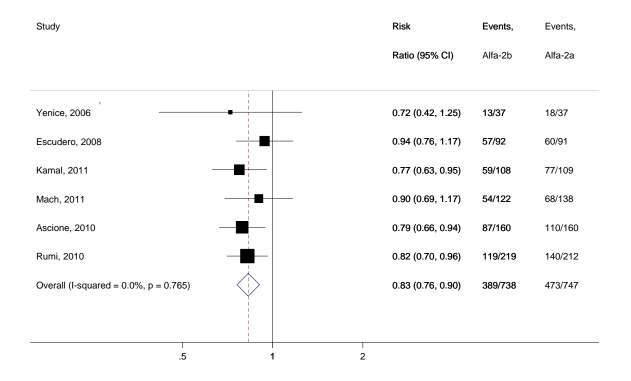
Figure 3. Sustained virologic response: Dual therapy with pegylated interferon alfa-2b plus ribavirin versus dual therapy with pegylated interferon alfa-2a plus ribavirin (standard doses of pegylated interferon only)

Study	Risk	Events,	Events,
	Ratio (95% CI)	Alfa-2b	Alfa-2a
Yenice, 2006	0.72 (0.42, 1.25)	13/37	18/37
Escudero, 2008	0.94 (0.76, 1.17)	57/92	60/91
Kamal, 2011	0.77 (0.63, 0.95)	59/108	77/109
Mach, 2011	0.90 (0.69, 1.17)	54/122	68/138
Ascione, 2010	0.79 (0.66, 0.94)	87/160	110/160
Rumi, 2010 —	0.82 (0.70, 0.96)	119/219	140/212
McHutchison, 2009	0.97 (0.88, 1.08)	406/1019	423/1035
Overall (I-squared = 27.4%, p = 0.220)	0.87 (0.80, 0.95)	795/1757	896/1782
.5 1	2		

The largest head-to-head trial was the Individualized Dosing Efficacy vs. Flat Dosing to Assess Optimal Pegylated Interferon Therapy (IDEAL) study (n=3070, compared with 66 to 477 in the other trials). It was rated fair quality because loss to followup exceeded 20 percent. A three-armed trial, IDEAL randomized patients with HCV genotype 1 infection to one of two doses of pegylated interferon alfa-2b (1.0 mcg/kg/week or 1.5 mcg/kg/week) plus ribavirin 800 to 1400 mg daily (800 mg 40 to 65 kg; 1000 mg >65 to 85 kg; 1200 mg >85 to 105 kg; 1400 >105 to 125 kg) or pegylated interferon alfa-2a 180 mcg/week plus ribavirin 1000 to 1200 mg/day (1000 mg <75 kg; 1200 mg \geq 75 kg). Overall, SVR rates were similar at 38–41 percent in

the three arms. However, differences in ribavirin dosing could have affected treatment comparability. Excluding IDEAL²² had little effect on the pooled estimate and eliminated statistical heterogeneity (six trials, pooled RR 0.83, 95% CI 0.76 to 0.90, $I^2 = 0\%$) (Figure 4).^{20, 21, 23, 53, 55, 58}

Figure 4. Sustained virologic response: Dual therapy with pegylated interferon alfa-2b plus ribavirin versus dual therapy with pegylated interferon alfa-2a plus ribavirin (excluding trials with differential ribavirin dosing or that evaluated triple therapy regimens)



Dual Therapy With Pegylated Interferon Alfa-2a or Alfa-2b Plus Ribavirin: Duration Effects

Table 3. Trials on effects of duration with dual therapy with pegylated interferon (alfa-2a or alfa-2b) plus ribavirin^a

_plus ribavirin ^a						
Trial Country N Quality	Population Characteristics	Percent Genotype 2 or 3	Weekly Pegylated Interferon Dose	Daily Ribavirin Dose	Duration	Sustained Virologic Response Among Patients with Genotype 2 or 3 Infection
	48 W	eeks vs. 24 V	Veeks			
Hadziyannis, 2004 ⁶³ World-wide N=492 with genotype 2 or 3 infection N(total) = 1284 Quality: Fair	A vs. B vs. C vs. D: Age (mean): 41 vs. 42 vs. 43 vs. 43. years Female: vs. 32% vs. 34% vs. 27% vs. 34% Non-White race: 13% vs. 10% vs. 12% vs. 9% Cirrhosis: 7% vs. 8% vs. 5% vs. 7% Minimal or no fibrosis: Not reported	38%	Alfa-2a 180 mcg	A. 800 mg B. 1200 mg C. 800 mg D. 1200 mg	A/B. 48 weeks C/D. 24 weeks	A/B. 75% C/D. 82%
Zeuzem, 2004 ⁷¹ (PEGASYS) Australia, Europe, New Zealand, North & South America N=117 with genotype 2 or 3 infection N(total) = 491 Quality: Fair	A vs. B Age (Mean): 44 vs. 44 Female: 61% vs. 58% Non-White race: 14% vs. 14% Cirrhosis: 1% vs. 0% Minimal or no fibrosis: 69% vs. 66%	28%	Alfa-2a 180 mcg	800 mg	A. 48 weeks B. 24 weeks	A. 78% B. 72%
Guanty: 1 an	24 We	eks vs. 12-16	Weeks	I		l
Lagging, 2008 ⁶⁴ Denmark & Finland N=382 Quality: Fair	A vs. B: Age (mean): 42 vs. 42 years Female: 44% vs. 37% Non-White race: Not reported Cirrhosis: 13% vs. 13% Minimal or no fibrosis: Not reported	100%	Alfa-2a 180 mcg	800 mg	A. 24 weeks B. 12 weeks	A. 78% B. 59%
Manns, 2011 ⁶⁶ International N=458 Quality: Poor	A vs. B Age (Mean): 40 vs. 40 years Female: 35% vs. 36% Non-White race: Not reported Cirrhosis: Not reported Minimal or no fibrosis: Not reported	100%	Alfa-2b 1.5 mcg	800-1400 mg	A. 24 weeksB. 16 weeks	A. 67% B. 57%

Table 3. Trials on effects of duration with dual therapy with pegylated interferon (alfa-2a or alfa-2b) plus ribavirin^a (continued)

plus ribavirin ^a (ce	ontinued)					
Trial Country N Quality	Population Characteristics	Percent Genotype 2 or 3	Weekly Pegylated Interferon Dose	Daily Ribavirin Dose	Duration	Sustained Virologic Response Among Patients with Genotype 2 or 3 Infection
	24 Weeks vs	. 12-16 Week	s (continued)	T	T
Shiffman, 2007 ⁶⁸ 132 Centers World-wide N=1465 Quality: Good	A vs. B: Age (mean): 45.6 vs. 46 years Female: 37% vs. 39% Non-White race: 13% vs. 13% Cirrhosis: Not reported (23% vs. 25% severe fibrosis or cirrhosis) Minimal or no fibrosis: Not reported	100%	Alfa-2a 180 mcg	800 mg	A. 24 weeks B. 16 weeks	A. 70% B. 62%
Yu, 2007 ⁷⁰ Taiwan N=150 Quality: Fair	A vs. B: Age (mean): 50 vs. 49 years Female: 34% vs. 40% Non-White race: Not reported Cirrhosis: Not reported (severe fibrosis or cirrhosis 22% vs. 20%) Minimal or no fibrosis: Not reported (mild, minimal, or no fibrosis 78% vs. 80%)	100%	Alfa-2a 180 mcg	1000- 1200 mg	A. 24 weeks B. 16 weeks	A. 95% B. 94%
24	Weeks vs. 12-16 Weeks Amo	ng Those Wi	th Undetecta	ble Virus by	Week 4	
Dalgard, 2008 ⁶² Denmark, Sweden, Norway N=298 Quality: Fair	A vs. B:: Age (median): 38 vs. 38 years Female: 35% vs. 36% Non-White race: Not reported Cirrhosis: Not reported Minimal or no fibrosis: Not reported	100%	Alfa-2b 1.5 mcg/kg	800-1400 mg	A. 24 weeks B. 14 weeks	A. 91% B. 81%
Mecenate, 2010 (CLEO) ⁶⁷ Italy N=143 Quality: Fair	Demographics reported overall only Age (mean): 43 years Female: 19% Non-White race: Not reported Cirrhosis: 10% (overall) Minimal or no fibrosis: Not reported	100%	Alfa-2a 180 mcg	800-1200 mg	A. 24 weeks B. 12 weeks	A. 75% B. 83%

Table 3. Trials on effects of duration with dual therapy with pegylated interferon (alfa-2a or alfa-2b)

plus ribavirin^a (continued)

Trial Country N Quality	Population Characteristics	Percent Genotype 2 or 3	Weekly Pegylated Interferon Dose	Daily Ribavirin Dose	Duration	Sustained Virologic Response Among Patients with Genotype
						2 or 3 Infection
24 Week	s vs. 12-16 Weeks Among Th	ose With Und	detectable Vi	rus by Week	4 (continue	ed)
von Wagner, 2005 ⁶⁹ Germany N=142 Quality: Fair	A vs. B: Age (mean): 39 vs. 38 Female: 42% vs. 26% Non-White race: Not reported Cirrhosis: Not reported Minimal or no fibrosis: Not reported	100%	Alfa-2a 180 mcg	800-1200 mg	A. 24 weeks B. 16 weeks	A. 80% B. 82%
	Other D	uration Com	parisons			
Andriulli, 2009 ^{61b} Italy N=120 Quality: Fair	A vs. B: Age (mean): 53 vs. 53 years Female: 51% vs. 41% Race: Not reported Cirrhosis: Not reported Minimal or no fibrosis: Not reported	100%	Alfa-2a 180 mcg	A. 1000- 1200 mg for 12 weeks B. 1000- 1200 mg for 6 weeks	12 weeks	A. 82% B. 54%
Mangia, 2005 ⁶⁵ Italy N=283 Quality: Fair	A vs. B: Age (mean): 47 vs. 50 years Female: 44% vs. 44% Race: Not reported Cirrhosis: Not reported (16% vs. 23% severe fibrosis or cirrhosis) Minimal or no fibrosis: Not reported	100%	Alfa-2b 1.0 mcg/kg	1000- 1200 mg	A. 24 weeks B. 12-24 weeks ^c	A. 77% B. 76%

Note: Cirrhosis = METAVIR F4, Ishak 5-6, or equivalent. Minimal or no fibrosis=METAVIR F0-F1, Ishak 0-2, or equivalent.

Six trials compared fixed-duration regimens of dual therapy with pegylated interferon (alfa-2a or alfa-2b) plus ribavirin in patients with genotype 2 or 3 infection. ^{63, 64, 66, 68, 70, 71} Two trials found no difference between 48 versus 24 weeks of dual therapy in likelihood of achieving an SVR (pooled RR 0.97, 95% CI 0.84 to 1.1, I²=43%) (Figure 5). ^{63, 71} Four other trials found 24 weeks of dual therapy associated with a higher likelihood of achieving an SVR than 12 to 16 weeks of dual therapy (pooled RR 1.2, 95% CI 1.0 to 1.3; I²=80%), ^{64, 66, 68, 70} but substantial statistical heterogeneity was present (I²=80%) (Figure 6). Of the four trials, three found 12 to 16 weeks of dual therapy associated with lower likelihood of SVR compared with 24 weeks. ^{64, 66, 68} The fourth trial, ⁷⁰ which found no difference between 16 versus 24 weeks of dual therapy (RR 1.0, 95% CI 0.93 to 1.1), used weight-based dosing of ribavirin starting at 1,000 mg (1,000-1,200 mg), compared with a flat dose of 800 mg or weight-based dosing starting at 800 mg (800-1,400 mg) in the other three trials. This trial also enrolled only patients with a genotype 2 infection, whereas the others enrolled genotype 2 or 3. It reported substantially higher overall SVR rates

^a Sample sizes and results restricted to patients with genotype 2 or 3 infection.

^b Patients who had undetectable HCV-RNA at 4 weeks randomized to 6 or 12 weeks of ribavirin.

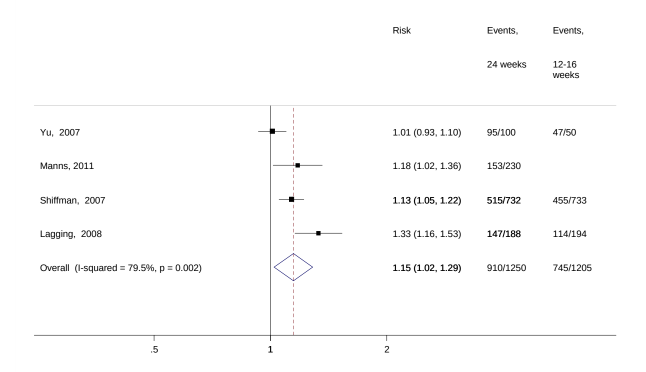
^cTreatment for 12 weeks if HCV RNA undetectable at 4 weeks, and for 24 weeks if detectable.

(94 vs. 95 percent) than the other trials (57–62 percent vs. 67–78 percent). Excluding this trial from the meta-analysis reduced statistical heterogeneity, with no appreciable impact on the pooled estimate of effect (three trials, pooled RR 1.2, 95% CI 1.1 to 1.3, I^2 =47%). Another potential source of heterogeneity was the evaluation of pegylated interferon alfa-2b and high attrition in one of the trials. However, excluding this trial did not affect the pooled estimate or reduce statistical heterogeneity (three trials, pooled RR 1.1, 95% CI 0.99 to 1.3, I^2 =86%).

Figure 5. Sustained virologic response: Dual therapy with pegylated interferon (alfa-2a or alfa-2b) plus ribavirin for 48 versus 24 weeks in patients with genotype 2 or 3 infection

	Risk	Events,	Events,
Study	Ratio (95% CI)	48 Weeks	24 Weeks
Hadziyannis, 2004 —■	0.92 (0.84, 1.01)	190/252	196/240
Zeuzem, 2004	1.08 (0.87, 1.33)	46/59	42/58
Overall (I-squared = 42.7%, p = 0.186)	0.97 (0.84, 1.11)	236/311	238/298

Figure 6. Sustained virologic response: Dual therapy with pegylated interferon (alfa-2a or alfa-2b) plus ribavirin for 24 weeks versus 12 to 16 weeks in patients with genotype 2 or 3 infection



Three trials of patients with genotype 2 or 3 infection who achieved a rapid virologic response (defined as undetectable HCV-RNA by week 4) found no difference between patients randomized to a total of 24 versus 12 to 16 weeks of dual therapy with pegylated interferon alfa-2a (two trials) or alfa-2b (one trial) plus ribavirin (pooled RR 0.99, 95% CI 0.86 to 1.1, I²=66%) (Figure 7).^{62, 67, 69} Although statistical heterogeneity was present, absolute differences were relatively small, ranging from 10 percentage points favoring 24 over 16 weeks of therapy⁶² to 9 percentage points favoring 12 over 24 weeks of therapy.⁶⁷ One trial used the alfa-2b form of pegylated interferon and a somewhat different weight-based ribavirin dosing algorithm, which might account for some of the heterogeneity.⁶²

Figure 7. Sustained virologic response: Dual therapy with pegylated interferon (alfa-2a or alfa-2b) plus ribavirin for a total of 24 versus 12 to 16 weeks in patients with genotype 2 or 3 infection with a rapid virologic response

	Risk	Events,	Events,
Study	Ratio (95% CI)	12-16 weel	ks 24 weeks
Dalgard, 2008 —■—	0.89 (0.81, 0.98)	120/148	136/150
Mecenate, 2010	1.12 (0.94, 1.32)	60/72	53/71
Von Wagner, 2005	1.01 (0.86, 1.19)	57/70	57/71
Overall (I-squared = 65.7%, p = 0.054)	0.99 (0.86, 1.14) 237/290	246/292
.5 1	2		

Two other trials evaluated other comparisons related to duration of dual therapy with pegylated interferon plus ribavirin in patients with HCV genotype 2 or 3 infection. One trial found fixed duration therapy with low dose (1.0 mcg/kg/week) pegylated interferon alfa-2b plus ribavirin for 24 weeks associated with nearly identical likelihood of achieving an SVR versus response-guided therapy for 12 or 24 weeks, based on absence or presence of a rapid virologic response (76 vs. 77 percent). A trial of patients who experienced a rapid virologic response found 12 weeks of pegylated interferon alfa-2a with early discontinuation of ribavirin after 6 weeks associated with lower likelihood of SVR than dual therapy for 12 weeks (54 vs. 82 percent; RR 0.66, 95% CI 0.51 to 0.86).

Dual Therapy With Pegylated Interferon Alfa-2a or Alfa-2b Plus Ribavirin: Dose Effects of Pegylated Interferon (Alfa-2a or Alfa-2b)

Six trials of dual therapy with pegylated interferon plus ribavirin compared lower versus higher doses of pegylated interferon alfa-2b in patients with genotype 2 or 3 infection (Table 4, Appendix H: Evidence Table 7). Three trials $^{66, 73, 77}$ restricted enrollment to patients with genotype 2 or 3 infection and three trials $^{73, 75, 76}$ enrolled other genotypes but reported results in the subgroup of patients with genotype 2 or 3 infection. Sample sizes ranged from 53 to 454 people with genotype 2 or 3 infection. Two trials $^{66, 76}$ were rated poor quality and the remainder fair quality (Appendix H: Evidence Table 8). Methodologic shortcomings included open-label or inadequately described blinding procedures $^{66, 73-77}$ and unclear randomization methods. Five trials compared standard dose pegylated interferon alfa-2b (1.5 mcg/kg/week) compared with lower doses (1.0 or 0.75 mcg/kg/week). The sixth trial evaluated an atypical pegylated interferon alfa-2b dosing regimen of 100-150 mcg weekly (100 mcg if <75 kg or 150 mcg if \geq 75 kg) compared with 50 mcg weekly.

Table 4. Dose effects of pegylated interferon, trials of with dual therapy with pegylated interferon (alfa-2a or alfa-2b) plus ribavirin^a

Author Country N Quality	Population Characteristics	Percent Genotype 2 or 3	Weekly Pegylated Interferon Dose	Daily Ribavirin Dose	Duration	Sustained Virologic Response Among Genotype 2 or 3
	Trials of Higher vs. Lower	Doses of Pe	gylated Interi	feron Alfa-2l	b	
Abergel, 2006 ⁷³ France N=78 with genotype 2 or 3 infection N(total)=203 Quality: Fair	A vs. B Age (mean): 50 vs. 52 years Female: 36% vs. 32% Race: Not reported Cirrhosis: 46% vs. 57% Minimal or no fibrosis: Not reported	38%	A. Alfa-2b 1.5 mcg/kg B. Alfa-2b 0.75 mcg/kg	800 mg	48 weeks	A. 73% B. 73%
Kawaoka, 2009 ⁷⁴ Japan N=53 Quality: Fair	A vs. B Age (median): 57 vs. 55 years Female: 65% vs. 44% Race: Not reported (study conducted in Japan) Cirrhosis: None Minimal or no fibrosis: 55% vs. 48%	100%	A. Alfa-2b 1.0 mcg/kg B. Alfa-2b 1.5 mcg/kg	600-1000 mg	24 weeks	A. 39% B. 74%
Krawitt, 2006 ⁷⁵ U.S. N=86 with genotype 2 or 3 infection N(total) = 301 Quality: Fair	A vs. B Age >50 years: 18% vs. 19% Female: 38% vs. 36% Non-White race: 4.6% vs. 3.1% Cirrhosis: 17% vs. 10% Minimal or no fibrosis: 30% vs. 33%	29%	A. Alfa-2b 50 mcg B. Alfa-2b 100-150 mcg	1000 mg	48 weeks	A. 56% B. 65%
Meyer-Wyss, 2006 ⁷⁶ Switzerland N=91 with genotype 2 or 3 infection N(total)=219 Quality: Poor	A vs. B Age (median): 39 vs. 42 years Female: 43% vs. 28% Race: Not reported Cirrhosis: None Minimal or no fibrosis: 58% vs. 49%	42%	A. Alfa-2b 1.0 mcg/kg B. Alfa-2b 1.5 mcg/kg	800 mg	24-48 weeks by genotype	A. 71% B. 81%
Sood, 2008 ⁷⁷ India N=103 Quality: Fair	A vs. B Age (mean): 43 vs. 37 years Female: 12% vs. 22% Race: Not reported Cirrhosis: Not reported Minimal or no fibrosis: Not reported	100%	A. Alfa-2b 1.0 mcg/kg B. Alfa-2b 1.5 mcg/kg	10-12 mg/kg	24 weeks	A. 79% B. 93%
Manns, 2011 ⁶⁶ International N=454 (24 week) N(total)=602 Quality: Poor	A vs. B Age (Mean): 40 vs. 39 vs. years Female: 35% vs. 40% vs. Non-White race: Not reported Cirrhosis: Not reported Minimal or no fibrosis: Not reported	100%	A: Alfa-2b 1.5 mcg B: Alfa-2b 1.0 mcg	800-1400 mg	24 weeks	A. 67% B. 64%

Table 4. Dose effects of pegylated interferon, trials of with dual therapy with pegylated interferon

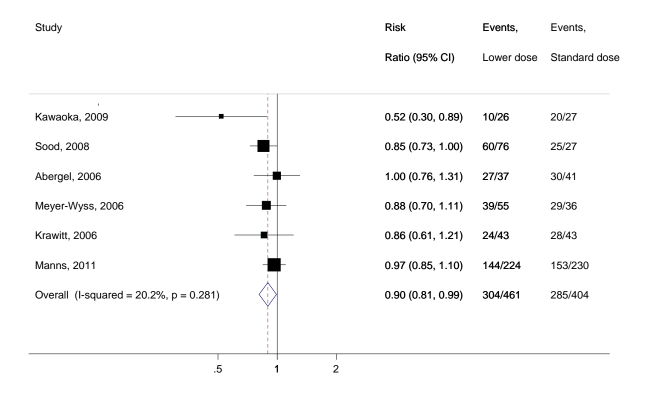
(alfa-2a or alfa-2b) plus ribavirin^a (continued)

Author Country N Quality	Population Characteristics	Percent Genotype 2 or 3	Weekly Pegylated Interferon Dose	Daily Ribavirin Dose	Duration	Sustained Virologic Response Among Genotype 2 or 3
	Trials of Ind	uction Dosin	g Regimens			
Manns, 2001 ⁷⁸ U.S. & UK N=1530 Quality: Fair	A vs. B Age (mean): Female: Non-White race: Cirrhosis: Not reported (29% vs. 30% severe fibrosis or cirrhosis) Minimal or no fibrosis: Not reported	29%	A. Alfa-2b 1.5 mcg/kg x 4 weeks, then 0.5 mcg/kg x 44 weeks B. Alfa-2b 1.5 mcg/kg x 48 weeks	A. 1000- 1200 mg B. 800 mg	48 weeks	A. 80% B. 88%
Mimidis, 2006 ⁷⁹ Greece N=120 Quality: Poor	A vs. B Age (mean): Not reported Female: : 49% vs. 51% Race: Not reported Cirrhosis: Not reported Minimal or no fibrosis: Not reported	51%	A. Alfa-2b 3.0 mcg/kg x 12 weeks, 1.5 mcg/kg x 36 weeks B. Alfa-2b 1.5 mcg/kg x 48 weeks	800-1200 mg	48 weeks	A. 48% B. 59%

HCV = hepatitis C virus; NA = not applicable; SD = standard deviation; U.K. = United Kingdom; U.S. = United States. Note: Cirrhosis = METAVIR F4, Ishak 5-6, or equivalent. Minimal or no fibrosis=METAVIR F0-F1, Ishak 0-2, or equivalent. a Sample sizes and results restricted to patients with genotype 2 or 3 infection.

Lower dose pegylated interferon alfa-2b as part of dual therapy with ribavirin was associated with lower likelihood of SVR than standard dose (six trials, pooled RR 0.90, 95% CI 0.81 to 0.99, I^2 =20%) (Figure 8). $^{66, 73-77}$ Excluding two poor quality trials $^{66, 76}$ (four trials, pooled RR 0.85, 95% CI 0.71 to 1.0, I^2 =38%) or the trial that compared atypical dosing regimens 75 (five trials, pooled RR 0.89, 95% CI 0.79 to 1.0, I^2 =35%) had little effect on the pooled estimates.

Figure 8. Sustained virologic response: Dual therapy with lower dose pegylated interferon alfa-2b plus ribavirin versus higher dose pegylated interferon alfa-2b plus ribavirin in patients with genotype 2 or 3 infection



Two other trials evaluated induction regimens of pegylated interferon alfa-2b (higher initial doses followed by lower doses until completion of therapy) plus ribavirin compared with standard fixed-dose regimens of pegylated interferon alfa-2b plus ribavirin. One good quality trial found dual therapy with pegylated interferon alfa-2b (3.0 mcg/kg/week) plus ribavirin for 12 weeks followed by 36 weeks of standard dose pegylated interferon alfa-2b plus ribavirin associated with a nonstatistically significant trend towards decreased likelihood of SVR versus standard fixed dose dual therapy for 48 weeks (48 vs. 59 percent, p>0.05). Another trial found no clear difference in likelihood of achieving an SVR between dual therapy with standard dose pegylated interferon alfa-2b plus ribavirin for 4 weeks followed by 0.5 mcg/kg/week for 44 weeks versus fixed dose dual therapy with standard doses of pegylated interferon alfa-2b for 48 weeks (82 vs. 80 percent), but results are difficult to interpret because ribavirin dosing was higher (1000 to 1200 mg daily) in the induction compared with the standard therapy arm (800 mg daily).

Ribavirin

Four trials compared effects of dual therapy with pegylated interferon plus ribavirin with different doses of ribavirin in patients with genotype 2 or 3 infection (Table 5, Appendix H: Evidence Table 5). ^{63, 80-82} One trial ⁸⁰ restricted enrollment to patients with genotype 2 or 3 infection and three trials ^{63, 81, 82} enrolled other genotypes but reported results in the subgroup of patients with genotype 2 or 3 infection. Sample sizes ranged from 60 to 1831 with genotype 2 or

3 infection. All four trials were rated fair quality (Appendix H: Evidence Table 6). Methodological shortcomings included open-label design or inadequately described blinding^{63, 80-82} and high loss to followup. Three trials^{63, 80, 81} evaluated ribavirin in combination with pegylated interferon alfa-2a and one trial in combination with pegylated interferon alfa-2b. 82

Table 5. Dose effects of ribavirin: Trials of with dual therapy with pegylated interferon (alfa-2a or

alfa-2b) plus ribavirina

alia-20) pius riba						Sustained
Author Country Study Name N Quality	Population Characteristics	Percent Genotype 2 or 3	Pegylated Interferon Dose	Ribavirin Dose	Duration	Virologic Response Among Genotype 2 or 3
_	A vs. B					
Ferenci, 2008 ⁸⁰ Austria N= 282 Quality: Poor	Age (mean): 37 vs. 36 years Female: 40% vs. 38% Race: Not reported Cirrhosis: Not reported Minimal or no fibrosis: Not reported	100%	Alfa-2a 180 mcg	A. 400 mg B. 800 mg	24 weeks	A. 64% B. 69%
Hadziyannis, 2004 (PEGASYS) ⁶³ Worldwide N=492 with genotype 2 or 3 infection Quality: Fair	A vs. B vs. C vs. D: Age (mean): 41 vs. 42 vs. 43 vs. 43 years Female: 32% vs. 34% vs. 27% vs. 34% Nonwhite race: 12% vs. 9% vs. 13% vs. 10% Cirrhosis: 5% vs. 7% vs. 7% vs. 8% Minimal or no fibrosis: Not reported	38%	Alfa-2a 180 mcg	A/C. 800 mg B/D. 1000- 1200 mg	A. 24 weeks B. 24 weeks C. 48 weeks D. 48 weeks	A/C. 82% B/D. 80%
Helbling, 2006 ⁸¹ Switzerland N= 60 (genotype 2 or 3) N(total)=97 Quality: Fair	A vs. B Age (median): 47 vs. 47 years Female: 30% vs. 40% Race: Not reported Cirrhosis: 57% vs. 52% Minimal or no fibrosis: 6% vs. 2%	48%	Alfa-2a 180 mcg	A. 1000- 1200 mg B. 600- 800 mg	24-48 weeks by genotype	A. 72% B. 45%
Jacobson, 2007a (WIN-R) ⁸² U.S. N=1831 with genotype 2 or 3 infection N(total)=4913 Quality: Fair	A vs. B Age (mean): 46 vs. 46 years Female - 37.7% vs. 36.2% Nonwhite race: 19% vs. 21% Cirrhosis: 10% vs. 10% Minimal or no fibrosis: Not reported (mild, minimal, or no fibrosis 70% vs. 70%)	37%	Alfa-2b 1.5 mcg	A. 800 mg B. 800- 1400 mg	24-48 weeks by genotype	A. 60% B. 62%

Cirrhosis = METAVIR F4, Ishak 5-6, or equivalent. Minimal or no fibrosis=METAVIR F0-F1, Ishak 0-2, or equivalent.

^a Sample sizes and results restricted to patients with genotype 2 or 3 infection.

The trials each evaluated a different ribavirin dose comparison, precluding pooled analyses. The two largest trials found no clear differences between lower flat doses of ribavirin versus higher or weight-based doses. ^{63, 82} One trial (n=492 with genotype 2 or 3 infection) randomized patients to dual therapy with pegylated interferon alfa-2a 180 mcg/week plus flat-dose ribavirin, in one of four regimens: 24 weeks with ribavirin 800 mg/day, 24 weeks with ribavirin 1000–1200 mg/day, 48 weeks with ribavirin 800 mg/day, and 48 weeks with ribavirin 1000–1200 mg/day. ⁶³ Rates of SVR were very similar in the combined 800 mg versus the combined 1200 mg arms (82 vs. 80 percent, RR 1.1, 95% CI 0.99 to 1.2). Another trial (n=1831 with genotype 2 or 3 infection) found no difference between dual therapy for 24 weeks with pegylated interferon alfa-2b 1.5 mcg/kg week and flat-dose ribavirin 800 mg versus weight-dosed ribavirin 800 to 1400 mg (60 vs. 62 percent, RR 0.96, 95% CI 0.89 to 1.0). ⁸² One other smaller trial (n=282) found no difference between dual therapy with pegylated interferon alfa-2a with flat doses of ribavirin 400 mg versus ribavirin 800 mg in likelihood of an SVR (64 vs. 69 percent, RR 0.92, 95% CI 0.78 to 1.1). ⁸⁰

One trial (n=60 with genotype 2 or 3 infection) of pegylated interferon alfa-2a found 600-800 mg daily of ribavirin associated with lower likelihood of SVR than 1000-1200 mg daily (45 vs. 72 percent, RR 0.62, 95% CI 0.40 to 0.98), but differed from the others in that it enrolled subjects primarily with advanced fibrosis or cirrhosis (Ishak stage F4-F6).⁸¹

Trials of Triple Therapy With Pegylated Interferon Alfa-2b, Ribavirin, and Boceprevir

Two randomized trials compared triple therapy with boceprevir, pegylated interferon alfa-2b and weight-based ribavirin with dual therapy with pegylated interferon alfa-2b plus ribavirin in antiviral treatment-naïve patients with chronic HCV genotype 1 infection (Table 6, Appendix H: Evidence Table 3). The Serine Protease Inhibitor Therapy (SPRINT-1) and SPRINT-2 trials (n=1088 and 520, respectively) were conducted in the U.S., Canada, and Europe. In SPRINT-1, percent of enrolled patients had cirrhosis at baseline and in SPRINT-2 about 10 percent had either severe fibrosis or cirrhosis. Both trials were rated fair quality (Appendix H: Evidence Table 4). SPRINT-1 was an open label trial, and in SPRINT-2, 24 percent of patients did not complete followup. Neither trial evaluated the FDA-recommended dosing regimen for boceprevir in antiviral-naïve patients without cirrhosis at baseline (4 weeks of dual therapy leadin with pegylated interferon alfa-2a or alfa-2b plus ribavirin, followed by triple therapy with the addition of boceprevir for either 24 or 32 weeks, based on virologic response at weeks 8 and 24), although both trials evaluated the FDA-recommended dosing regimen for boceprevir in antiviral treatment-naïve patients with cirrhosis at baseline (4 weeks of dual therapy lead-in, followed by triple therapy for the final 44 weeks).

Table 6. Trials of triple therapy with pegylated interferon alfa-2b, ribavirin, and boceprevir versus

dual therapy with pegylated interferon alfa-2b plus ribavirin

Trial Country Study Name N Quality	Population characteristics	Boceprevir Dose / Duration	Weekly Pegylated interferon dose	Daily Ribavirin Dose	Overall Duration of Therapy (weeks)	Sustained Virologic Response
Kwo, 2010 ³⁰ U.S., Canada, Europe Serine Protease Inhibitor Therapy 1 (SPRINT-1) Trial N(total)=520 Quality: Fair	A vs. B vs. C vs. D vs. E Age (mean): 47 vs. 46 vs. 48 vs. 48. vs. 48 years Female: 39% vs. 41% vs. 44% vs. 50% vs. 33% Nonwhite race: 16% vs. 20% vs. 17% vs. 17% vs. 20% Genotype 1: 100% Cirrhosis: 7% (overall) Minimal or no fibrosis: Not reported Elevated transaminases: Not reported	A. BCP 800 mg tid weeks 1-48 B. BCP 800 mg tid weeks 1-28 C. BCP 800 mg tid weeks 5-48 ^a D. BCP 800 mg tid weeks 5-28 E. placebo	Alfa-2b 1.5 mcg/kg	800-1400 mg	A. 48 B. 28 C. 48 D. 28 E. 48	A. 67% B. 54% C. 75% ^a D. 56% E. 38%
Poordad, 2011 ³² U.S. and Europe Serine Protease Inhibitor Therapy 2 (SPRINT-2) N=1,088 Quality: Fair	A vs. B vs. C Age (mean) 49 vs. 50 vs. 49 years Female: 40% vs. 38% vs. 43% Nonwhite race: 19% vs. 17% vs. 18% Genotype 1: 100% Cirrhosis: Not reported (Severe fibrosis or cirrhosis 11% vs. 9% vs. 7%) Minimal or no fibrosis: Not reported Elevated transaminases:	A. 800 mg tid weeks 5-48 B. 800 mg tid weeks 5-28 C. placebo	Alfa-2b 1.5 mcg/kg	A. 600- 1400 mg weeks 5-48 B. 600- 1400 mg weeks 5-28 C. 600- 1400 mg	A. 48 B. 28/48 ^b C. 48	A. 66% ^a B. 63% C. 38%

BCP = boceprevir; bid = twice daily; eRVR = extended rapid virologic response; TCP = telaprevir; tid = three times daily Note: Cirrhosis=METAVIR F4, Ishak 5-6, or equivalent. Minimal or no fibrosis=METAVIR F0-F1, Ishak 0-2, or equivalent. ^a Dosing recommended by the U.S. Food and Drug Administration for boceprevir in antiviral-naïve patients with cirrhosis at baseline

SPRINT-1 randomized patients to five different antiviral regimens: (1) 4-week dual therapy lead-in with pegylated interferon alfa-2b plus ribavirin followed by the addition of boceprevir for 24 weeks (total 28 weeks); (2) 28 weeks of triple therapy with pegylated interferon alfa-2b, ribavirin, and boceprevir with no lead-in; (3) 4-week dual therapy lead-in followed by triple therapy for 44 weeks (total 48 weeks); (4) 48 weeks of triple therapy with no lead-in; or (5) dual therapy for 48 weeks. Takes were 56 percent and 54 percent in the 28-week boceprevir treatment arms and 75 percent and 67 percent in the 48-week boceprevir treatment arms (with and without dual therapy lead- in, respectively), versus 38 percent with dual therapy (p<0.01 for

^b Response-guided duration: 28 weeks of pegylated interferon/ribavirin if HCV-RNA negative from week 8 through week 24. Patients not meeting these criteria continued until week 48.

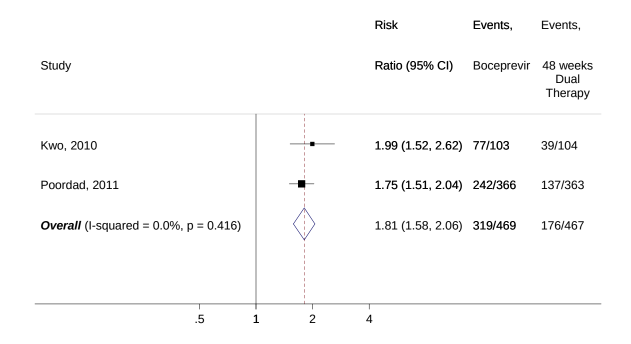
each triple therapy arm vs. dual therapy), for an absolute risk difference for triple compared with dual therapy that ranged from 19–37 percent. Versus dual therapy, the relative risk for achieving an SVR for the two 48-week triple therapy arms combined was 1.9 (95% CI 1.5 to 2.5), and for the two 28-week triple therapy arms combined was 1.5 (95% CI 1.1 to 1.9). Four-week dual therapy lead-in was associated with an increased absolute rate of achieving an SVR versus no lead-in of 2 percent for the 28-week regimens and 8 percent for the 48-week regimens.

SPRINT-2 compared a fixed duration triple therapy regimen, a response-guided triple therapy regimen, and dual therapy.³² The fixed duration regimen consisted of four weeks of dual therapy lead-in with pegylated interferon alfa-2b plus ribavirin followed by the addition of boceprevir for 44 weeks (48 weeks total). The response-guided approach consisted of a four-week dual therapy lead-in, followed by triple therapy for 24 weeks. Patients with undetectable serum HCV-RNA from weeks 8 through 24 completed their antiviral treatment at week 28. Patients with detectable HCV-RNA at any time between weeks 8 and 24 continued dual therapy for another 20 weeks (48 weeks total). The third (control) arm consisted of dual therapy for 48 weeks. SVR rates for the three regimens were 66, 63, and 38 percent, respectively, (p<0.001 for either boceprevir regimen vs. dual therapy), with an absolute risk difference of 25–28 percent for triple compared with dual therapy. Compared with dual therapy, the relative risk for achieving an SVR for the two regimens with boceprevir combined was 1.7 (95% CI 1.5 to 2.0).

The only treatment regimen evaluated in both SPRINT trials was the 48-week regimen with dual therapy lead-in for the first 4 weeks and boceprevir added for the final 44 weeks. Based on data from both trials, triple therapy was associated with a higher likelihood of SVR than dual therapy (pooled RR 1.8, 95% CI 1.6 to 2.1, I^2 =0%), with a pooled absolute increase in SVR of 31 percentage points (95% CI 23 to 39) (Figure 9).

SPRINT-1 also included a separate trial of 75 patients randomized to weight-based low dose (400–1000 mg) or standard dose (800–1400 mg) ribavirin as part of 48 weeks of triple therapy with boceprevir without dual therapy lead in Low dose ribavirin was associated with a non–statistically significant trend towards lower likelihood of SVR (36 vs. 50%, RR 0.71, 95% CI 0.39 to 1.3).

Figure 9. Sustained virologic response: 48 weeks of triple therapy with boceprevir (4 weeks of dual therapy lead-in with pegylated interferon alfa-2b followed by the addition of 44 weeks boceprevir) versus 48 weeks of dual therapy in patients with genotype 1 infection



Trials of Triple Therapy With Pegylated Interferon (Alfa-2a or Alfa-2b), Ribavirin, and Telaprevir

Six randomized trials compared triple therapy with telaprevir, pegylated interferon alfa-2a or alfa-2b and weight-based ribavirin compared with dual therapy with pegylated interferon alfa-2a or alfa-2b and ribavirin for antiviral treatment-naïve patients with chronic HCV genotype 1 infection (Table 7, Appendix H: Evidence Table 3). 31, 51, 59, 85-87 A seventh, small trial was excluded because it evaluated patients with HCV genotype 2 or 3 (telaprevir is only approved for use in genotype 1 infection). 88 One trial 31 was rated good quality and the remainder fair quality (Appendix H: Evidence Table 4). The proportion of patients with cirrhosis at baseline in the trials ranged from 0–10 percent. Methodological shortcomings included open-label design or unclear blinding procedures, ^{59, 85, 87} unclear randomization methods, ^{31, 85} and unclear reporting of attrition. 31, 86 Three trials (n=189 to 323) evaluated fixed duration triple compared with dual therapy regimens (12, 24, or 48 weeks). Two other trials (n=161 and 1088) evaluated response-guided duration triple therapy regimens, including one trial⁵¹ that compared the FDArecommended telaprevir dosing regimen (12 weeks of triple therapy followed by 12 or 36 weeks of dual therapy, depending on early virologic response) with dual therapy. 84 The sixth trial (n=322) compared different durations of antiviral therapy in patients who experienced an extended rapid virologic response. 87 In all evaluated triple therapy regimens, telaprevir was administered with pegylated interferon (alfa-2a or alfa-2b) plus ribavirin for the first 8 to 12 weeks. For regimens longer than 12 weeks, dual therapy with pegylated interferon alfa-2a or alfa-2b plus ribavirin without telaprevir was continued for the remainder of the regimen.

Table 7. Trials of triple therapy with pegylated interferon alfa-2b, ribavirin, and telaprevir

Table 7. Trials of triple therapy with pegylated interferon alfa-2b, ribavirin, and telaprevir						
Author Country Study Name N Quality	Population Characteristics	Telaprevir Dose / Duration	Weekly Pegylated interferon Dose	Daily Ribavirin Dose	Overall Duration of Therapy (weeks)	Sustained Virologic Response
Hezode, 2009 ⁸⁵ Europe N=323 Quality: Fair	A vs. B vs. C vs. D Age (median): 46 vs. 44 vs. 45 vs. 45 years Female: 33% vs. 40% vs. 45% vs. 44% Non White race: 7% vs. 7% vs. 1% vs. 7% Genotype 1: 100% Cirrhosis: 0% vs. 0% vs. 1% vs. 0% Minimal or no fibrosis: 43% vs. 37% vs. 40% vs. 34% Elevated transaminases: Not reported	A. 750 mg tid weeks 1- 12 B. 750 mg tid weeks 1- 12 C. 750 mg tid weeks 1- 12 D. placebo	Alfa-2a 180 mcg	A. 1000- 1200 mg B. 1000- 1200 mg C. placebo D. 1000- 1200 mg	A. 12 B. 24 C. 12 D. 48	A. 69% B. 60% C. 36% D. 46%
Jacobson, 2011 ⁵¹ Worldwide N=1088 Quality: Good	A vs. B vs. C Age (median): 49 vs. 49 vs. 49 years Female: 41% vs. 42% vs. 42% Non White race: 10% vs. 13% vs. 12% Genotype 1: 100% Cirrhosis: 6% overall Minimal or no fibrosis: 28% overall Elevated transaminases: Not reported	A. 750 mg tid weeks 1- 8 B. 750 mg tid weeks 1- 12 ^b C. placebo	Alfa-2a 180 mcg	1000- 1200 mg	A. 24/48 ^a B. 24/48 ^a C. 48	A. 69% B. 75% ^b C. 44%
Kumada, 2012 ⁸⁶ Japan N=189 Quality: Fair	A vs. B Age (mean): 53 vs. 55 Female: 48% vs. 48% Non White: Not reported (conducted in Japan) Minimal or no fibrosis: Not reported Elevated transaminases: Not reported	A. 750 mg tid weeks 1- 12 B. placebo	Alfa-2b 1.5 mcg/kg	600-1000 mg	A. 24 B. 48	A. 73% B. 49%
Marcellin, 2011 ⁵⁹ Europe N=161 Quality: Fair	A vs. B vs. C vs. D Age (median): 47 vs. 46 vs. 40 vs. 49 years Female: 50% vs. 52% vs. 48% vs. 51% Non White race: 10% vs. 10% vs. 10% vs. 8% Genotype 1: 100% Cirrhosis: 2.5% vs. 2.4% vs. 0 vs. 5.1% Minimal or no fibrosis: 39% overall Elevated transaminases: Not reported	A. 750 mg tid weeks 1- 12 B. 750 mg tid weeks 1- 12 C. 1125 mg bid weeks 1- 12 D. 1125 mg bid weeks 1- 12	A. Alfa-2a 180 mcg B. Alfa-2b 1.5 mcg/kg C. Alfa-2a 180 mcg D. Alfa-2a 1.5 mcg/kg	A. 1000- 1200 mg B. 800- 1200 mg C. 1000- 1200 mg D. 800- 1200 mg	24/48 ^c	A. 85% B. 81% C. 83% D. 82%

Table 7. Trials of triple therapy with pegylated interferon alfa-2b, ribavirin, and telaprevir

(continued)

Author Country Study Name N Quality	Population Characteristics	Telaprevir Dose / Duration	Weekly Pegylated interferon Dose	Daily Ribavirin Dose	Overall Duration of Therapy (weeks)	Sustained Virologic Response
McHutchison, 2009 ³¹ U.S. PROVE1 N=250 Quality: Fair	A vs. B vs. C vs. D Age (median): 49 vs. 50 vs. 49 vs. 49 years Female: 32% vs. 39% vs. 29% vs. 43% Non White race: 24% vs. 24% vs. 24% vs. 21% Cirrhosis: 0% Minimal or no fibrosis: 31% (overall) Elevated transaminases: Not reported	A. 750 mg tid weeks 1- 12 B. 750 mg tid weeks 1- 12 C. 750 mg tid weeks 1- 12 D. placebo	Alfa-2a 180 mcg	1000- 1200 mg	A. 12 B. 24 C. 48 D. 48	A. 35% B. 61% C. 67% D. 41%
Sherman, 2011 ⁸⁷ U.S. Name: ILLUMINATE N=322 ^d Quality: Fair	A vs. B Age (median): 51 vs. 50 years Female: 36% vs. 39% Non White race: 17% vs. 18% Cirrhosis: 11% vs. 8% Minimal or no fibrosis: 27% (overall) Elevated transaminases: Not reported	A. 750 mg tid weeks 1- 12 B. 750 mg tid weeks 1- 12	Alfa-2a 180 mcg	1000- 1200 mg	A. 24 B. 48	A. 92% B. 88%

 $bid = two\ times\ daily;\ eRVR = extended\ rapid\ virologic\ response;\ HCV = hepatitis\ C\ virus;\ NA = not\ applicable;\ TCP = telaprevir;\ tid = three\ times\ daily$

Three trials found the 24-week fixed duration triple therapy with pegylated interferon alfa-2a or alfa-2b, ribavirin, and telaprevir associated with higher likelihood of achieving an SVR than 48 weeks of dual therapy (pooled RR 1.5, 95% CI 1.3 to 1.8, I²=0%) (Figure 10). 31, 85, 86 The pooled absolute increase in SVR rates was 22 percentage points (95% CI 13 to 31). Two of the trials found no difference between the 12-week fixed duration triple therapy regimen versus 48 weeks of dual therapy (pooled RR 1.2, 95% CI 0.86 to 1.6, I²=14%) (Figure 11). 31, 85 One of the trials also found a 48-week triple therapy regimen with telaprevir associated with similar likelihood of SVR versus a 24-week triple therapy regimen (RR 1.1, 95% CI 0.87 to 1.4). The other trial also found a 12-week triple therapy regimen of telaprevir plus pegylated interferon without ribavirin associated with a non–statistically significant trend towards lower likelihood of achieving an SVR than pegylated interferon alfa-2a plus ribavirin for 48 weeks (RR 0.77, 95% CI 0.53 to 1.1). So One trial of 24-week fixed duration triple therapy with telaprevir was conducted in Japan, while the other two were conducted in the United States and Europe. Additionally, the Japanese trial studied telaprevir with pegylated interferon alfa-2b, compared

Note: Cirrhosis=METAVIR F4, Ishak 5-6, or equivalent. Minimal or no fibrosis=METAVIR F0-F1, Ishak 0-2, or equivalent.

^a Response-guided duration: 24 weeks of pegylated interferon/ribavirin if HCV-RNA negative from week 4 through week 12. Patients not meeting these criteria continued until week 48.

^b Dosing regimen recommended by the U.S. Food and Drug Administration for telaprevir.

^c Response-guided duration: 24 weeks of pegylated interferon/ribavirin if HCV-RNA negative from week 4 through week . Patients not meeting these criteria continued until week 48.

^d Patients with undetectable HCV RNA at week 4 and week 12 randomized to either 24 or 48 weeks of dual therapy.

with pegylated interferon alfa-2a in the other fixed duration trials. Excluding this trial did not change the pooled result for SVR (two trials, pooled RR 1.5, 95% CI 1.20 to 1.8, $I^2=0\%$). 31,85

Figure 10. Sustained virologic response: Triple therapy with pegylated interferon alfa-2a, ribavirin, and telaprevir for 12 weeks followed by dual therapy for 12 weeks versus dual therapy with pegylated interferon alfa-2a plus ribavirin for 48 weeks in patients with genotype 1 infection

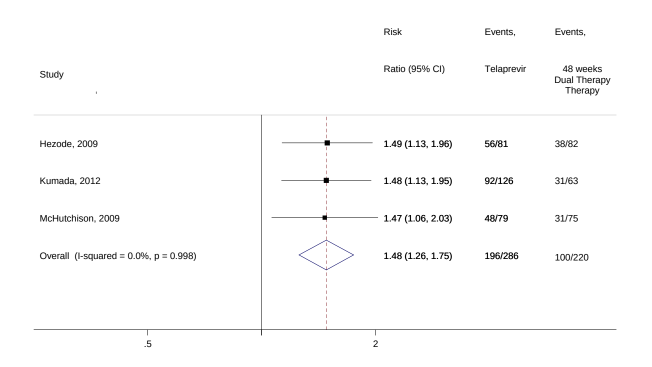
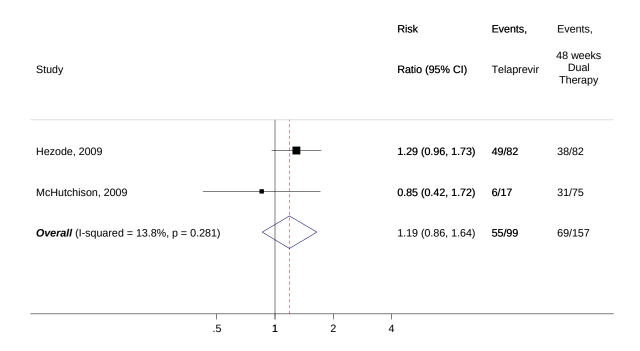


Figure 11. Sustained virologic response: Triple therapy with pegylated interferon alfa-2a, ribavirin, and telaprevir for 12 weeks versus dual therapy with pegylated interferon alfa-2a plus ribavirin for 48 weeks in patients with genotype 1 infection



One trial compared response-guided duration triple therapy with telaprevir compared with dual therapy. Patients were randomized to 8 weeks of initial triple therapy with pegylated interferon alfa-2a, ribavirin, and telaprevir, 12 weeks of initial triple therapy, or dual therapy with pegylated interferon alfa-2a plus ribavirin. In the telaprevir arms, patients with an extended rapid viral response (HCV-RNA undetectable between weeks 4 and 12) continued pegylated interferon plus ribavirin for a total of 24 weeks, while those without an extended rapid viral response continued dual therapy for a total of 48 weeks. Patients randomized to dual therapy received pegylated interferon alfa-2a plus ribavirin for a fixed duration of 48 weeks. Both telaprevir treatment-guided response regimens were associated with higher SVR rates than dual therapy (69, 75, and 44 percent for 8 weeks of telaprevir, 12 weeks of telaprevir, and dual therapy, respectively; p<0.001 for either telaprevir regimen vs. dual therapy), with an absolute increase in SVR ranging from 25–31 percent for triple therapy compared with dual therapy. The relative risk for achieving an SVR in the combined telaprevir arms versus dual therapy was 1.6 (95% CI 1.4 to 1.9).

One trial of response-guided triple therapy with telaprevir (24 or 48 weeks, based on absence or presence of HCV-RNA from weeks 4 through 20) found similar SVR rates (81–85 percent) for regimens that varied on telaprevir dose (750 mg three times daily vs. 1,125 mg two times daily) and type of pegylated interferon (alfa-2a or alfa-2b). Another trial of patients with an extended rapid virologic response to initial triple therapy with telaprevir reported similar, high (92 and 88 percent) SVR rates in patients randomized to a total of 24 or 48 weeks of therapy, meeting the study's predefined noninferiority threshold.

Key Question 2b. How does the comparative effectiveness of antiviral treatment for intermediate outcomes vary according to patient subgroup characteristics, including but not limited to HCV genotype, age, race, sex, stage of disease, or genetic markers?

- The largest randomized trial (n=3070) of dual therapy with pegylated interferon alfa-2a plus ribavirin compared with dual therapy with pegylated interferon alfa-2b plus ribavirin found no clear differences in relative risk estimates for SVR in genotype 1 patients stratified by race, sex, age, baseline fibrosis stage, or baseline viral load. Characteristics associated with lower absolute SVR rates across dual therapy regimens were older age, Black race, advanced fibrosis or cirrhosis, and high baseline viral load (strength of evidence: low).
- Four randomized trials of dual therapy with pegylated interferon alfa-2a plus ribavirin versus dual therapy with pegylated interferon alfa-2b plus ribavirin found no clear differences in relative risk estimates for SVR in patients stratified by genotype. Genotype 1 infection was associated with a lower absolute SVR rate than genotypes 2 or 3 (strength of evidence: moderate).
- Two trials of triple therapy with boceprevir for 48 weeks (4 weeks of dual therapy lead-in with pegylated interferon plus ribavirin followed by 44 weeks of triple therapy with pegylated interferon, ribavirin, and boceprevir) found no difference in relative risk estimates for SVR in men versus women, and no clear difference in relative risk estimates for Black versus non-Black patients. Black race was associated with a lower absolute SVR rate than non-Black race (strength of evidence: moderate).
- Two trials found triple therapy with pegylated interferon alfa-2b, ribavirin, and boceprevir associated with higher likelihood of achieving SVR than dual therapy with pegylated interferon alfa-2b plus ribavirin in patients with high baseline HCV-RNA viral load (>600,000 or ≥800,000 IU/mL), but found no difference in likelihood of SVR in patients with lower viral load (strength of evidence: moderate).
- One trial of response-guided triple therapy with telaprevir (12 weeks of pegylated interferon alfa-2a, ribavirin, and telaprevir followed by response-guided dual therapy with pegylated interferon alfa-2a and ribavirin) versus dual therapy with pegylated interferon plus ribavirin for 48 weeks found no clear differences in relative risk estimates in patients stratified by age, sex, race, baseline fibrosis status, or body mass index. Characteristics associated with lower absolute rates of SVR were older age, Black race, advanced fibrosis or cirrhosis, and higher body mass index. One other trial of 24-week fixed duration triple therapy with telaprevir, pegylated interferon alfa-2b, and ribavirin versus 48 weeks of dual therapy found no differences in estimates of effect in patients stratified by sex or age (strength of evidence: moderate).
- Two trials of triple therapy with pegylated interferon (alfa-2a or alfa-2b), ribavirin, and telaprevir versus dual therapy depending reported inconsistent findings for differential relative risk estimates according baseline viral load (strength of evidence: insufficient).

Dual Therapy With Pegylated Interferon Alfa-2a Plus Ribavirin Compared With Pegylated Interferon Alfa-2b Plus Ribavirin

Five trials of dual therapy with pegylated interferon alfa-2a plus ribavirin versus pegylated interferon alfa-2b evaluated SVR rates in patients subgroups defined by demographic and clinical characteristics (Appendix H: Evidence Table 1 and Evidence Table 2). 20-23, 56 The largest study (n=3070), the IDEAL trial, which only enrolled patients with genotype 1 infection, reported no clear differences in relative risk estimates for SVR dual therapy with pegylated interferon alfa-2b plus ribavirin versus dual therapy with pegylated interferon alfa-2a plus ribavirin in patients stratified by race (RR 0.88, 95% CI 0.59 to 1.3 for Black patients and RR 0.98, 95% CI 0.84 to 1.2 for white patients), sex (RR 0.92, 95% CI 0.77 to 1.1 for males and RR 1.1, 95% CI 0.86 to 1.3 for females), age (RR 0.95, 95% CI 0.77 to 1.2 for <40 years and RR 0.99, 95% CI 0.88 to 1.1 for age >40 years), baseline fibrosis (RR 0.88, 95% CI 0.53 to 1.4 for METAVIR F3 or F4 and RR 0.97, 95% CI 0.87 to 1.1 for METAVIR F0 to F2), and baseline viral load (RR 0.99, 95% CI 0.87 to 1.1 for HCV-RNA >600,000 IU/mL and RR 0.93, 95% CI 0.79 to 1.1 for HCV-RNA \(\leq 600,000\) IU/mL). 22 However, overall absolute SVR rates across dual therapy regimens were lower in older (38 percent) versus younger (53–56 percent) patients, Black patients (23–26 percent) versus white patients (53–55 percent), patients with F3 or F4 (21– 24 percent) versus F0 to F2 fibrosis (42–44 percent), and patients with high (35–36 percent) versus low viral load (61–66 percent). The relative risk estimate was somewhat lower for patients 75 to 85 kg (RR 0.80, 95% CI 0.65 to 0.98) than other weight groups (RR ranged from 0.89 to 1.1) but the confidence intervals for the estimates overlapped, and results were potentially confounded by differential ribayirin dosing according to weight.

Four smaller (n=183 to 431) trials found no clear differences in relative risk estimates in patients stratified by genotype, although rates of SVR were lower by 24–42 percent for genotype 1 infection than genotypes 2 and 3 infection. One of these trials also found no clear differences in relative risk estimates in patient groups stratified by presence or absence of cirrhosis, or high or low viral load. One of these trials also found no clear differences in relative risk estimates in patient groups stratified by presence or absence of cirrhosis, or high or low viral load.

Two trials that compared different durations of therapy in patients with genotype 2 or 3 infection reported risk estimates for SVR stratified by patient characteristics. ^{68, 70} They found no differences in relative risk estimates for 16 weeks of therapy compared with 24 weeks of therapy when patients were stratified according to fibrosis stage, body mass index, sex, or age (all RR estimates close to 1). Although the pooled estimates suggested lower likelihood of SVR with 16 compared with 24 weeks of therapy in patients with HCV-RNA >800,000 IU/mL (pooled RR 0.84, 95% CI 0.77 to 0.93, I²=0%) and no difference in those with a viral load less than 800,000 IU/mL (pooled RR 0.99, 95% CI 0.93 to 1.06, I²=0%), the estimates were imprecise and the confidence intervals overlapped. ^{68, 70}

Another large trial that compared 48 weeks with 24 weeks of dual therapy with pegylated interferon alfa-2a plus ribavirin found similar rates of SVR in patients with genotype 2 or 3 infection regardless of baseline viral load.⁶³

Triple Therapy With Pegylated Interferon (Alfa-2a or Alfa-2b), Ribavirin, and Boceprevir or Telaprevir

Boceprevir

Two trials (n=520 and 1097) of triple therapy with boceprevir for a total of 48 weeks (4 weeks dual therapy lead-in with pegylated interferon alfa-2b plus ribavirin followed by the addition of 44 weeks of boceprevir) versus 48 weeks of dual therapy with pegylated interferon alfa-2b plus ribavirin found no difference in relative risk estimates for SVR in men (pooled RR 1.8, 95% CI 1.6 to 2.2, $I^2=0\%$) versus women (pooled RR 1.9, 95% CI 1.3 to 2.8, $I^2=57\%$). There was also no clear difference in the relative risk estimates for Black (pooled RR 2.5, 95% CI 1.5 to 4.2, $I^2=0\%$) and non-Black patients (pooled RR 1.7, 95% CI 1.5 to 2.0, $I^2=0\%$). although the overall absolute SVR rate across regimens was lower in Black (53 percent) compared with non-Black (63–78 percent) patients. The relative risk estimate was higher for patients with HCV-RNA viral load >600-800,000 IU/mL at baseline (pooled RR 2.0, 95% CI 1.7 to 2.3, $I^2=0\%$) than those with a lower viral load (pooled RR 1.3, 95% CI 1.0 to 1.5, $I^2=0\%$), with an absolute SVR rate of 63–73 percent in individuals with a high viral load and 85–91 percent in individuals with a lower viral load. Although triple therapy with boceprevir was associated with no difference in likelihood of SVR in the subgroup of patients with advanced fibrosis or cirrhosis, the number of patients randomized to triple therapy was small (n=30) and the estimate was imprecise (pooled RR 1.1, 95% CI 0.55 to 2.1, $I^2=0\%$).

Telaprevir

One trial (n=1088) of response-guided duration triple therapy with telaprevir (12 weeks of pegylated interferon alfa-2a, ribavirin, and telaprevir followed by response-guided duration dual therapy) versus 48 weeks of dual therapy with pegylated interferon alfa-2a plus ribavirin found no clear differences in relative risk estimates in patients stratified by age, sex, race, baseline fibrosis status, or body mass index. Absolute SVR rates were higher in patients younger than 45 years versus those older (83 vs. 70 percent), white patients versus Black patients (75 vs. 62 percent), patients with no or minimal fibrosis versus those with advanced fibrosis or cirrhotics (81 vs. 62 percent), and those with body mass index <25 versus those with higher body mass index (83 vs. 69 percent). Triple therapy was more effective than dual therapy in patients with a baseline HCV-RNA viral load ≥800,000 IU/mL (RR 2.0, 95% CI 1.7 to 2.4), but there was no difference in likelihood of achieving an SVR in those with a baseline viral load <800,000 IU/mL (RR 1.1, 95% CI 0.93 to 1.3), with triple therapy associated with similar absolute SVR rates across viral load strata (78 and 74 percent). In a second trial, SVR rates were similar among men (76 percent) and women (70 percent), age less than or greater than 50 (85 vs. 67 percent), and high versus low baseline viral load (69 vs. 74 percent).

Another trial of patients with an extended rapid virologic response on triple therapy with telaprevir reported similar, high (80–90 percent) SVR rates with either 12 versus 36 additional weeks of dual therapy in patients stratified by race, body mass index, or fibrosis stage.⁸⁷

Key Question 3a. What are the comparative harms associated with antiviral treatments?

• Dual therapy with pegylated interferon alfa-2b was associated with slightly greater risk of headache (three trials, pooled RR 1.1, 95% CI 1.1 to 1.2, I²=0%), lower risk of serious

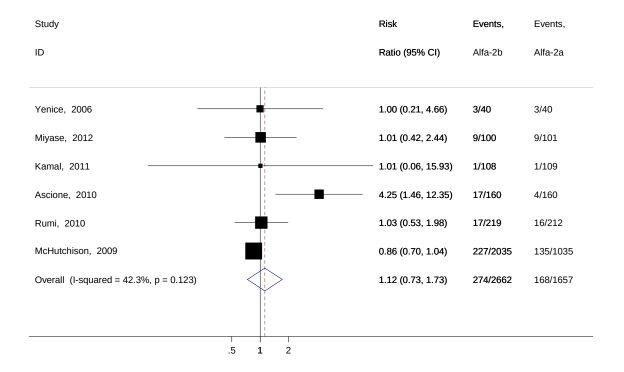
- adverse events (two trials, pooled RR 0.76, 95% CI 0. 71 to 0. 88, I^2 =0%), lower risk of neutropenia (five trials, pooled RR 0.61, 95% CI 0.46 to 0.83, I^2 =38%), and lower risk of rash (two trials, pooled RR 0.79, 95% CI 0.71 to 0.88, I^2 =0%) than dual therapy with pegylated interferon alfa-2a plus ribavirin, with no differences in withdrawals due to adverse events (strength of evidence: moderate).
- Triple therapy with boceprevir for 48 weeks (pegylated interferon alfa-2b plus ribavirin for 4 weeks followed by addition of boceprevir for 44 weeks) was associated with increased risk of neutropenia (two trials, pooled RR 1.8, 95% CI 1.5 to 2.3, I²=0%), dysgeusia (two trials, pooled RR 2.5, 95% CI 2.0 to 3.2, I²=0%), anemia (two trials, pooled RR 2.0, 95% CI 1.4 to 2.8, I²=0%), and thrombocytopenia (two trials, pooled RR 3. 2, 95% CI 1. 2 to 8.2, I²=0%) than dual therapy with pegylated interferon alfa-2b plus ribavirin. The incidence of anemia was about 25 percent with triple therapy and the incidence of neutropenia about 33 percent, with severe anemia in 4–5 percent and severe neutropenia in 8–15 percent. There was no difference in the overall risk of withdrawal due to adverse events (strength of evidence: moderate).
- In two trials, there were no statistically significant differences between a 12-week regimen of triple therapy with pegylated interferon alfa-2a, ribavirin, and telaprevir versus dual therapy with pegylated interferon alfa-2a plus ribavirin in risk of any assessed adverse event (strength of evidence: moderate).
- In three trials, a 24-week regimen of triple therapy with telaprevir (pegylated interferon alfa-2a or alfa-2b, ribavirin, and telaprevir for 12 weeks followed by pegylated interferon alfa-2a plus ribavirin for 12 weeks) was associated with increased risk of anemia (three trials, pooled RR 1.3, 95% CI 1.1 to 1.5, I²=0%) and rash (three trials, pooled RR 1.4, 95% CI 1.1 to 1.7, I²=0%) versus dual therapy with pegylated interferon alfa-2a plus ribavirin for 48 weeks. Among patients randomized to the 24-week telaprevir regimen, one to two-thirds experienced rash (7–10 percent experienced severe rash) and 27–91 percent experienced anemia (4–11 percent experienced severe anemia). There was no difference in risk of withdrawal due to adverse events (strength of evidence: moderate).
- In one trial, response-guided triple therapy with telaprevir (pegylated interferon alfa-2a, ribavirin, and telaprevir for 8 or 12 weeks followed by response-guided duration pegylated interferon alfa-2a and ribavirin) was associated with increased risk of withdrawal due to adverse events (27 vs. 7.2 percent, RR 3.8, 95% CI 2.6 to 5.7), anemia (38 vs. 19 percent, RR 2.0, 95% CI 1.6 to 2.5), any rash (36 vs. 24 percent, RR 1.5, 95% CI 1.2 to 1.8), and severe rash (5 vs. 1 percent, RR 4.6, 95% CI 1.6 to 13) versus dual therapy with pegylated interferon alfa-2a plus ribavirin for 48 weeks (strength of evidence: low).

Dual Therapy With Pegylated Interferon Alfa-2b Plus Ribavirin Compared With Dual Therapy With Pegylated Interferon Alfa-2a Plus Ribavirin

Seven head-to-head randomized trials of dual therapy with pegylated interferon alfa-2b plus ribavirin versus dual therapy with pegylated interferon alfa-2a plus ribavirin reported adverse events (Table 8, Appendix H: Evidence Table 1 and Evidence Table 2). 20-23, 53, 57, 58 Characteristics of the trials were described earlier (see Key Question 2a).

There was no difference between dual therapy with pegylated interferon alfa-2b and dual therapy with pegylated interferon alfa-2a in risk of withdrawal due to adverse events (six trials, pooled RR 1.1, 95% CI 0.73 to 1.7, I²=42%) (Figure 12). 20, 22, 23, 53, 57, 58 In the largest study, the IDEAL trial, about 13 percent of patients randomized to dual therapy with standard doses of pegylated interferon alfa-2b or pegylated interferon alfa-2a plus ribavirin withdrew due to adverse events, versus about 10 percent in those randomized to low-dose pegylated interferon alfa-2b plus ribavirin. Excluding the low-dose pegylated interferon alfa-2b arm of IDEAL from the pooled analysis resulted in a similar pooled estimate (six trials, RR 1.2, 95% CI 0.8 to 1.7, I²=30%). 20, 22, 23, 53, 57, 58 One outlier trial found dual therapy with pegylated interferon alfa-2b associated with substantially higher risk of withdrawal due to adverse events than dual therapy with pegylated interferon alfa-2a (RR 4.2, 95% CI 1.5 to 12). Excluding it eliminated statistical heterogeneity, but the association remained non–statistically significant (five trials, pooled RR 0.88, 95% CI 0.7 to 1.1, I²=0%). 22, 23, 53, 57, 58

Figure 12. Withdrawal due to adverse events: Dual therapy with pegylated interferon alfa-2b plus ribavirin versus dual therapy with pegylated interferon alfa-2a plus ribavirin



Two trials found dual therapy with pegylated interferon alfa-2b plus ribavirin associated with lower risk of serious adverse events than dual therapy with pegylated interferon alfa-2a plus ribavirin (pooled RR 0. 76, 95% CI 0. 61 to 0.95, I²=0%).^{22, 23} In the IDEAL trial, serious treatment-related adverse events occurred in about 4 percent of patients.²² There were no statistically significant differences between regimens in risk of anemia, thrombocytopenia, depression, fatigue, myalgia, or flulike symptoms (Table 8). Dual therapy with pegylated interferon alfa-2b plus ribavirin was associated with slightly greater risk of headache (three trials, pooled RR 1. 1, 95% CI 1.1 to 1. 2, I²=0%)^{20, 22, 57} and slightly lower risk of rash (two trials,

pooled RR 0.79, 95% CI 0.71 to 0.88, $I^2=0\%$)^{22,57} and neutropenia (five trials, pooled RR 0.61, 95% CI 0.46 to 0.83, $I^2=38\%$)^{20-23,57} than dual therapy with pegylated interferon alfa-2a plus ribavirin. In the IDEAL trial, dual therapy with either pegylated interferon (alfa-2a or alfa-2b) was associated with fatigue in about 65 percent of patients, headache in about 45 percent, nausea in about 40 percent, and myalgia in about 25 percent, neutrophil count <500/mm³ in about 5 percent, and hemoglobin <8.5 g/dL in about 3 percent.

Table 8. Harms: Dual therapy with pegylated interferon alfa-2b plus ribavirin versus dual therapy with pegylated interferon alfa-2a plus ribavirin

Outcome	Relative Risk (95% CI); I ²	Number of Trials
All-cause mortality	RR 0.85 (95% CI 0.26 to 2.8)	1 ²²
Serious adverse events	RR 0.76 (0. 61 to 0.95); I ² =0%	2 ^{22, 23}
Withdrawal due to adverse events	RR 1.1 (0.73 to 1.7); I ² =42%	6 ^{20, 22, 23, 53, 57, 58}
Neutropenia	RR 0.61 (0.46 to 0.83); I ² =38%	5 ^{20-23, 57}
Anemia	RR 0.97 (0.72 to 1.3); I ² =64%	4 ^{20, 22, 23, 57}
Thrombocytopenia	RR 0.87 (0.59 to 1.3); I ² =0%	3 ^{20, 23, 57}
Depression	RR 1.1 (0.92 to 1.2); I ² =0%	3 ^{20, 22, 57}
Fatigue	RR 1.0 (0.96 to 1.1): I ² =7%	3 ^{20, 22, 57}
Flulike symptoms	RR 0.98 (0.85 to 1.1)	1 ²³
Headache	RR 1.1 (1.1 to 1.2); I ² =0%	3 ^{20, 22, 57}
Myalgia	RR 1.1 (0.86 to 1.5); I ² =33%	3 ^{20, 22, 57}
Rash	RR 0.79 (0.71 to 0.88); I ² =0%	2 ^{22,57}

RR = relative risk

Excluding data from the IDEAL trial 22 for patients who received pegylated interferon alfa-2b at a lower dose of 1.0 mcg/kg/week had little effect on pooled results, except the pooled estimate for depression became greater and statistically significant in favor of dual therapy with pegylated interferon alfa-2a (three trials, pooled RR 1.2, 95% CI 1.0 to 1.4, I^2 =0%) $^{20, 22, 57}$ There was also reduced statistical heterogeneity in the analysis of neutropenia, but the risk estimate was unchanged (five trials, pooled RR 0.64, 95% CI 0.51 to 0.80, I^2 =0%). $^{20-23, 57}$ Excluding two poorquality trials $^{21, 58}$ from the pooled analysis also had little effect on estimates.

Trials of Triple Therapy With Pegylated Interferon (Alfa-2a or Alfa-2b), Ribavirin, and Boceprevir or Telaprevir

Five trials of triple therapy with pegylated interferon (alfa-2a or alfa-2b), ribavirin, and either boceprevir or telaprevir versus dual therapy with pegylated interferon (alfa-2a or alfa-2b) plus ribavirin without a protease inhibitor in patients with genotype 1 infection reported adverse events (Appendix H: Evidence Table 3 and Evidence Table 4). Characteristics of the trials were described earlier (see Key Question 2a).

Boceprevir

For boceprevir, two trials evaluated a 48-week fixed duration regimen consisting of dual therapy lead-in for 4 weeks with pegylated interferon alfa-2b plus ribavirin, with the addition of boceprevir from weeks 5 through 48.^{30, 32} Triple therapy was associated with increased risk of neutropenia (two trials, pooled RR 1.8, 95% CI 1.5 to 2.3, I²=0%), dysgeusia (two trials, pooled RR 2.5, 95% CI 2.0 to 3.2, I²=0%), anemia (two trials, pooled RR 2.0, 95% CI 1.4 to 2.8,

 I^2 =0%), and thrombocytopenia (two trials, pooled RR 3.2, 95% CI 1.2 to 8.2, I^2 =0%) versus dual therapy with pegylated interferon alfa-2b plus ribavirin (Table 9). About 25 percent of patients on triple therapy experienced anemia and about 33 percent neutropenia, with an incidence of severe neutropenia (neutrophil count <500 cells per μ L) that ranged from 8–15 percent and an incidence of severe anemia (hemoglobin <80 or <85 g/L) of 4–5 percent. In addition, more patients randomized to boceprevir triple therapy used erythropoietin (43 and 87 percent) than those randomized to dual therapy (24 and 33 percent). One of the trials reported similar use of granulocyte stimulating agents with boceprevir triple therapy and dual therapy (8 vs. 6 percent). There were no statistically significant differences between triple therapy and dual therapy in risk of withdrawal due to adverse events, serious adverse events, depression, fatigue, headache, myalgia, chills/rigors, rash, or flulike symptoms (Table 9).

Table 9. Harms: Triple therapy with boceprevir, pegylated interferon alfa-2b, and ribavirin versus

dual therapy with pegylated interferon alfa-2b plus ribavirin

Outcome	Triple Therapy With Pegylated Interferon and Ribavirin for 48 Weeks With Boceprevir From Weeks 5 to 48 vs. Dual Therapy for 48 Weeks: Relative Risk (95% CI); I ²	Number of Trials
Serious adverse events	RR 1.4 (0.93 to 2.2)	1 ³²
Withdrawal due to adverse events	RR 1.1 (0.77 to 1.4); I ² =0%	2 ^{30, 32}
Neutropenia	RR 1.8 (1.5 to 2.3); I ² =0%	2 ^{30, 32}
Anemia	RR 2.0 (1.4 to 2.8); I ² =0%	2 ^{30, 32}
Thrombocytopenia	RR 3.2 (1.2 to 8.2); I ² =0%),	2 ^{30, 32}
Depression	RR 0.87 (0.65 to 1.2)	1 ³²
Fatigue	RR 1.1 (0.82 to 1.5); I ² =82%	2 ^{30, 32}
Flulike symptoms	RR 0.80 (0.58 to 1.1); I ² =27%	2 ^{30, 32}
Headache	RR 1.1 (0.96 to 1.3); I ² =0%	2 ^{30, 32}
Myalgia	RR 0.97 (0.76 to 1.2)	1 ³²
Rash	RR 1.1 (0.81 to 1.4)	1 ³²
Dysgeusia	RR 2.5 (2.0 to 3.2); I ² =0%	2 ^{30, 32}

RR = relative risk

Telaprevir

For fixed duration triple therapy with telaprevir (administered during the first 12 weeks in combination with pegylated interferon and ribavirin), we focused on 12- or 24-week regimens, as 48 week triple therapy regimens have not been shown to be more effective than 24 weeks. There were no differences between a 12-week regimen of triple therapy with telaprevir versus dual therapy with pegylated interferon alfa-2a plus ribavirin for 48 weeks in risk of serious adverse events, neutropenia, anemia, depression, fatigue, headache, myalgia, chills/rigors, rash, or flulike symptoms (Table 10). Rash was reported in 44–77 percent of patients randomized to 12 weeks of triple therapy with telaprevir, with 6 percent of patients reporting severe rash. 31,85

Table 10. Harms: Triple therapy with telaprevir, pegylated interferon (alfa-2a or alfa-2b), and ribavirin versus dual therapy with pegylated interferon (alfa-2a or alfa-2b) plus ribavirin

Outcome	12-Week Regimen With Telaprevir vs. Dual Therapy for 48 Weeks: Relative Risk (95% CI); I ²	Number of Trials	24-Week Regimen With Telaprevir vs. Dual Therapy for 48 Weeks: Relative Risk (95% CI); I ²	Number of Trials
All-cause mortality	No deaths reported	No deaths reported	No deaths reported	No deaths reported
Serious adverse events	RR 1.3 (0.68 to 2.5)	1 ⁸⁵	RR 1.0, 95% CI 0.50 to 2.0)	1 ⁸⁵
Withdrawal due to adverse events	RR 1.5 (0.56 to 4.0)	1 ⁸⁵	RR 1.1 (0.45 to 2.6); I ² =60%	2 ^{85, 86}
Neutropenia	RR 0.11 (0.01 to 1.8)	1 ³¹	RR 0.81 (0.51 to 1.3); I ² =53%	2 ^{31, 86}
Anemia	RR 1.2 (0.72 to 1.9); I ² =0%	2 ^{31, 85}	RR 1.3 (1.1 to 1.5); I ² =0%	3 ^{31, 85, 86}
Thrombocytopenia	Not reported	Not reported	RR 1.8 (1.2 to 2.5)	1 ⁸⁶
Depression	RR 0. 90 (0.53 to 1.5); I ² =0%	2 ^{31, 85}	RR 1.0 (0.66 to 1.6); I ² =0%	2 ^{31, 85}
Fatigue	RR 0.94 (0.63 to 1.4); I ² =61%	2 ^{31, 85}	RR 0.96 (0.74 to 1.2); I ² =53%	3 ^{31, 85}
Flulike symptoms	RR 0.76 (0.56 to 1.0); I ² =0%	231,85	RR 0.87(0.63 to 1.2); I ² =50%	3 ^{31, 85, 86}
Headache	RR 0.87 (0.65 to 1.2); I ² =0%	2 ^{31, 85}	RR 0.83 (0.69 to 1.0); I ² =0%	3 ^{31, 85, 86}
Myalgia	RR 0.71 (0.40 to 1.3); I ² =0%	2 ^{31, 85}	RR 0.76 (0.43 to 1.3); I ² =57%	3 ^{31, 85}
Rash	RR 1.2 (0.92 to 1.7); I ² =0%	2 ^{31, 85}	RR 1.4 (1.1 to 1.7);I ² =0%	3 ^{31, 85, 86}

CI = confidence interval; RR = relative risk

A 24-week regimen of triple therapy with telaprevir was associated with increased risk of anemia (three trials, pooled RR 1.3, 95% CI 1.1 to 1.5, $I^2=0\%$) and increased risk of rash (three trials, pooled RR 1.4, 95% CI 1.1 to 1.7, I²=0%) versus dual therapy for 48 weeks, but there were no statistically significant differences in risk of serious adverse events, neutropenia, depression, fatigue, headache, chills/rigors, or flulike symptoms (Table 10). 31, 85, 86 Triple therapy was also associated with increased risk of thrombocytopenia, but this outcome was only evaluated in one trial (RR 1.8, 95% CI 1.2 to 2.5). 86 One-third to two-thirds of patients randomized to the 24week regimen with telaprevir experienced a rash, with the incidence of severe rash ranging from 7–10 percent. 31, 85, 86 The incidence of anemia with telaprevir was 27–91 percent, 31, 85, 86 with two trials^{31,85} reporting severe anemia in 4–9 percent of patients and another trial⁸⁶ reporting grade 3 anemia (hemoglobin <8 g/dl) in 11 percent of patients. Two trials found no difference in risk of withdrawal due to adverse events (RR 1.1, 95% CI 0.45 to 2.6, I²=60%). 85, 86 The third trial did not report withdrawal due to adverse events separately for the 24 week telaprevir regimen, but reported a similar trend towards higher risk of withdrawal due to adverse events for all telaprevir regimens combined (12, 24, or 48 weeks) versus dual therapy (21 vs. 11 percent, RR 2.0, 95% CI 0.97 to 4.1).³¹

One trial evaluated triple therapy with telaprevir for 8 or 12 weeks followed by response-guided dual therapy for 12 or 36 weeks versus dual therapy for 48 weeks.⁵¹ Since the two telaprevir regimens were associated with similar rates of harms, results were combined. The trial found response-guided therapy with telaprevir associated with increased risk of withdrawal due to adverse events (27 vs. 7.2 percent, RR 3.8, 95% CI 2.6 to 5.7), anemia (38 vs. 19 percent, RR

2.0, 95% CI 1.6 to 2.5), any rash (36 vs. 24 percent, RR 1.5, 95% CI 1.2 to 1.8), and severe rash (5 vs. 1 percent, RR 4.6, 95% CI 1.6 to 13).

A trial of extended early virologic responders (undetectable HCV-RNA levels at weeks 4 and 12) to telaprevir triple therapy reported very similar rates of adverse events in patients randomized after 20 weeks of therapy to 4 weeks versus 28 more weeks of dual therapy.⁸⁷ The overall incidence of rash was 38 percent (severe rash 5 percent) and the incidence of anemia 42 percent (severe anemia 6 percent).

Key Question 3b. Do these harms differ according to patient subgroup characteristics, including HCV genotype, age, race, sex, stage of disease, or genetic markers?

- No trial of dual therapy with pegylated interferon alfa-2b plus ribavirin versus dual therapy with pegylated interferon alfa-2a plus ribavirin reported harms in patients stratified by factors such as HCV genotype, age, race, sex, stage of disease, or genetic markers. Three trials that restricted enrollment to patients with genotype 1 infection reported risk estimates for risk of harms that were similar to the risk estimates based on all trials (strength of evidence: insufficient).
- No trial evaluated harms associated with triple therapy with pegylated interferon, ribavirin, and boceprevir or telaprevir versus dual therapy with pegylated interferon plus ribavirin in patient subgroups. All trials evaluated patients with genotype 1 infection (strength of evidence: insufficient).

No trial of dual therapy with pegylated interferon alfa-2b plus ribavirin versus dual therapy with pegylated interferon alfa-2a plus ribavirin reported harms in patients stratified by factors such as HCV genotype, age, race, sex, stage of disease, or genetic markers. A subgroup of three trials of dual therapy with pegylated interferon alfa-2a versus pegylated interferon alfa-2b that restricted enrollment to patients with genotype 1 infection reported pooled estimates for risk of harms that were similar to the risk estimates based on all trials. ^{22, 58, 59} All trials of triple therapy including protease inhibitors restricted enrollment to patients with genotype 1 infection.

Key Question 4. Have improvements in intermediate outcomes (SVR, histologic changes) been shown to reduce the risk or rates of adverse health outcomes from HCV infection?

- A large Veterans Affairs (VA) study that controlled well for potential confounders found an SVR after antiviral therapy associated with lower risk of all-cause mortality versus no SVR (adjusted HR 0.71 [0.60–0.86], 0.62 [0.44–0.87] and 0.51 [0.35–0.75] for genotypes 1, 2, and 3, respectively). Eighteen other cohort studies found an SVR associated with decreased risk of all-cause mortality, liver-related mortality, HCC, and other complications of end-stage liver disease versus no SVR, with stronger effect estimates than the VA study (adjusted HRs generally ranged from around 0.10 to 0.33). However, the studies had methodological shortcomings, including inadequate handling of confounders, and 10 were conducted in Asia (strength of evidence: moderate).
- Nine studies found an SVR associated with greater improvement in measures related to quality of life (generic or disease-specific) 24 weeks after the end of antiviral treatment versus no SVR, with differences averaging less than 5 to 10 points on various SF-36 domains. All studies were poor quality and were characterized by failure to adjust for

confounders, high loss to followup, and failure to blind patients to SVR status (strength of evidence: low).

All-Cause Mortality, Liver-Related Mortality, and Complications Related to Chronic Hepatitis C Virus Infection

Nineteen cohort studies evaluated the association between achieving an SVR following interferon-based antiviral therapy and mortality (all-cause or liver-related) or complications related to chronic HCV infection, such as HCC, ascites, hepatic encephalopathy, or gastrointestinal bleeding, and these 19 studies reported risk estimates adjusted for potential confounders (Table 11, Appendix F, Appendix H: Evidence Table 9). 8, 9, 89-105 Sample sizes ranged from 105 to 16,864 subjects and duration of followup ranged from 3 to 9 years. Ten studies were conducted in Asia. 89, 95-100, 102, 104, 105 Four studies focused on patients who received pegylated interferon (alfa-2a or alfa-2b) plus ribavirin. Four studies focused on patients who received nonpegylated interferon plus ribavirin, or either pegylated or nonpegylated interferon monotherapy. Ten studies 8, 89, 92, 97-101, 104, 105 evaluated general populations of HCV patients treated with antiviral therapy (baseline rate of cirrhosis ranged from 3–20 percent) and nine studies 9, 90, 91, 93-96, 102, 103 focused on patients with advanced fibrosis or cirrhosis at the time of antiviral treatment. Six studies 90, 93-96, 102 enrolled patients with cirrhosis only, and the baseline rate of cirrhosis ranged from 21–77 percent in three others.

All studies had methodological shortcomings (Appendix H: Evidence Table 10). Eight studies ^{92-94, 100, 102-105} were rated poor quality and the remainder fair quality. Although all of the studies reported adjusted risk estimates, only eight^{8, 89, 91, 95-98, 101} of the 19 studies evaluated five key potential confounders (age, sex, genotype, viral load, and fibrosis stage). No study clearly described assessment of outcomes blinded to SVR status and only five studies^{8, 94, 97, 98, 102} reported the number of patients who met inclusion criteria but were excluded due to missing data or loss to followup.

For general populations of HCV patients treated with antiviral therapy, the largest study (n=16,864) had the fewest methodological shortcomings and was also conducted in the United States. (Appendix H: Evidence Table 11). It adjusted for multiple potential confounders, including age, sex, viral load, presence of cirrhosis, multiple comorbidities, aspartate aminotransferase (AST) and alanine aminotransferase (ALT) levels, and others; and stratified results by genotype. In a predominantly male (>95 percent) population of veterans, the study found SVR after antiviral therapy associated with decreased risk of all-cause mortality versus no SVR, after a median followup of 3.8 years (adjusted HR 0.71 [0.60 to 0.86], 0.62 [0.44 to 0.87] and 0.51 [0.35 to 0.75] for genotypes 1, 2, and 3, respectively). Although point estimates showed somewhat smaller effects for genotype 1 compared with genotypes 2 or 3, the confidence intervals for the three genotypes overlapped. The very rapid (within 3 months after assessing for SVR for genotype 3) separation of mortality curves suggests possible residual confounding, given the expected duration required to observe benefits in long-term clinical outcomes. Clinical outcomes other than mortality were not assessed.

Nine other studies also evaluated the association between achieving an SVR and mortality or hepatic complications in general populations of HCV patients (Appendix H: Evidence Table 11). ^{89, 92, 97-101, 104, 105} One fair-quality study from Scotland found an SVR after antiviral therapy associated with decreased risk of liver-related mortality (adjusted HR 0.22, 95% CI 0.09 to 0.58) and liver-related hospital episodes (adjusted HR 0.22, 95% CI 0.15 to 0.34) versus no SVR. ⁹⁸ One Australian study (poor quality) found no statistically significant association between

virologic response status (SVR, response-relapse, or nonresponse) and all-cause mortality, liver-related mortality, or HCC, although trends favored the SVR group. The other seven studies (three poor quality), all conducted in Asia, each found an SVR after antiviral therapy associated with substantially lower risk of all-cause mortality, liver-related mortality, or HCC versus no SVR. Sy, 97, 99-101, 104, 105 Six studies reported substantially lower risk for all-cause mortality than the U.S. study described above (adjusted HR range 0.12 to 0.39). Sy, 97, 100, 101, 104, 105 For liver-related mortality, four studies Sy, 97, 100, 104 reported adjusted HRs that ranged from 0.04 to 0.17 and for HCC, four studies reported adjusted HRs that ranged from of 0.12 to 0.36.

Six studies of European or North American populations (two poor quality) evaluated the association between achieving an SVR after antiviral therapy and clinical outcomes in patients with advanced fibrosis and cirrhosis prior to antiviral treatment (Appendix H: Evidence Table 11). 9,90,91,93,94,103 One study (fair quality) found an SVR after antiviral therapy associated with decreased risk of all-cause mortality or liver transplantation versus no SVR (adjusted HR 0.17, 95% CI 0.06 to 0.46). Another study (poor quality) found an SVR associated with decreased risk of all-cause mortality (adjusted HR 0.31, 95% CI 0.07 to 1.4). Four studies found an SVR associated with decreased risk of liver-related mortality and HCC versus no SVR (adjusted HRs ranged from 0.12 to 0.27 and from 0.19 to 0.46, respectively). For complications of chronic HCV infection (variably defined), six studies reported adjusted HRs that ranged from 0.13 to 0.38. Results from three Asian studies for one poor quality) were consistent with the North American and European studies. One study found an SVR associated with lower risk of all-cause mortality versus no SVR (adjusted HR 0.07, 95% CI 0.09 to 0.56) and three studies found an SVR associated with lower risk of All-cause mortality versus no SVR (adjusted HR 0.07, 95% CI 0.09 to 0.56) and three studies found an SVR associated with lower risk of HCC versus no SVR (adjusted HR range 0.18 to 0.40).

One study stratified results according to presence or absence of cirrhosis of baseline. Although effects of an SVR versus no SVR on all-cause mortality appeared more favorable in patients with cirrhosis compared with those without cirrhosis, estimates were imprecise and confidence intervals overlapped substantially, precluding strong conclusions. ¹⁰⁴

The only study to evaluate the association between improvement in histological outcomes and clinical outcomes did not meet inclusion criteria because it did not report adjusted risk estimates. ¹⁰⁶ In 96 patients with chronic HCV infection and cirrhosis, it found regression of cirrhosis (defined as a decrease in METAVIR fibrosis score from 4 to ≤2) after interferon-based therapy associated with decreased risk of liver-related events (ascites, hepatic encephalopathy, variceal bleeding, spontaneous bacterial peritonitis, HCC, or liver transplantation) or death (0 vs. 4 events/100 patients-years, p=0.002) after a median followup of 10.5 years. Transplantation-free survival was 100 percent in patients with regression of cirrhosis compared with 74 percent in those without regression (p=0.02). In addition to failure to analyze potential confounders, the study only included patients who underwent a post-treatment biopsy, which could have resulted in selection bias, and cirrhosis regression only occurred in 13 patients, resulting in low precision.

Table 11. Sust	Comparison	esponse and clinical outcon	103	
Author Country N Quality	Definition of Sustained Virologic Response	Population Characteristics	Treatments	Results
Arase, 2007 ⁸⁹ Japan N=500 Quality: Fair	SVR vs. no SVR SVR=Undetecta ble HCV-RNA 6 months after completion of long-term IFN therapy	SVR (n=140) vs. no SVR (n=360) Mean age (years): 63 vs. 64 (p=0.07) Female: 41% vs. 53% (p=0.01) Race: Not reported Genotype 1b: 34% vs. 71% (p<0.0001) Viral load (kIU/ml): 172 vs. 661 (p<0.0001) Cirrhosis (Knodell F4): 9% vs. 16% (p=0.009)	Interferon-2a or Interferon- 2b monotherapy: 94% Interferon plus ribavirin combination therapy: 6%	SVR vs. no SVR HCC: Adjusted HR 0.19 (0.08-0.45) All-cause mortality: Adjusted HR 0.39 (0.16-0.93) Liver-related mortality: Adjusted HR 0.13 (0.03- 0.59)
Backus, 2011 ⁸ U.S. N=16,864 Quality: Fair	SVR vs. no SVR SVR=Undetecta ble HCV-RNA 6 months after completion of antiviral therapy	SVR vs. no SVR (genotypes 1 [n=12,166], 2 [n=2904], and 3 [n=1794], respectively) Mean age (years): 51 vs. 52, 53 vs. 53. and 51 vs. 51 Female: 5% vs. 4%, 4% vs. 3%, and 4% vs. 3% Non-White: 40% vs. 51%, 33% vs. 31%, and 30% vs. 29% Genotype: Results stratified by genotype Viral load ≥500,000 IU/mL: 70% vs. 82%, 78% vs. 83%, and 64% vs. 68% Cirrhosis: 9% vs. 15%, 7% vs. 12%, and 12% vs. 20%	Pegylated interferon (alfa-2a or alfa-2b) plus ribavirin	SVR vs. no SVR (genotypes 1, 2, and 3, respectively) All-cause mortality: Adjusted HR 0.71 (0.60-0.86), 0.62 (0.44-0.87), and 0.51 (0.35-0.75)
Bruno, 2007 ⁹⁰ Italy N=883 Quality: Fair	SVR vs. no SVR SVR=Undetecta ble HCV-RNA 6 months after completion of antiviral therapy	SVR (n=124) vs. no SVR (n=759) Mean age (years): 53 vs. 44 (p=0.004) Female: 27% vs. 38% (p<0.001) Non White: 0 (0%) vs. 0 (0%) Race: Not reported Genotypes 1 and 4: 37% vs. 63% (p<0.001) Viral load: Not reported Cirrhosis: All (inclusion criterion)	Interferon monotherapy	SVR vs. no SVR Ascites, encephalopathy, or gastrointestinal bleeding: Not calculated, 0 events/1061 person-years vs. 107 events/5703 person-years (1.88 events/100 person- years) HCC: Adjusted HR 0.39 (0.17-0.88) Liver-related mortality: 0.14 (0.04-0.59)
Cardoso, 2010 ⁹¹ France N=307 Quality: Fair	SVR vs. no SVR SVR=Undetecta ble HCV-RNA 6 months after completion of antiviral therapy	SVR (n=103) vs. no-SVR (n=204) Mean age (years): 55 vs. 55 (p=0.93) Female: 30% vs. 34% (p=0.51) Race: Not reported Genotype 1: 36% vs. 72% (p<0.001) Viral load (log ₁₀ l/ml): 5.5 vs. 5.7 (p=0.08) Cirrhosis (METAVIR F4): 53% vs. 61% (p=0.19)	Pegylated interferon (alfa-2a or alfa-2b) and ribavirin: 252 (82%) Pegylated interferon monotherapy: 22 (7%) Nonpegylated interferon with or without ribavirin: 33 (11%)	SVR vs. no SVR HCC: Adjusted HR 0.33 (0.23-0.89) Ascites or variceal bleeding: Adjusted HR 0.21 (0.05- 0.92) Liver-related mortality: Adjusted HR 0.27 (0.08- 0.95)

Author Country N Quality	Comparison Definition of Sustained Virologic Response	Population Characteristics	Treatments	Results
Coverdale, 2004 ⁹² Australia N=343 Quality: Poor	SVR vs. response relapse vs. nonresponse SVR=Undetecta ble HCV-RNA on at least 2 occasions at least 2 years after completion of therapy	Demographics for all treated patients (not reported by SVR status) Median age (years): 37 Female: 33% Race: Not reported Genotype 1: 38% Viral load: Not reported Median fibrosis score (Scheuer): 2	Interferon-2a or Interferon- 2b	SVR vs. response-relapse vs. nonresponse Liver-related complications (hepatic decompensation, complications of portal hypertension, HCC, liver transplantation, and liver-related mortality) at 10 years: Not statistically significant in multivariate analysis, adjusted HR not reported (p=0.06) HCC at 10 years: Not statistically significant in multivariate analysis, adjusted HR and p value not reported Liver transplant or liver-related death at 10 years: Not statistically significant in multivariate analysis, adjusted HR not reported (p=0.20)
El Braks, 2007 ⁹³ France N=113 Quality: Poor	SVR vs. no SVR SVR=Undetecta ble HCV-RNA 6 months after completion of antiviral therapy	SVR (n=37) vs. no SVR (n=76) Mean age (years): 51 vs. 56 (p=0.02) Female: 16% vs. 50% (p=0.0005) Race: Not reported HCV genotype 1: 36% vs. 73% (p=0.0001) Viral load: Not reported Cirrhosis: All (inclusion criterion)	Interferon monotherapy: 35/113 (31%) Interferon + ribavirin: 40/113 (35%) Pegylated interferon + ribavirin: 38/113 (34%)	SVR (n=37) vs. no SVR (n=76) Clinical events (HCC, ascites, hepatic encephalopathy, or death): Adjusted HR 0.14 (0.04-0.45)
Fernandez- Rodriguez, 2010 ⁹⁴ Spain N=509 Quality: Poor	SVR vs. no SVR SVR=Undetecta ble HCV-RNA 6 months after completion of antiviral therapy	SVR (n=174) vs. no SVR (n=394) Mean age (years): 51 vs. 52 (p=0.31) Female: 69% vs. 73%, p=0.37 Genotype 1: 24% vs. 55% (p=0.001) Race: Not reported Viral load (10 ⁶ IU/ml): 1.7 vs. 3.1 (p=0.001) Cirrhosis: All (inclusion criterion)	Pegylated interferon-2a or 2b	SVR vs. no SVR Combined clinical endpoint (hepatic decompensation, upper gastrointestinal bleeding secondary to rupture of esophageal or gastric varices, HCC, liver transplantation, and liver- related or liver-unrelated mortality): Adjusted HR 0.38 (0.18-0.76)

Author Country N Quality	Comparison Definition of Sustained Virologic Response	Population Characteristics	Treatments	Results
Hasegawa, 2007 ⁹⁵ Japan N=105 Quality: Fair	SVR vs. no SVR SVR=Sustained undetectable HCV-RNA after completion of antiviral therapy (duration of undetectability not specified)	SVR (n=48) vs. no SVR (n=58) Age >56 years: 60% vs. 55% (p>0.05) Female: 35% vs. 34% (p>0.05) Race: Not reported Genotype 1b: 19% vs. 21% (p>0.05) Viral load ≥100 KIU/ml or ≥1 mg/mL: 25% vs. 62% (p<0.001) Cirrhosis: All (inclusion criterion)	Natural or recombinant interferon alfa: 67% Natural interferon- beta: 31% Both: 1.6%	SVR vs. no SVR HCC: Adjusted HR 0.18 (0.04-0.81)
Hung, 2006 ⁹⁶ Taiwan N=132 Quality: Fair	SVR vs. no SVR SVR=Undetecta ble HCV-RNA 6 months after completion of antiviral therapy	SVR (n=73) vs. no SVR (n=59) Mean age (years): 55 vs. 58 (p=0.07) Female: 43% vs. 54% (p=0.12) Race: Not reported Genotype 1b: 27% vs. 78% (p<0.001) Viral load ≥2 x 10 ⁶ copies/ml: 21% vs. 51% (p<0.001) Cirrhosis: 100% (inclusion criterion)	Interferon-2b plus ribavirin	SVR vs. no SVR HCC: Adjusted HR 0.28 (0.09-0.92)
Imazeki, 2003 ⁹⁷ Japan N=459 Quality: Fair	SVR vs. no SVR SVR=Undetecta ble HCV-RNA 6 months after completion of antiviral therapy	Demographics for all treated patients (not reported by SVR status) Mean age (years): 49 Female: 36% Race: Not reported Genotype 1: 74% Viral load: Not reported Cirrhosis (Desmet F4): 13%	Interferon-2a: 84% Interferon-2b: 12% Both: 4%	SVR vs. no SVR ^a Liver-related mortality: Adjusted HR 0.11 (0.01- 0.96) All-cause mortality: Adjusted HR 0.12 (0.01-1.3)
Innes, 2011 ⁹⁸ UK N=1,215 Quality: Fair	SVR vs. no SVR SVR=Undetecta ble HCV-RNA > 6 months after completion of antiviral therapy	SVR (560) vs. no SVR (655) Mean age (years): 42 overall Female: 34% vs. 28% Non-White: 10% vs. 6% Genotype 1: 19% vs. 50% Viral load: Not reported Cirrhosis: 10% vs. 18%	Pegylated interferon plus ribavirin: 61% Pegylated interferon monotherapy: 1% Interferon plus ribavirin: 21% Interferon monotherapy: 18%	SVR vs. no SVR Liver-related mortality: Adjusted HR 0.22 (0.09- 0.58) Liver-related hospital episode: Adjusted HR 0.22 (0.15-0.34)

Author Country N Quality	Comparison Definition of Sustained Virologic Response	Population Characteristics	Treatments	Results
Izumi 2005 ⁹⁹ Japan N=495 Quality: Fair	SVR vs. no SVR SVR=Undetecta ble HCV-RNA 6 months after completion of antiviral therapy	Demographics for patients treated with interferon monotherapy and interferon plus ribavirin combination therapy, respectively (not reported by SVR status) Mean age (years): 52 and 58 Female: 43% and 44% Race: Not reported Genotype 1b: 71% and 80% Median viral load (kIU/mI): 470 and 680 Cirrhosis: 7% and 2%	Interferon monotherapy: 69% Interferon-2b plus ribavirin combination therapy: 34%	SVR vs. no SVR HCC: Adjusted HR 0.36 (0.04-0.83)
Kasahara 2004 ¹⁰⁰ Japan N=2,698 Quality: Poor	SVR vs. no SVR SVR=Undetecta ble HCV-RNA 6 months after completion of antiviral therapy	SVR (n=738) vs. no-SVR (n=1930) Median age (years): 51 vs. 54 (p=0.12) Female: 31% vs. 37% (p=0.32) Race: Not reported Genotype 1: Not reported Viral load: Not reported Cirrhosis (Desmet F4): 3.0% vs. 5.4% (p=0.34)	Interferon	SVR vs. no SVR Liver-related mortality: Adjusted HR 0.04 (0.005- 0.30) All-cause mortality: Adjusted HR 0.14 (0.06-0.35)
Maruoka 2012 ¹⁰¹ Japan N=577 Quality: Fair	SVR vs. no SVR SVR=Undetecta ble HCV-RNA >6 months after completion of antiviral therapy	For all treated patients (not reported by SVR status) Mean age (years): 50 Female: 36% Non-White: Not reported Genotype 1: 73% Viral load high (≥100 KIU, 100 kc, 1.0 Meq, 10⁴/50 microL, or 30 core antigens): 69% Cirrhosis: 10%	Interferon-alfa or -beta monotherapy: 83% Interferon-alfa or -beta sequential therapy: 3.3% Interferon-alfa plus ribavirin combination therapy: 14%	SVR vs. no SVR ^a All-cause mortality: Adjusted HR 0.20 (0.08-0.54) HCC: Adjusted HR: 0.12 (0.04-0.40)

Table 11. Sustained virologic response and clinical outcomes (continued)					
Author Country N Quality	Comparison Definition of Sustained Virologic Response	Population Characteristics	Treatments	Results	
Morgan, 2010 ⁹ U.S. Name: HALT-C N=526 Quality: Fair	SVR vs. no SVR SVR=Undetecta ble HCV-RNA 6 months after completion of antiviral therapy	SVR (n=140) vs. breakthrough/relapse (n=77) vs. no SVR (n=309) Mean age (years): 49 vs. 49 vs. 50 (p=0.23) Female: 24% vs. 26% vs. 30% (p=0.30) Non-White: 20% vs. 20% vs. 32% (p=0.001) Genotype 1: 72% vs. 86% vs. 94% (p<0.0001) Viral load: Not reported Cirrhosis (Ishak 5 or 6): 21% vs. 31% vs. 43% (p<0.0001)	Pegylated interferon-2a- 180 µg/week + ribavirin 1000-12000 mg/day for 24weeks	SVR vs. no SVR All-cause mortality or liver transplantation: Adjusted HR 0.17 (0.06-0.46) Any liver-related outcome (decompensated liver disease [ascites, variceal bleeding, hepatic encephalopathy, spontaneous bacterial peritonitis], HCC, liver transplantation, liver-related mortality): Adjusted HR 0.15 (0.06-0.38) Decompensated liver disease: Adjusted HR 0.13 (0.03-0.53) HCC: Adjusted HR 0.19 (0.04-0.80) Liver-related mortality or liver transplantation: Adjusted HR 0.12 (0.03-0.48)	
Shiratori, 2005 ¹⁰² Japan N=271 Quality: Poor	SVR vs. no SVR SVR=Undetecta ble HCV-RNA 6 months after completion of antiviral therapy	For all treated patients (not reported by SVR status) Mean age (years): 57 Female: 62% Race: Not reported Genotype 1: 75% Viral load (log ₁₀ copies/ml): 5.8 Cirrhosis: 100% (inclusion criterion)	Interferon alfa-2a: 58% Natural interferon alfa: 42%	SVR vs. no SVR ^a HCC: Adjusted HR 0.40 (0.18-0.89) All-cause mortality: Adjusted HR 0.07 (0.01-0.56)	
Veldt, 2007 ¹⁰³ Europe and Canada N=479 Quality: Poor	SVR vs. no SVR SVR=Undetecta ble HCV-RNA 6 months after completion of antiviral therapy	SVR (n=142) vs. no-SVR (n=337) Mean age (years): 48 vs. 49 (p=0.45) Female: 27% vs. 32% (p=0.23) Race: Not reported Genotype 1: 39% vs. 67% (p<0.001) Viral load (x10 ⁵ IU/mL): 8.5 vs. 8.0 (p=0.75) Cirrhosis (Ishak 5 or 6): 71% vs. 77% (p=0.45)	Interferon monotherapy: 27% Interferon and ribavirin: 27% Pegylated interferon monotherapy: 2.1% Pegylated interferon and ribavirin: 43%	SVR vs. no SVR Any event (death, liver failure, and HCC): Adjusted HR 0.20 (0.07-0.58) All-cause mortality: Adjusted HR 0.31 (0.07-1.4) Liver-related mortality: Adjusted HR 0.19 (0.02-1.4) HCC: Adjusted HR 0.46 (0.12-1.70)	
Yoshida, 2002 ¹⁰⁴ Japan N=459 Quality: Poor	SVR vs. no SVR SVR=Undetecta ble HCV-RNA 6 months after completion of antiviral therapy	SVR (817) vs. non-SVR (1613) Mean age (years): 48 vs. 51 Female: 30% vs. 40% Race: Not reported Genotype: Not reported Viral load: Not reported Cirrhosis (Desmet F4): 6.5% vs. 11%	Interferon- alfa: 84% Interferon- beta: 14% Both: 2%	SVR vs. no SVR ^a Liver-related mortality: Adjusted HR 0.13 (0.02- 0.66) All-cause mortality Adjusted HR 0.32 (0.12-0.86)	

Author Country N Quality	Comparison Definition of Sustained Virologic Response	Population Characteristics	Treatments	Results
Yu, 2006 ¹⁰⁵ Taiwan N=1,057 Quality: Poor	SVR vs. no SVR SVR=Undetecta ble HCV-RNA 6 months after completion of antiviral therapy	For all treated patients (not reported by SVR status) Mean age (years): 47 Female: 40% Race: Not reported Genotype 1: 46% Viral load: Not reported Cirrhosis (criteria not reported): 16%	Interferon monotherapy: 28% Interferon plus ribavirin combination therapy: 72%	SVR vs. no SVR ^a All-cause mortality: Adjusted HR 0.28 (0.08-1.0) HCC: Adjusted HR 0.25 (0.13-0.50)

HCV = hepatitis C virus; HR = hazard ratio; NA = not applicable; HCC = hepatocellular carcinoma; HCV-RNA = hepatitis C virus-ribonucleic acid; SVR = sustained virologic response

Quality of Life

Nine cohort studies evaluated the association between an SVR following interferon-based antiviral therapy and outcomes related to quality of life (Appendix H: Evidence Table 12). ¹⁰⁷⁻¹¹⁵ Sample sizes ranged from 138 to 1121. Only one study ¹⁰⁷ reported adjusted risk estimates, thus we included studies that reported unadjusted risk estimates. Eight studies ^{107, 108, 110-115} evaluated patients originally enrolled in randomized trials ^{27, 41, 116-119} of antiviral treatments. Two studies evaluated the same cohort of patients ^{113, 115} and one study ¹¹⁴ evaluated a cohort of patients included in a study ¹⁰⁸ that reported results for three pooled cohorts. One study included patients randomized to pegylated interferon (alfa-2a or alfa-2b) plus ribavirin, although results were not stratified according to what type of antiviral therapy was received. ¹¹¹ The remainder of the studies evaluated nonpegylated interferon plus ribavirin combination therapy, or nonpegylated or pegylated interferon monotherapy.

All studies were rated poor quality (Appendix H: Evidence Table 13). One study adjusted for potential confounders, ¹⁰⁷ one study reported low loss to followup, ¹¹² and one study reported blinding of patients to virologic outcomes. ¹¹⁰ Followup was at 24 weeks after treatment (typically 72 weeks from start of treatment) in all studies. No study evaluated longer term quality of life according to SVR status.

All of the studies found patients with an SVR experienced better improvement from baseline on individual SF-36 domains as well as SF-36 physical and mental component summary scores compared with those with no SVR (Appendix H: Evidence Table 14). In most studies, differences between patients with and without an SVR on various SF-36 domains were less than 5 to 10 points. Patients with an SVR also reported greater improvements from baseline on hepatitis C specific quality of life measures (health distress and limitations) and measures related to fatigue and sleep somnolence. However, results are subject to the methodological limitations of the studies.

One study also found achieving an overall response (defined as SVR plus 2-point improvement in the Histological Activity Index) associated with improved quality of life compared with those without an overall response. ¹¹³

^a Calculated from estimates of SVR compared with untreated and no SVR compared with untreated.

Discussion

The evidence reviewed in this study is summarized in Table 12. The specific domain scores used to determine the overall strength of evidence for each body of evidence are shown in Appendix G.

Antiviral therapy for chronic hepatitis C virus (HCV) infection continues to evolve. No study has evaluated comparative effectiveness of current antiviral regimens on long-term clinical outcomes such as mortality, complications of chronic HCV infection, or quality of life. Such trials would be difficult to design and carry out due to the long time required for complications of chronic HCV infection to develop in most patients. The first pegylated interferon was approved by the FDA only in 2001, and the initial major trials of pegylated interferon plus ribavirin were published in 2002. The protease inhibitors telaprevir and boceprevir were approved only in 2011. Although some trials reported short-term (prior to 1 year after the end of antiviral therapy) mortality, ^{22, 32, 51} few adverse events were reported, precluding reliable conclusions.

Dual Therapy Regimens With Pegylated Interferon and Ribavirin

In lieu of direct evidence on long-term clinical outcomes, sustained virologic response (SVR) rates are the primary outcome to assess comparative benefits of different antiviral regimens. In trials of treatment-naïve patients, the likelihood of achieving an SVR was slightly lower with dual therapy with pegylated interferon alfa-2b plus ribavirin compared with dual therapy with pegylated interferon alfa-2a plus ribavirin (pooled RR 0.87, 95% CI 0.80 to 0.95; I²=27%), with a difference in absolute SVR rates of about 8 percentage points. 20-23, 53, 55, 58 Although the largest study, the IDEAL trial, found no difference in SVR rates between dual therapy with pegylated interferon alfa-2a compared with dual therapy with pegylated interferon alfa-2b, excluding the IDEAL trial from pooled analyses resulted in similar effect estimates. 22 Although there was no difference between dual therapy regimens in risk of withdrawals due to adverse events, dual therapy with pegylated interferon alfa-2b plus ribavirin was associated with a lower risk of serious adverse events than dual therapy with pegylated interferon alfa-2a plus ribavirin (pooled RR 0.76, 95% CI 0. 61 to 0.95, $I^2=0\%$), suggesting a potential tradeoff between greater benefits and harms. However, serious adverse events were only reported in two trials;^{22, 23} the rate of serious adverse events was relatively low (about 4 percent overall in IDEAL), with an absolute difference of about one percent; and adverse events with antiviral treatments generally resolve following discontinuation of therapy.

Trials found no clear differences in estimates of relative effectiveness of dual therapy with pegylated interferon alfa-2a compared with dual therapy with pegylated interferon alfa-2b in patient subgroups stratified by age, sex, race, viral load, fibrosis stage, and genotype, although absolute response rates were lower in older patients, Black patients, patients with high viral load, patients with more advanced fibrosis or cirrhosis, and genotype 1 infection. SVR rates ranged from 24–42 percent lower in patients with genotype 1 infection compared with patients with genotype 2 or 3.

In patients with genotype 2 or 3 infection, dual therapy for 12 to 16 weeks appears to be associated with lower likelihood of SVR compared with dual therapy for 24 weeks, with no differences between 24 weeks and longer courses of therapy. Standard doses of pegylated interferon alfa-2b were more effective than lower doses (no trials compared different

doses of pegylated interferon alfa-2a).^{66, 73-77} Although trials comparing different ribavirin doses found no clear differences, with the exception of one trial that found lower doses associated with lower SVR rates in patients with advanced fibrosis,⁸¹ they evaluated different dose comparisons, precluding firm conclusions.^{63, 80, 82}

Key Question	Outcome	Summary of Evidence	Strength of
Key Question 1a What is the comparative effectiveness of antiviral treatment in improving health outcomes in patients with HCV infection?	Long-term clinical outcomes	No evidence.	Evidence Insufficient
	Short-term mortality	Three trials that compared current antiviral regimens ^a found no differences in risk of short-term mortality, but reported very few (20 total) events.	Low
	Short-term quality of life	One open-label randomized trial of patients with genotype 4 infection found dual therapy with pegylated interferon alfa-2a plus ribavirin associated with statistically significant, slightly better short-term scores on some quality of life assessments compared with dual therapy with pegylated interferon alfa-2b plus ribavirin.	Low
Key Question 1b How does the comparative effectiveness of antiviral treatment for health outcomes vary according to patient subgroup characteristics?	Any clinical outcome	No evidence.	Insufficient
Key Question 2a	Dual Therapy With Pegylated Interferon Alfa-2b Plus Ribavirin vs. Dual Therapy With Pegylated Interferon Alfa-2a Plus Ribavirin		
What is the comparative effectiveness of antiviral treatments on intermediate outcomes?	Sustained virologic response	Seven trials found dual therapy with standard doses of pegylated interferon alfa-2b plus ribavirin associated with lower likelihood of achieving an SVR than pegylated interferon alfa-2a plus ribavirin (pooled RR 0.87, 95% CI 0.80 to 0.95; I ² =27%), with an absolute difference in SVR rates of 8 percentage points (95% CI 3 to 14).	Moderate

(continued) Key Question	Outcome	Summary of Evidence	Strength of Evidence	
	Dual Therapy With Pegylated Interferon Alfa-2a or Alfa-2b Plus Ribavirin: Duration Effects			
	Sustained virologic response	Two trials of patients with genotype 2 or 3 infection found no difference in likelihood of achieving an SVR between 48 vs. 24 weeks of dual therapy with pegylated interferon alfa-2a plus ribavirin (pooled RR 0.97, 95% CI 0.84 to 1.1; I ² =43%).	Moderate	
	Sustained virologic response	Four trials of patients with genotype 2 or 3 infection found 24 weeks of dual therapy with pegylated interferon (alfa-2a or alfa-2b) more effective than 12-16 weeks for achieving an SVR (pooled RR 1.2, 95% Cl 1.0 to 1.3; l²=80%). Relative risk estimates ranged from 1.0 to 1.3 in the four trials and may have varied in part due to differences across studies in ribavirin dosing.	Moderate	
Key Question 2a What is the comparative effectiveness of antiviral	Sustained virologic response	Three trials of patients with genotype 2 or 3 infection with a rapid virologic response (undetectable HCV-RNA by week 4) found no differences between 24 vs. 12-16 weeks of dual therapy with pegylated interferon (alfa-2a or alfa-2b) plus ribavirin (pooled RR 0.99, 95% CI 0.86 to 1.1, I ² =66%). Relative risk estimates ranged from 0.89 to 1.1.	Moderate	
treatments on	Dual Therapy With Pegylated Interferon Alfa-2a or Alfa-2b Plus Ribavirin: Dose Effects			
	Sustained virologic response	Six trials of patients with genotype 2 or 3 infection found lower doses of pegylated interferon alfa-2b (0.75-1.0 mcg/kg or 50 mcg) associated with lower likelihood of achieving an SVR than higher doses (1.5 mcg/kg or 100-150 mcg) (pooled RR 0.90; 95% CI 0.81 to 0.99; I ² =20%).	Moderate	
	Sustained virologic response	Three trials of patients with genotype 2 or 3 infection who did not specifically have advanced fibrosis or cirrhosis found no clear difference in likelihood of SVR between lower doses of ribavirin (400 or 800 mg flat dose or 600 to 800 mg weight-based dose) vs. higher doses (800 or 1,200 mg flat dose or 800 to 1400 mg weight-based dose).	Moderate	
	Sustained virologic response	One small trial of patients with genotype 2 or 3 infection (N=60) and advanced fibrosis or cirrhosis (Ishak stage 4-6) found 600 to 800 mg daily of ribavirin associated with lower likelihood of SVR than 1000 to 1200 mg daily (45 vs. 72 percent, RR 0.62, 95% C I 0.40 to 0.98).	Low	

(continued) Key Question	Outcome	Summary of Evidence	Strength of Evidence	
	Triple Therapy With Pegylated Interferon Alfa-2b, Ribavirin, and Boceprevir vs. Dual Therapy With Pegylated Interferon Alfa-2b Plus Ribavirin			
	Sustained virologic response	Two trials of patients with genotype 1 infection found triple therapy with boceprevir (pegylated interferon alfa-2b plus ribavirin for 4 weeks, followed by the addition of boceprevir for 44 weeks) associated with higher likelihood of SVR than dual therapy with pegylated interferon alfa-2b plus ribavirin therapy for 48 weeks (pooled RR 1.8; 95% CI 1.6 to 2.1; I ² =0%), with an absolute increase in SVR rate of 31% (95% CI 23 to 39).	Moderate	
	Sustained virologic response	One trial of patients with genotype 1 infection found 48 weeks of triple therapy with boceprevir using a low dose of ribavirin (400-1000 mg daily) associated with a non—statistically significant trend toward lower likelihood of SVR compared with 48 weeks of triple therapy with a standard ribavirin dose (800-1400 mg daily) (36% vs. 50%, RR 0.71, 95% CI 0.39 to 1.3).	Low	
		egylated Interferon Alfa-2a or Alfa-2b, Ribavirin, With Pegylated Interferon Alfa-2a or Alfa-2b Plus		
Key Question 2a	vs. Duai Trierapy	Three trials of patients with genotype 1	S KIDAVIIII	
What is the comparative effectiveness of antiviral treatments on intermediate outcomes? (continued)	Sustained virologic response	infection found triple therapy with telaprevir for 24 weeks (12 weeks of pegylated interferon alfa-2a, ribavirin, and telaprevir followed by 12 weeks of pegylated interferon alfa-2a plus ribavirin) associated with a higher likelihood of SVR than dual therapy with pegylated interferon alfa-2a plus ribavirin for 48 weeks (pooled RR 1.5, 95% CI 1.3 to 1.8; I ² =0%), with an absolute increase in SVR rate of 22% (95% CI 13 to 31).	Moderate	
	Sustained virologic response	One trial of patients with genotype 1 infection found no difference in likelihood of SVR between triple therapy with pegylated interferon, ribavirin, and telaprevir for 12 weeks vs. dual therapy with pegylated interferon alfa-2a plus ribavirin for 48 weeks.	Moderate	
	Sustained virologic response	One trial of patients with genotype 1 infection found response-guided triple therapy with telaprevir (pegylated interferon alfa-2a, ribavirin, and telaprevir for 8 or 12 weeks followed by a response-guided dual therapy with pegylated interferon alfa-2a plus ribavirin for an additional 12 or 36 weeks) associated with a higher likelihood of SVR than dual therapy with pegylated interferon alfa-2a plus ribavirin for 48 weeks (RR 1.6, 95% CI 1.4 to 1.9), with an absolute increase in SVR rate ranging from 25% to 31%. The regimen with 8 weeks of telaprevir was associated with a slightly lower SVR rate than the 12 week telaprevir regimen (69% vs. 75%).	Low	

(continued)	T	т		
Key Question	Outcome	Summary of Evidence	Strength of Evidence	
	Triple Therapy With Pegylated Interferon Alfa-2a or Alfa-2b, Ribavirin, and Telaprevir vs. Dual Therapy With Pegylated Interferon Alfa-2a or Alfa-2b			
Key Question 2a What is the comparative effectiveness of antiviral treatments on	Sustained virologic response	Plus Ribavirin (continued) One trial of patients with genotype 1 infection found no difference in likelihood of SVR between triple therapy with telaprevir for 48 weeks (12 weeks of triple therapy with pegylated interferon alfa-2a, ribavirin, and telaprevir followed by 36 weeks of dual therapy with pegylated interferon alfa-2a plus ribavirin) vs. triple therapy with telaprevir for 24 weeks (12 weeks of triple therapy followed by 12 weeks of dual therapy).	Low	
intermediate		th Pegylated Interferon Alfa-2a, Ribavirin, and		
outcomes?	Dose Effects of Pe	gylated Interferon Alfa-2a vs. Alfa-2b and Durat	ion Effects	
(continued)	Sustained virologic response	One trial of response-guided triple therapy with telaprevir (24 or 48 weeks, based on absence or presence of HCV-RNA from weeks 4 through 20) found similar SVR rates (81–85%) for regimens that varied on telaprevir dose (750 mg tid vs. 1125 mg bid) and type of pegylated interferon (alfa-2a or alfa-2b).	Low	
	Sustained virologic response	One trial of patients with an extended rapid virologic response to initial triple therapy with telaprevir reported similar, high (92% and 88%) SVR rates in patients randomized to a total of 24 or 48 weeks of therapy.	Low	
	Dual Therapy With Pegylated Interferon Alfa-2b Plus Ribavirin vs. Dual Therapy With			
Key Question 2b How does the comparative effectiveness of antiviral treatment for intermediate outcomes vary	Sustained virologic response	The largest randomized trial (n=3070) of dual therapy with pegylated interferon alfa-2a plus ribavirin vs. dual therapy with pegylated interferon alfa-2b plus ribavirin found no clear differences in relative risk estimates for SVR in genotype 1 patients stratified by race, sex, age, baseline fibrosis stage, or baseline viral load. Characteristics associated with lower absolute SVR rates across dual therapy regimens were older age, Black race, advanced fibrosis or cirrhosis, and high baseline viral load.	Low	
according to patient subgroup characteristics?	Sustained virologic response	Four randomized trials of dual therapy with pegylated interferon alfa-2a plus ribavirin vs. dual therapy with pegylated interferon alfa-2b plus ribavirin found no clear differences in relative risk estimates for SVR in patients stratified by genotype. Genotype 1 infection was associated with a lower absolute SVR rate than genotypes 2 or 3.	Moderate	

(continued) Key Question	Outcome	Summary of Evidence	Strength of Evidence
Key Question 2b How does the comparative effectiveness of antiviral treatment for	Triple Therapy With Pegylated Interferon Alfa-2b, Ribavirin, and Boceprevir vs. Dual Therapy With Pegylated Interferon Alfa-2b Plus Ribavirin		
	Sustained virologic response	Two trials of triple therapy with boceprevir for 48 weeks (4 weeks of dual therapy lead-in with pegylated interferon plus ribavirin followed by 44 weeks of triple therapy with pegylated interferon, ribavirin, and boceprevir) found no difference in relative risk estimates for SVR in men vs. women, and no clear difference in relative risk estimates for Black vs. non-Black patients. Black race was associated with a lower absolute SVR rate than non-Black race.	Moderate
	Sustained virologic response	Two trials found triple therapy with pegylated interferon alfa-2b, ribavirin, and boceprevir associated with higher likelihood of achieving SVR than dual therapy with pegylated interferon alfa-2b plus ribavirin in patients with high baseline HCV-RNA viral load (>600,000 or ≥800,000 IU/mL), but found no difference in likelihood of SVR in patients with lower viral load.	Moderate
intermediate	Triple Therapy With Pegylated Interferon Alfa-2a or Alfa-2b, Ribavirin, and Telaprevir vs. Dual Therapy With Pegylated Interferon Alfa-2a or Alfa-2b Plus Ribavirin		
according to patient subgroup characteristics? Sustained virologic response Sustained virologic response Sustained virologic response Sustained virologic response Sustained virologic response	One trial of response-guided triple therapy with telaprevir (12 weeks of pegylated interferon alfa-2a, ribavirin, and telaprevir followed by response-guided dual therapy with pegylated interferon alfa-2a and ribavirin) vs. dual therapy with pegylated interferon plus ribavirin for 48 weeks found no clear differences in relative risk estimates in patients stratified by age, sex, race, baseline fibrosis status, or body mass index. Characteristics associated with lower absolute rates of SVR were older age, Black race, advanced fibrosis or cirrhosis, and higher body mass index. One other trial of 24-week fixed duration triple therapy with telaprevir, pegylated interferon alfa-2b, and ribavirin vs. 48 weeks of dual therapy found no differences in estimates of effect in patients stratified by sex or age.	Moderate (for age and sex) Low (for other factors)	
	•	Two trials of triple therapy with pegylated interferon (alfa-2a or alfa-2b), ribavirin, and telaprevir vs. dual therapy depending reported inconsistent findings for differential relative risk estimates according baseline viral load.	Insufficient

(continued) Key Question	Outcome	Summary of Evidence	Strength of Evidence
	Dual Therapy With Pegylated Interferon Alfa-2b Plus Ribavirin vs. Dual Therapy With Pegylated Interferon Alfa-2a Plus Ribavirin		
	Harms	Dual therapy with pegylated interferon alfa-2b was associated with slightly greater risk of headache (three trials, pooled RR 1.1, 95% Cl 1.1 to 1.2, l ² =0%), and a lower risk of serious adverse events (two trials, pooled RR 0.76; 95% Cl 0.71 to 0.88; l ² =0%), lower risk of neutropenia (five trials, pooled RR 0.61, 95% Cl 0.46 to 0.83, l ² =38%), and lower risk of rash (two trials, pooled RR 0.79, 95% Cl 0.71 to 0.88, l ² =0%) than dual therapy with pegylated interferon alfa-2a plus ribavirin, with no differences in withdrawals due to adverse events.	Moderate
		egylated Interferon Alfa-2b, Ribavirin, and Boce With Pegylated Interferon Alfa-2b Plus Ribavirii	
Key Question 3a What are the comparative harms associated with antiviral treatments?	Harms	Triple therapy with boceprevir for 48 weeks (pegylated interferon alfa-2b plus ribavirin for 4 weeks followed by addition of boceprevir for 44 weeks) was associated with increased risk of neutropenia (two trials, pooled RR 1.8, 95% CI 1.5 to 2.3, I²=0%), dysgeusia (two trials, pooled RR 2.5, 95% CI 2.0 to 3.2, I²=0%), anemia (two trials, pooled RR 2.0, 95% CI 1.4 to 2.8, I²=0%), and thrombocytopenia (two trials, pooled RR 3.2, 95% CI 1.2 to 8.2; I²=0%) than dual therapy with pegylated interferon alfa-2b plus ribavirin. The incidence of anemia was about 25% with triple therapy and the incidence of neutropenia about 33%, with severe anemia in 4–5% and severe neutropenia in 8–15%.	Moderate
		gylated Interferon Alfa-2a or Alfa-2b, Ribavirin, With Pegylated Interferon Alfa-2a or Alfa-2b Plus In two trials, there were no statistically significant differences between a 12-week regimen of triple therapy with pegylated	
		interferon alfa-2a, ribavirin, and telaprevir vs. dual therapy with pegylated interferon alfa-2a plus ribavirin in risk of any assessed adverse event.	

(continued)			Strength of
Key Question	Outcome	Summary of Evidence	Evidence
		With Pegylated Interferon Alfa-2a or Alfa-2b, Ri	
	and Telaprevir vs.	Dual Therapy With Pegylated Interferon Alfa-2a	or Alfa-2b
		Plus Ribavirin (continued) In three trials, a 24-week regimen of triple	
		therapy with telaprevir (pegylated interferon	
		alfa-2a or alfa-2b, ribavirin, and telaprevir for	
Kay Ouastian 2a		12 weeks followed by pegylated interferon alfa-	
Key Question 3a What are the		2a plus ribavirin for 12 weeks) was associated	
comparative		with increased risk of anemia (three trials,	
harms associated		pooled RR 1.3, 95% CI 1.1 to 1.5, I ² =0%) and	
with antiviral	Hormo	rash (three trials, pooled RR 1.4, 95% CI 1.1 to	Madarata
treatments?	Harms	1.7; l ² =0%) vs. dual therapy with pegylated interferon alfa-2a plus ribavirin for 48 weeks.	Moderate
(continued)		Among patients randomized to the 24-week	
		telaprevir regimen, one to two-thirds	
		experienced a rash (7–10% experienced	
		severe rash) and 27–91% experienced anemia	
		(4–11% experienced severe anemia). There	
		was no difference in risk of withdrawal due to	
		adverse events.	
		In one trial, response-guided triple therapy with telaprevir (pegylated interferon alfa-2a,	
		ribavirin, and telaprevir for 8 or 12 weeks	
		followed by response-guided duration	
		pegylated interferon alfa-2a and ribavirin) was	
		associated with increased risk of withdrawal	
	Harms	due to adverse events (27% vs. 7.2%, RR 3.8,	Low
		95% CI 2.6 to 5.7), anemia (38% vs. 19%, RR	
		2.0, 95% Cl 1.6 to 2.5), any rash (36% vs.	
		24%, RR 1.5, 95% CI 1.2 to 1.8), and severe rash (5% vs. 1%, RR 4.6, 95% CI 1.6 to 13) vs.	
		therapy with pegylated interferon alfa-2a plus	
		ribavirin for 48 weeks.	
		ylated Interferon Alfa-2b Plus Ribavirin vs. Dua	l Therapy With
	Pe	egylated Interferon Alfa-2a Plus Ribavirin	T
		No trial of dual therapy with pegylated	
		interferon alfa-2b plus ribavirin vs. dual therapy with pegylated interferon alfa-2a plus ribavirin	
		reported harms in patients stratified by factors	
		such as HCV genotype, age, race, sex, stage	
Key Question 3b	Harms	of disease, or genetic markers.	Insufficient
Do these harms		Three trials that restricted enrollment to	
differ according		patients with genotype 1 infection reported risk	
to patient		estimates for risk of harms that were similar to	
subgroup characteristics?	Triple Theren:	the risk estimates based on all trials.	havirin
CHALACIEHISHUS!		With Pegylated Interferon Alfa-2a or Alfa-2b, Ri evir or Boceprevir vs. Dual Therapy With Pegyla	
		terferon Alfa-2a or Alfa-2b Plus Ribavirin	
		No trial evaluated harms associated with triple	
		therapy with pegylated interferon, ribavirin, and	
	Harms	boceprevir or telaprevir vs. dual therapy with	Insufficient
	- I GITTIO	pegylated interferon plus ribavirin in patient	oamoion
		subgroups. All trials evaluated patients with	
		genotype 1 infection.	

(continued)

Key Question	Outcome	Summary of Evidence	Strength of Evidence
Key Question 4 Have improvements in intermediate outcomes been shown to reduce the risk or rates of adverse health outcomes from HCV infection?	Mortality and long-term hepatic complications	A large VA hospital study that controlled well for potential confounders found an SVR after antiviral therapy associated with lower risk of all-cause mortality vs. no SVR (adjusted HR 0.71 [0.60-0.86], 0.62 [0.44-0.87] and 0.51 [0.35-0.75] for genotypes 1, 2, and 3, respectively). Eighteen other cohort studies found an SVR associated with decreased risk of all-cause mortality, liver-related mortality, HCC, and other complications of ESLD compared with no SVR, with stronger effect estimates than the VA study (adjusted HRs generally ranged from around 0.10 to 0.33). However, the studies had methodological shortcomings, including inadequate handling of confounders, and 10 were conducted in Asia.	Moderate
	Short-term quality of life	Nine studies found an SVR associated with greater improvement in measures related to quality of life (generic or disease-specific) 24 weeks after the end of antiviral treatment vs. no SVR, with differences averaging less than 5 to 10 points on various SF-36 domains. All studies were poor-quality and were characterized by failure to adjust for confounders, high loss to followup, and failure to blind patients to SVR status.	Low

CI = confidence interval; ESLD = end-stage liver disease; HCC = hepatocellular carcinoma; HCV = hepatitis C virus; HCV-RNA = hepatitis C virus ribonucleic acid; HR = hazard ratio; I²=index measures the extent of true heterogeneity in a metaanalysis, RR = relative risk; SVR = sustained virologic response

Triple Therapy Regimens With Pegylated Interferon, Ribavirin, and Either Boceprevir or Telaprevir

The relatively low SVR rates with pegylated interferon plus ribavirin dual therapy for genotype 1 infection (present in about three-quarters of U.S. patients with HCV infection) has led to ongoing efforts to identify more effective treatment alternatives. Recent trials found triple therapy regimens with pegylated interferon (alfa-2a or alfa-2b), ribavirin, and either boceprevir or telaprevir associated with substantially higher SVR rates than standard dual therapy with pegylated interferon (alfa-2a or alfa-2b) plus ribavirin in treatment-naïve patients with genotype 1 infection. ^{30-32, 51, 59, 85-87}SVR rates with triple therapy approached the 70–80 percent rates observed with dual therapy in patients with genotype 2 or 3 infection. Trials that evaluated the telaprevir regimen recommended by the FDA (12 weeks of triple therapy with telaprevir followed by response-guided duration of 12 or 36 weeks of dual therapy) reported SVR rates of 75–80 percent. 51,59 Trials that evaluated the FDA-recommended boceprevir regimen for antiviral-naïve patients with cirrhosis (4 weeks of dual therapy lead-in followed by 44 weeks of triple therapy with boceprevir) reported SVR rates of 66–75 percent. 30, 32 Trials that evaluated other regimens in antiviral naïve patients, including fixed duration telaprevir regimens, shorter fixed duration triple therapy with boceprevir, and boceprevir without dual therapy lead-in,

a "Current antiviral treatment regimen" refers to dual therapy with pegylated interferon (alfa-2a or alfa-2b) plus ribavirin, or triple therapy with pegylated interferon (alfa-2a or alfa-2b) plus ribavirin and boceprevir or telaprevir.

reported similar or lower SVR rates. The SVR rates with various antiviral regimens or placebo are summarized in Table 13.

Table 13. Sustained virologic response rates with different antiviral regimens for hepatitis C virus infection

Regimen	Sustained Virologic Response Rate 6 Months after Treatment (%)	Approximate Number Needed To Treat To Achieve One Sustained Virologic Response Compared With Placebo	References
Placebo	<2	Not applicable	Poynard et al., 1996 ¹²
Interferon monotherapy	6-16	7-25	Chander, 2002 ¹⁰ Kjaegard, 2000 ¹¹ Poynard, 1996 ¹² Shepherd., 2000 ¹³
Interferon plus ribavirin	33-41	2.6-3.2	Chander, 2002 ¹⁰ Kjaergard, 2001 ¹¹ Shepherd, 2000 ¹³
Pegylated interferon plus ribavirin	54-61 overall, 42-52 in patients with genotype 1 infection	1.7-1.9 overall; 2.0-2.5 for genotype 1 infection	Shepherd, 2005 ¹²⁰ Sieber, 2005 ¹²¹ Zaman, 2003 ¹²²
Pegylated interferon plus ribavirin plus boceprevir or telaprevir ^a	66-80 (genotype 1 infection only)	1.3-1.6 (genotype 1 infection only)	Jacobson, 2011 ⁵¹ Kwo, 2010 ³⁰ Marcellin, 2011 ⁵⁹ Poordad, 2011 ³²

^a Based on regimens recommended by the U.S. Food and Drug Administration evaluated in trials of antiviral-naïve patients.

The boceprevir regimen recommended by the FDA⁸⁴ for antiviral-naïve patients without baseline cirrhosis (4 weeks of dual therapy lead-in with pegylated interferon plus ribavirin followed by the addition of boceprevir for 24 weeks for virologic responders at weeks 8 to 24, or 4 weeks of dual therapy lead-in followed by the addition of boceprevir for 32 weeks and then 12 additional weeks of dual therapy for late virologic responders) has not been evaluated in a trial of antiviral-naïve patients. Rather, the FDA recommendation was based on a trial of previous partial responders to pegylated interferon plus ribavirin which found slightly higher SVR rates in late virologic responders who received 32 weeks of triple therapy followed by 12 weeks of dual therapy versus those who received 44 weeks of triple therapy, each following 4 weeks of dual therapy lead-in (SVR rates 79 vs. 73 percent).¹²³

As in the head-to-head trials of dual therapy with pegylated interferon alfa-2a plus ribavirin versus pegylated interferon alfa-2b plus ribavirin, relative risk estimates were similar (or there was no clear difference) in patient subgroups based on age, sex, and race, although absolute SVR rates were lower in older patients and Black patients. Triple therapy with either boceprevir or telaprevir was no more effective than dual therapy in the subgroup of patients with lower HCV-RNA viral load (<600,000 or <800,000 IU/mL). There was insufficient evidence to evaluate relative effectiveness of triple compared with dual therapy based on fibrosis stage.

In addition to higher likelihood of SVR, another advantage of triple therapy regimens in patients with genotype 1 infection is the potential for shorter duration (24 or 28 weeks in patients with early virologic response compared with the standard 48 weeks of dual therapy with pegylated interferon plus ribavirin). Shorter courses of treatment would probably be appealing to patients of high relevance to patients, given the high frequency of bothersome flulike symptoms associated with interferon-based therapy. Triple therapy regimens were associated with increased risk of certain harms, in particular hematological adverse events (neutropenia, anemia, and thrombocytopenia) with boceprevir and anemia and rash (including severe rash in <10 percent of

patients that could result in treatment discontinuation) with telaprevir. However, there was no clear increase in risk of serious adverse events with use of protease inhibitors, and the adverse events appear to be self-limited following drug discontinuation.

Sustained Virologic Response After Antiviral Therapy and Clinical Outcomes

The strongest evidence on the association between an SVR after antiviral therapy and improved clinical outcomes is a large VA cohort study (n=16,864) that adjusted for many confounders. The VA study found decreased risk of all-cause mortality in patients who achieved an SVR compared with those who didn't achieve an SVR across groups stratified by genotype (adjusted HR 0.71 [0.60–0.86], 0.62 [0.44–0.87] and 0.51 [0.35–0.75] for genotypes 1, 2, and 3, respectively). Despite controlling for important confounders, the possibility of residual confounding is suggested by the very rapid separation of mortality curves for patients with an SVR versus those without an SVR, which was observed at three months after assessment for SVR. This is more rapid than expected given the typically prolonged natural history of HCV infection. Therefore, estimates of effects of SVR on clinical outcomes from this study may be exaggerated, though it is not possible to determine to what degree.

Eighteen other cohort studies also found an SVR after antiviral therapy associated with decreased risk of all-cause mortality and complications of chronic HCV infection, including studies specifically of patients with baseline cirrhosis, but had more methodological shortcomings. In addition, 10 of the 19 studies were conducted in Asia, where the incidence of HCC in patients with chronic HCV infection is higher than in the United States, ⁴⁶ potentially limiting their generalizability. Other studies found an SVR after antiviral therapy associated with better scores on various measures of quality of life than no SVR, but those studies focused on short-term outcomes, and typically did not adjust for confounders or blind patients to SVR status when assessing outcomes.

Findings in Relationship to What Is Already Known

Our findings regarding the comparative effectiveness of dual therapy with pegylated interferon alfa-2b plus ribavirin compared with dual therapy with pegylated interferon alfa-2a plus ribavirin are consistent with recent systematic reviews that also found the former associated with a lower likelihood of SVR. ^{18, 124} Our findings of no clear difference in comparative effectiveness between 12 to 16 weeks compared with 24 weeks of response-guided dual therapy with pegylated interferon plus ribavirin in hepatitis C genotype 2 or 3 infection with rapid virologic response are discordant with a recent systematic review, which found a shorter duration of treatment associated with a lower likelihood of achieving an SVR. ¹²⁵ The discrepancy may be explained by the inclusion in the other systematic review of a study that we excluded because it evaluated a nonstandard dose of pegylated interferon, ⁶⁵ as well as its inclusion of subgroup analyses from trials of patients randomized to different fixed durations of therapy prior to assessment of rapid virologic response, ^{64, 68, 70} which we considered separately because they did not represent randomized comparisons of response-guided treatment.

Because telaprevir and boceprevir are so new, we are unaware of other published systematic reviews on the comparative benefits and harms of regimens including these drugs, compared with standard dual therapy. Our findings on the association between achieving an SVR and

reduced risk of mortality or complications associated with chronic HCV infection are consistent with a recent review that used some systematic methods. 126

Applicability

The trials included in this review generally met criteria for efficacy studies, based on the exclusion of patients with common comorbidities (such as serious psychiatric conditions or recent or ongoing substance abuse) who may receive treatments in clinical practice. In addition, the trials may have overestimated efficacy compared with what would be seen in typical practice due to improved adherence as a result of closer followup, effects of trial participation, selection of patients, or other factors. A separate review funded by AHRQ will focus on issues related to adherence in the treatment of HCV infection. ¹²⁷

The severity of baseline liver disease in the patients enrolled in the trials suggests that they enrolled a broad range of patients. In trials of triple therapy with boceprevir or telaprevir, the proportion of patients with cirrhosis at enrollment ranged from <1–11 percent. ^{30-32, 51, 59, 85, 87} Trials that reported the proportion of patients with minimal or no fibrosis reported rates of 27–39 percent. ^{31, 51, 59, 87}

Evidence to evaluate potential differences in comparative benefits or harms in patient subgroups based on age, sex, race, and other clinical factors was relatively limited, precluding strong conclusions in these specific subgroups. The strongest evidence on the association between an SVR versus no SVR after antiviral therapy and reduced mortality comes from a study performed in a VA population, which might limit generalizability to other settings. As described above, studies conducted in Asia on the association between an SVR after antiviral therapy and risk of clinical outcomes may be of limited applicability to U.S. populations because of a higher incidence of HCC in Asian patients with chronic HCV infection. However, HCC incidence is increasing in the United States in HCV-infected patients, which may attenuate such concerns regarding applicability.

The results of this CER are not applicable to populations excluded from the review, including patients previously treated with antiviral therapies and excluded populations such as patients with HIV coinfection, post-transplant patients, or hemodialysis patients. Antiviral therapy is not recommended in patients following kidney transplant, and ribavirin is not recommended in those with more severe (stage 3 to 5) kidney disease since it is cleared via renal function and associated with increased risk of hemolytic anemia in this setting. Such patients were typically excluded from randomized trials of antiviral treatment.¹⁵

Implications for Clinical and Policy Decisionmaking

Our review has potential implications for clinical and policy decisionmaking. For patients with genotype 1 infection, triple therapy regimens with pegylated interferon (alfa-2a or alfa-2b), ribavirin, and telaprevir or boceprevir may be considered an alternative to dual therapy with pegylated interferon alfa-2a or alfa-2b plus ribavirin as standard treatment due to substantially superior efficacy for achieving SVR compared with dual therapy with pegylated interferon alfa-2a or alfa-2b, as well as a shorter duration of treatment. Factors that may affect decisions to utilize regimens with boceprevir or telaprevir include cost and specific harms associated with use of these drugs (such as hematologic adverse events with boceprevir and anemia and rash with telaprevir). Dual therapy with pegylated interferon alfa-2a plus ribavirin appears to be associated with higher likelihood of achieving SVR compared with dual therapy with pegylated interferon alfa-2b plus ribavirin, but absolute differences were relatively small and may be offset in part by

a small increase in serious (but generally self-limited) adverse events. Therefore, decisions about which pegylated interferon to use may be affected by other considerations, such as cost, patient preferences, or other factors. For genotype 2 or 3 infection, standard doses and duration (24 weeks) of pegylated interferon as part of dual therapy are more effective than shorter regimens or lower doses, lending support to dosing guidance from the FDA and clinical practice guidelines. ^{15, 33, 34} Evidence on differential effects of ribavirin dose are too limited to draw strong conclusions about optimal dosing of this component of antiviral regimens, though differences appeared relatively small.

The findings that absolute SVR rates are lower in certain subgroups (such as older patients, Black patients, patients with worse baseline fibrosis, and patients with high viral load) can be used to inform individualized decisionmaking. Patients who are less likely to achieve a SVR may make different informed decisions about therapy compared to those more likely to achieve an SVR, given the adverse effects associated with treatment.

The findings of the review are also relevant to screening recommendations, which are based in part on the effectiveness of treatments in patients found through screening to have HCV infection. Important new evidence that may affect assessments regarding potential benefits of screening include stronger evidence on the link between achieving an SVR and improvement in clinical outcomes, as well as evidence showing substantially higher SVR rates with newer triple therapy regimens with boceprevir or telaprevir in patients with genotype 1 infection, the predominant type of HCV infection in the United States.

Limitations of the Comparative Effectiveness Review Process

Our review had some potential limitations. We excluded non–English-language articles, which could result in language bias, although a recent systematic review found little empirical evidence that exclusion of non–English-language articles leads to biased estimates for noncomplementary or alternative medicine interventions. ¹²⁹

We did not formally assess for publication bias with funnel plots due to small numbers (<10) of studies for all comparisons. Small numbers of studies can make interpretation of funnel plots unreliable, and experts suggest 10 studies as the minimum number of studies to perform funnel plots. ⁵⁰ We included some studies which were published only as abstracts and found that their inclusion or exclusion from analyses did not change conclusions. In addition, we searched trial registries and solicited drug manufacturers for additional unpublished trials and identified none.

Another potential limitation is that we included cohort studies to evaluate the association between SVR and mortality or hepatic complications associated with chronic HCV infection. Such studies are susceptible to confounding if factors associated with SVR (such as age, race, viral load, or fibrosis stage) are also associated with these outcomes. Therefore, we only included studies that reported adjusted risk estimates, and we evaluated how well studies addressed key potential confounders as part of our quality assessment. Nonetheless, residual confounding is a possibility even in cohort studies that adjust for potential confounding.

Limitations of the Evidence Base

We identified several important limitations of the evidence base. First, studies assessing important long-term clinical outcomes associated with current antiviral treatments for chronic HCV infection are not available. In the case of antiviral regimens involving newly approved

antiviral drugs, such studies are not possible yet because of the extended followup required to adequately evaluate effects on clinical outcomes. Second, no trials directly compared regimens with boceprevir compared with regimens with telaprevir. Given the increased efficacy of these regimens in patients with genotype 1 infection, trials directly comparing their effects would be helpful for informing treatment choices between these drugs. In addition, few trials have evaluated the specifically FDA-approved regimens for these drugs, limiting confidence in conclusions regarding estimates of benefits and harms for the regimens likely to be used in clinical practice. Third, almost all of the randomized trials were funded by pharmaceutical companies. Studies have shown that such studies tend to report more favorable results for drugs produced by the funder than studies funded by governmental or other sources. [130, 131] Fourth, there was relatively limited information on effects of newer triple therapy regimens with a protease inhibitor in subgroups defined by age, body weight, baseline fibrosis stage, and other important factors. Such information would be helpful for individualizing treatment decisions with these regimens. Finally, few methodologically rigorous studies conducted in settings applicable to U.S. populations evaluated the association between achieving an SVR and improvements in clinical outcomes. Such studies would be very helpful for confirming the results of the recent, large, well-conducted VA cohort study showing an association between achieving an SVR and reduced mortality risk.8

Future Research

Evaluating the comparative effectiveness of current antiviral regimens on clinical outcomes in randomized trials or cohort studies is a challenge due to the long lead-time and large samples necessary to adequately assess these outcomes. This might be more feasible if the studies were to focus on populations at higher risk for complications from chronic HCV infection (e.g., patients with baseline cirrhosis, high viral load, or other risk factors for progression).

For all trials of antiviral treatments, studies that enroll broader populations with medical and psychological comorbidities, as frequently encountered in clinical practice, are needed to better understand comparative effectiveness, rather than just comparative efficacy. Studies designed using an effectiveness paradigm would also be helpful for understanding real-world effects of antiviral regimens, including effects related to the poorer treatment adherence than expected from efficacy trials. Studies that evaluate the usefulness of genomics and other methods for individualizing treatment decisions in patients with HCV infection are also needed.

Trials directly comparing triple therapy with telaprevir compared with triple therapy with boceprevir would be very helpful for understanding comparative effectiveness of these two protease inhibitors. In addition, trials evaluating the boceprevir regimen by the FDA in antiviral-naïve patients without baseline cirrhosis are needed to verify that results from studies of previously treated patients were appropriately generalized. Prolonged followup of patients exposed to telaprevir and boceprevir is needed to understand the long-term harms associated with these medications. A number of other protease inhibitors and other newer drugs for treatment of hepatitis C virus infection are currently in active development and further studies with new drugs and drug regimens are expected, including regimens without interferon. 88

It is critical that future studies that evaluate clinical outcomes in patients with an SVR versus no SVR after antiviral therapy adequately control for other factors that influence clinical outcomes in chronic HCV infection. Studies on effects of achieving an SVR on long-term quality of life would be very helpful for understanding other potential clinical benefits of antiviral

therapy, but a significant challenge is whether it is possible to ethically blind patients to virologic status, which may have an important impact on assessments of quality of life.

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Abbreviations and Acronyms

AHRQ Agency for Healthcare Research and Quality

ALT Alanine aminotransferase

CER Comparative effectiveness review
CHIP Children's Health Insurance Program

CI Confidence Interval

EPC Evidence-based Practice Center

ESLD End-stage liver disease HCC Hepatocellular carcinoma

HCV Hepatitis C virus

HCV-RNA Hepatitis C virus ribonucleic acid HIV Human immunodeficiency virus

HR Hazard ratio

PCR Polymerase chain reaction

PEG Pegylated

PICOTS Populations, Interventions, Comparators, Outcomes, Timing, and Setting

RR Relative risk

SVR Sustained virologic response TEP Technical Expert Panel

USPSTF U.S. Preventive Services Task Force

Appendix A. Exact Search Strategy

The following databases have been searched for relevant information:

Database Searches: Hepatitis C: Treatment

Name	Date Limits	Platform Provider
Medline	2002 through August 2012	OvidSP
Embase	2002 through April 2012	Embase (Elsevier)
Cochrane Library:	2002 through August 2012	Cochrane Library
CDSR, DARE, CCRCT		
Clinical Trials.gov	2002 through August 2012	
Drugs@FDA	2002 through August 2012	
Health Canada Drug Products	2002 through August 2012	
Database		
European Public Assessment	2002 through August 2012	
Reports (European Medicine		
Agency)		
Scopus	2002 through August 2012	Scopus
PsycINFO	2002 through August 2012	OvidSP

Hand Search of Journals & Supplements - Topic-Specific Search Terms

Concept	Controlled Vocabulary	Keywords
Hepatitis C	Hepatitis C/	hcv.mp
	Hepatitis C,` Chronic/	hepacivirus\$.mp
	Hepacivirus/ OR	
Treatment	Antiviral agents/	Interferon\$
	Interferons/	interferon alpha-2a
	Interferon-alpha/	interferon alpha-2b
	Interferon Alfa-2a/	IFNalpha2a
	Interferon Alpha-2b/	IFNalpha2b
	Exp Polyethylene Glycols/	interferon alpha 2a
	Ribavirin/	interferon alpha 2b
	Exp Protease Inhibitors/	pegasys
		Peg-intron
		peginterferon alpha-2a peginterferon
		alpha-2b peginterferon alpha 2a
		peginterferon alpha 2b
		pegylated interferon\$
		IFN\$
		PEG IFN\$
		Ribavirin
		RBV
		protease inhibitor\$ polymerase

		inhibit\$ HCV protease\$ Telaprevir boceprevir
Harms -	AE.fs	Unsafe
treatment	MO.fs	Safety
	PO.fs	harm\$
	TO.fs	complication\$
	CT.fs	poison\$
		risk\$
		side-effect\$
		side effect\$
		(undesirable ADJ1 effect\$)
		(treatment ADJ1 emergent) tolerab\$ toxic\$
		adrs
		(adverse ADJ2 (effect or effects or reaction or reactions or event or events or outcome or outcomes)) (undesirable ADJ1 effect\$)
	AE=adverse effects	(treatment ADJ1 emergent) tolerab\$
	CT=contraindications	toxic\$
	MO=mortality	adrs
	PO=poisoning	(adverse ADJ2 (effect or effects or
	TO=toxicity	reaction or reactions or event or
		events or outcome or outcomes))

Original Search: 12/16/2011

Ovid MEDLINE(R) and Ovid OLDMEDLINE(R) 1946 to November Week 3 2011, Ovid MEDLINE(R) In-Process & Other Non-Indexed Citations December 15, 2011

1	Hepatitis C/ or Hepatitis C, Chronic/ or Hepacivirus/ or Hepatitis C.mp. or hepacivirus\$.mp. or HCV.mp.	58901
2	Antiviral agents/ or Interferons/ or Interferon-alpha/ or Interferon Alfa-2a/ or Interferon Alpha-2b/ or Interferon\$.mp. or interferon alpha-2a.mp. or interferon alpha-2a.mp. or interferon alpha-2b.mp. or interferon alpha 2a.mp. or interferon alpha 2b.mp. or exp Polyethylene Glycols/ or pegasys.mp. or Peginteron.mp. or peginterferon alpha-2a.mp. or peginterferon alpha-2b.mp. or peginterferon alpha-2b.mp. or peginterferon alpha-2b.mp. or pegylated	379981

	interferon\$.mp. or IFN\$.mp. or PEG IFN\$.mp. or Ribavirin/ or ribavirin.mp. or RBV.mp. or exp Protease Inhibitors/ or protease inhibitor\$.mp. or polymerase inhibit\$.mp. or HCV protease\$.mp. or telaprevir.mp. or boceprevir.mp.	
3	1 and 2	17670
4	(randomized controlled trial or controlled clinical trial or meta analysis or review).pt. or clinical trials as topic/ or cohort studies/ or randomized.ab. or randomly.ab. or placebo.ab. or (systematic adj1 review).ti,ab.	2498350
5	3 and 4	5896
6	limit 5 to (yr="2002 -Current" and ("adult (19 to 44 years)" or "middle age (45 to 64 years)" or "all aged (65 and over)"))	1382
7	(unsafe or safety or harm\$ or complication\$ or poison\$ or risk\$).mp. or AE.fs. or MO.fs. or PO.fs. or TO.fs. or CT.fs. or side-effect\$.mp. or (undesirable adj1 effect\$).mp. or (treatment adj1 emergent).mp. or tolerab\$.mp. or toxic\$.mp. or adrs.mp. or (adverse adj2 (effect or effects or reaction or reactions or event or events or outcome or outcomes)).mp.	3892024
8	1 and 2 and 7	7401
9	4 and 8	3168
10	limit 9 to (yr="2002 -Current" and ("adult (19 to 44 years)" or "middle age (45 to 64 years)" or "all aged (65 and over)"))	885
11	Counseling/ or Sex Counseling/ or Health Education/ or Patient Education as Topic/ or Psychotherapy/ or Behavior Therapy/ or Cognitive Therapy/ or Immunization/ or Immunotherapy/ or Psychotherapy, Brief/ or Socioenvironmental Therapy/	268601
12	1 and 11	662
13	6 and (201102* or 201103* or 201104* or 201105* or 201106* or 201107* or 201108* or 201109* or 201110* or 201111* or 201112*).ed.	132
	201106 of 201107 of 201111 of 2011112).ed.	
14	10 and (201109* or 201103* or 201110* or 2011105* or 201106* or 201107* or 201108* or 201109* or 201110* or 201111* or 201112*).ed.	90

Additional Treatment Search: 2/28/2011

Ovid MEDLINE (R) and Ovid OLDMED (R) 1947 to February Week 3 2011 Ovid MEDLINE(R) In-Process & Other Non-Indexed Citations February 28, 2011

1	Hepatitis C/ or Hepatitis C, Chronic/ or Hepacivirus/ or Hepatitis C.mp. or hepacivirus\$.mp. or HCV.mp.	58837
2	Antiviral agents/ or Interferons/ or Interferon-alpha/ or Interferon Alfa-2a/ or Interferon Alpha-2b/ or Interferon\$.mp. or interferon alpha-2a.mp. or interferon alpha-2b.mp. or IFNalpha2a.mp. or IFNalpha2b.mp. or interferon alpha 2a.mp. or interferon alpha 2b.mp. or exp Polyethylene Glycols/ or pegasys.mp. or Peginteron.mp. or peginterferon alpha-2a.mp. or peginterferon alpha-2b.mp. or peginterferon alpha-2b.mp. or peginterferon alpha-2b.mp. or pegylated interferon\$.mp. or IFN\$.mp. or PEG IFN\$.mp. or Ribavirin/ or ribavirin.mp. or RBV.mp. or exp Protease Inhibitors/ or protease inhibitor\$.mp. or polymerase inhibits.mp. or HCV protease\$.mp. or telaprevir.mp. or boceprevir.mp.	379770
3	1 and 2	17643
4	(randomized controlled trial or controlled clinical trial or meta analysis or review).pt. or clinical trials as topic/ or cohort studies/ or randomized.ab. or randomly.ab. or placebo.ab. or (systematic adj1 review).ti,ab.	2497187
5	3 and 4	5889
6	limit 5 to (yr="2002 -Current" and ("adult (19 to 44 years)" or "middle age (45 to 64 years)" or "all aged (65 and over)"))	1380
7	(unsafe or safety or harm\$ or complication\$ or poison\$ or risk\$).mp. or AE.fs. or MO.fs. or PO.fs. or TO.fs. or CT.fs. or side-effect\$.mp. or (undesirable adj1 effect\$).mp. or (treatment adj1 emergent).mp. or tolerab\$.mp. or toxic\$.mp. or adrs.mp. or (adverse adj2 (effect or effects or reaction or reactions or event or events or outcome or outcomes)).mp.	3889277
8	1 and 2 and 7	7391
9	4 and 8	3164
10	limit 9 to (yr="2002 -Current" and ("adult (19 to 44 years)" or "middle age (45 to 64 years)" or "all aged (65 and over)"))	883
11	Counseling/ or Sex Counseling/ or Health Education/ or Patient Education as	268554

	Topic/ or Psychotherapy/ or Behavior Therapy/ or Cognitive Therapy/ or Immunization/ or Immunotherapy/ or Psychotherapy, Brief/ or Socioenvironmental Therapy/	
12	1 and 11	660

Updated Search after Peer Review: 4/04/2012

Ovid MEDLINE Search Strategy 1947 to February Week 3 2011 Searched February 28, 2011; Update Search April 04, 2012		
1	Hepatitis C/ or Hepatitis C, Chronic/ or Hepacivirus/ or Hepatitis C.mp. or hepacivirus\$.mp. or HCV.mp.	
2	Antiviral agents/ or Interferons/ or Interferon-alpha/ or Interferon Alfa-2a/ or Interferon Alpha-2b/ or Interferons.mp. or interferon alpha-2a.mp. or interferon alpha-2b.mp. or IFNalpha2a.mp. or IFNalpha2b.mp. or interferon alpha 2a.mp. or interferon alpha 2b.mp. or exp Polyethylene Glycols/ or pegasys.mp. or Peg-intron.mp. or peginterferon alpha-2a.mp. or peginterferon alpha-2b.mp. or peginterferon alpha-2b.mp. or peginterferon alpha-2b.mp. or PEG IFNs.mp. or Ribavirin/ or ribavirin.mp. or RBV.mp. or exp Protease Inhibitors/ or protease inhibitors.mp. or polymerase inhibits.mp. or HCV proteases.mp. or telaprevir.mp. or boceprevir.mp.	
3	1 and 2	
4	(randomized controlled trial or controlled clinical trial or meta analysis or review).pt. or clinical trials as topic/ or cohort studies/ or randomized.ab. or randomly.ab. or placebo.ab. or (systematic adj1 review).ti,ab.	
5	3 and 4	
6	limit 5 to (yr="2002 -Current" and ("adult (19 to 44 years)" or "middle age (45 to 64 years)" or "all aged (65 and over)"))	
7	(unsafe or safety or harm\$ or complication\$ or poison\$ or risk\$).mp. or AE.fs. or MO.fs. or PO.fs. or TO.fs. or CT.fs. or side-effect\$.mp. or (undesirable adj1 effect\$).mp. or (treatment adj1 emergent).mp. or tolerab\$.mp. or toxic\$.mp. or adrs.mp. or (adverse adj2 (effect or effects or reaction or reactions or event or events or outcome or outcomes)).mp.	
8	1 and 2 and 7	
9	4 and 8	

10	limit 9 to (yr="2002 -Current" and ("adult (19 to 44 years)" or "middle age (45 to 64 years)" or "all aged (65 and over)"))
	Counseling/ or Sex Counseling/ or Health Education/ or Patient Education as Topic/ or Psychotherapy/ or Behavior Therapy/ or Cognitive Therapy/ or Immunization/ or Immunotherapy/ or Psychotherapy, Brief/ or Socioenvironmental Therapy/
12	1 and 11

EM	EMBASE Search Strategy 1976 – 2011		
Sea	rched April 11, 2011; Update Search April 4, 2012		
13	#12 AND (2002:py OR 2003:py OR 2004:py OR 2005:py OR 2006:py OR 2007:py OR 2008:py OR 2009:py OR 2010:py OR 2011:py)		
12	#3 AND #11		
11	#1 AND #10		
10	'counseling'/exp OR 'patient guidance'/exp OR 'patient counseling'/exp OR 'sexual counseling'/exp OR 'psychotherapy'/exp OR 'cognitive therapy'/exp OR 'behavior therapy'/exp OR 'sex therapy'/exp OR 'patient education'/exp OR 'immunization'/exp OR 'virus vaccine'/exp OR 'immunotherapy'/exp OR counsel* OR 'socioenvironmental therapy'/de AND [embase]/lim		
9	#8 AND (2002:py OR 2003:py OR 2004:py OR 2005:py OR 2006:py OR 2007:py OR 2008:py OR 2009:py OR 2010:py OR 2011:py)		
8	#3 AND #7		
7	#1 AND #2 AND #6		
6	'adverse drug reaction'/exp OR 'adverse outcome'/exp OR 'toxicity'/exp OR 'drug toxicity'/exp OR 'drug tolerability'/exp OR 'drug safety'/exp OR 'patient safety'/exp OR unsafe OR 'safety'/exp OR harm* OR complication* OR poison* OR 'side effect'/exp OR 'side effects' OR undesirable NEAR/1 effect* OR treatment NEAR/1 emergen* OR tolerab* OR toxic* OR adrs OR adverse NEAR/2 (effect OR effects OR reaction OR reactions OR event OR events OR outcome OR outcomes) AND [embase]/lim		
5	#4 AND (2002:py OR 2003:py OR 2004:py OR 2005:py OR 2006:py OR 2007:py OR 2008:py OR 2009:py OR 2010:py OR 2011:py)		
4	#1 AND #2 AND #3		
3	'cohort analysis'/exp OR 'meta analysis'/exp OR 'randomized controlled trial'/exp OR 'systematic review'/exp OR 'controlled clinical trial'/exp OR 'placebo'/exp OR 'clinical trial'/exp OR 'controlled study'/exp OR randomized.ab OR randomly.ab AND [embase]/lim		
2	'antivirus agent'/exp OR 'antivirus agent' OR 'interferon'/exp OR interferon OR 'alpha interferon'/exp OR 'alpha interferon' OR 'alpha2a interferon'/exp OR 'alpha2a interferon' OR 'alpha2b interferon'/exp OR 'alpha2b interferon' OR 'macrogol derivative'/exp OR 'macrogol derivative' OR 'peginterferon'/exp OR peginterferon OR 'peginterferon alpha2a'/exp OR 'peginterferon alpha2a' OR 'peginterferon alpha2b'/exp OR 'peginterferon alpha2b' OR		

'ribavirin'/exp OR ribavirin OR 'protease inhibitor'/exp OR 'protease inhibitor' OR 'rna directed dna polymerase inhibitor' OR 'rna directed rna polymerase inhibitor'/exp OR 'rna directed rna polymerase inhibitor' OR 'telaprevir'/exp OR telaprevir OR 'boceprevir'/exp OR boceprevir OR 'antiviral agent':ab,ti OR interferon*:ab,ti OR 'interferon-alpha2a':ab,ti OR 'interferon-alpha2b':ab,ti OR 'interferon alpha':ab,ti OR 'peginterferon alpha 2a':ab,ti OR 'peginterferon alpha 2b':ab,ti OR 'polyethylene glycols':ab,ti OR pegasys:ab,ti OR 'peg intron':ab,ti OR 'peginterferon alpha 2a':ab,ti OR 'peginterferon alpha 2b':ab,ti OR 'pegylated interferon':ab,ti OR ifn:ab,ti OR 'peg ifn':ab,ti OR 'peg ifns':ab,ti OR ribavirin:ab,ti OR rbv:ab,ti OR 'protease inhibitor':ab,ti OR 'polymerase inhibitor':ab,ti OR 'polymerase inhibitors':ab,ti OR 'polymerase inhibit

'hepatitis c virus':de OR 'hepatitis c':de OR 'chronic active hepatitis':de OR 'hepatitis non a non b':de AND [embase]/lim

Cochrane Library:

Cochrane Database of Systematic Reviews & Database of Abstracts of Reviews of Effects 2002-2011

Searched April 11, 2011, Update Search April 4, 2012

"Hepatitis C" OR Hepacivirus OR HCV (Title, Abstract, Keyword)

Limit to reviews, published 2002-2011

Cochrane Library:

Cochrane Central Register of Controlled Trials 2002-2011

Searched April 11, 2011; Update Search April 04, 2012

"Interferon-alpha" OR "Interferon Alfa-2a" OR "Interferon Alpha-2b" OR "IFNalpha2a" OR "IFNalpha2b" OR "Interferon alpha 2a" OR "interferon alpha 2b" OR "Polyethylene Glycol*" OR pegasys OR Peg-intron OR "peginterferon alpha-2a" OR "peginterferon alpha-2b" OR "peginterferon alpha 2a" OR "peginterferon alpha 2b" OR "pegylated interferon*" OR IFN* OR "PEG IFN*" OR Ribavirin OR RBV OR "protease inhibitor*" OR "polymerase inhibit*" OR "HCV protease*" OR telaprevir OR boceprevir (Title, Abstract, Keyword)

SCOPUS Search Strategy 1960-2011

Searched April 11, 2011; Update Search April 04, 2012

(TITLE-ABS-KEY("hepatitis c" OR hepacivirus OR hcv)) AND (TITLE-ABS-KEY(cohort* OR "meta analysis" OR "randomized controlled trial*" OR "systematic review*" OR "controlled clinical trial*" OR "placebo" OR "clinical trial*" OR randomized OR randomly)) AND (TITLE-ABS-KEY(counseling OR "health education" OR "patient education" OR psychotherapy OR "behavior therapy" OR "cognitive therapy" OR immuniz* OR immunotherapy OR "socioenvironmental therapy" OR "cognitive behavior* therapy" OR vaccine*))

- TITLE-ABS-KEY(counseling OR "health education" OR "patient education" OR psychotherapy OR "behavior therapy" OR "cognitive therapy" OR immuniz* OR immunotherapy OR "socioenvironmental therapy" OR "cognitive behavior* therapy" OR vaccine*)
- (TITLE-ABS-KEY("hepatitis c" OR hepacivirus OR hcv)) AND ((TITLE-ABS-KEY("antiviral agent*" OR interferon* OR interferon-alpha OR "interferon alfa-2a" OR "interferon alpha-2b" OR ifnalpha2a OR ifnalpha2b OR "interferon alpha 2a" OR "interferon alpha 2b" OR "polyethylene glycols" OR pegasys OR peg-intron) OR TITLE-ABS-KEY("peginterferon alpha-2a" OR "peginterferon alpha-2b" OR "peginterferon alpha 2a" OR "peginterferon alpha 2b" OR "pegylated interferon*" OR ifn* OR peg ifn* OR ribavirin OR rbv OR "protease inhibitor*" OR "polymerase inhibitor*" OR "hcv protease*" OR telapr))) AND (TITLE-ABS-KEY(cohort* OR "meta analysis" OR "randomized controlled trial*" OR "systematic review*" OR "controlled clinical trial*" OR "placebo" OR "clinical trial*" OR randomized OR randomly)) AND (TITLE-ABS-KEY(unsafe OR safety OR harm* OR complication* OR poison* OR risk* OR side-effect* OR "side effect*" OR "undesirable effect* OR "treatment emergent" OR tolerab* OR toxic* OR "adverse effect*" OR "adverse reaction*" OR "adverse event*" OR "adverse outcome*")) AND (LIMIT-TO(PUBYEAR, 2011) OR LIMIT-TO(PUBYEAR, 2010) OR LIMIT-TO(PUBYEAR, 2009) OR LIMIT-TO(PUBYEAR, 2008) OR LIMIT-TO(PUBYEAR, 2007) OR LIMIT-TO(PUBYEAR, 2006) OR LIMIT-TO(PUBYEAR, 2005) OR LIMIT-TO(PUBYEAR, 2004) OR LIMIT-TO(PUBYEAR, 2003))
- KEY("antiviral agent*" OR interferon* OR interferon-alpha OR "interferon alfa-2a" OR "interferon alpha-2b" OR ifnalpha2a OR ifnalpha2b OR "interferon alpha 2a" OR "interferon alpha 2b" OR "polyethylene glycols" OR pegasys OR peg-intron) OR TITLE-ABS-KEY("peginterferon alpha-2a" OR "peginterferon alpha-2b" OR "peginterferon alpha-2a" OR "peginterferon alpha-2b" OR "peginterferon alpha 2a" OR "peginterferon alpha 2b" OR "pegylated interferon*" OR ifn* OR peg ifn* OR ribavirin OR rbv OR "protease inhibitor*" OR "polymerase inhibitor*" OR "hcv protease*" OR telapr))) AND (TITLE-ABS-KEY(cohort* OR "meta analysis" OR "randomized controlled trial*" OR "systematic review*" OR "controlled clinical trial*" OR "placebo" OR "clinical trial*" OR randomized OR randomly)) AND (TITLE-ABS-KEY(unsafe OR safety OR harm* OR complication* OR poison* OR risk* OR side-effect* OR "side effect*" OR "undesirable effect* OR "treatment emergent" OR tolerab* OR toxic* OR "adverse effect*" OR "adverse reaction*" OR "adverse event*" OR "adverse outcome*"))
- 7 TITLE-ABS-KEY(unsafe OR safety OR harm* OR complication* OR poison* OR risk* OR side-effect* OR "side effect*" OR "undesirable effect* OR "treatment emergent" OR tolerab* OR toxic* OR "adverse effect*" OR "adverse reaction*" OR "adverse event*" OR "adverse outcome*")
- 6 (TITLE-ABS-KEY("hepatitis c" OR hepacivirus OR hcv)) AND ((TITLE-ABS-KEY("antiviral agent*" OR interferon* OR interferon-alpha OR "interferon alfa-2a" OR "interferon alpha-2b" OR ifnalpha2a OR ifnalpha2b OR "interferon alpha 2a" OR "interferon alpha 2b" OR "polyethylene glycols" OR pegasys OR peg-intron) OR TITLE-ABS-KEY("peginterferon alpha-2a" OR "peginterferon alpha-2b" OR "peginterferon alpha 2a" OR "peginterferon alpha 2b" OR "pegylated interferon*" OR ifn* OR peg ifn* OR ribavirin OR rbv OR "protease inhibitor*" OR "polymerase inhibitor*" OR "hcv protease*"

OR telapr))) AND (TITLE-ABS-KEY(cohort* OR "meta analysis" OR "randomized controlled trial*" OR "systematic review*" OR "controlled clinical trial*" OR "placebo" OR "clinical trial*" OR randomized OR randomly)) AND (LIMIT-TO(PUBYEAR, 2011) OR LIMIT-TO(PUBYEAR, 2010) OR LIMIT-TO(PUBYEAR, 2009) OR LIMIT-TO(PUBYEAR, 2008) OR LIMIT-TO(PUBYEAR, 2007) OR LIMIT-TO(PUBYEAR, 2006) OR LIMIT-TO(PUBYEAR, 2005) OR LIMIT-TO(PUBYEAR, 2004) OR LIMIT-TO(PUBYEAR, 2003) OR LIMIT-TO(PUBYEAR, 2002))

- (TITLE-ABS-KEY("hepatitis c" OR hepacivirus OR hcv)) AND ((TITLE-ABS-KEY("antiviral agent*" OR interferon* OR interferon-alpha OR "interferon alfa-2a" OR "interferon alpha-2b" OR ifnalpha2a OR ifnalpha2b OR "interferon alpha 2a" OR "interferon alpha 2b" OR "polyethylene glycols" OR pegasys OR peg-intron) OR TITLE-ABS-KEY("peginterferon alpha-2a" OR "peginterferon alpha-2b" OR "peginterferon alpha 2a" OR "peginterferon alpha 2b" OR "pegylated interferon*" OR ifn* OR peg ifn* OR ribavirin OR rbv OR "protease inhibitor*" OR "polymerase inhibitor*" OR "hcv protease*" OR telapr))) AND (TITLE-ABS-KEY(cohort* OR "meta analysis" OR "randomized controlled trial*" OR "systematic review*" OR "controlled clinical trial*" OR "placebo" OR "clinical trial*" OR randomized OR randomly))
- (TITLE-ABS-KEY("hepatitis c" OR hepacivirus OR hcv)) AND ((TITLE-ABS-KEY("antiviral agent*" OR interferon* OR interferon-alpha OR "interferon alfa-2a" OR "interferon alpha-2b" OR ifnalpha2a OR ifnalpha2b OR "interferon alpha 2a" OR "interferon alpha 2b" OR "polyethylene glycols" OR pegasys OR peg-intron) OR TITLE-ABS-KEY("peginterferon alpha-2a" OR "peginterferon alpha-2b" OR "peginterferon alpha 2a" OR "peginterferon alpha 2b" OR "pegylated interferon*" OR ifn* OR peg ifn* OR ribavirin OR rbv OR "protease inhibitor*" OR "polymerase inhibitor*" OR "hcv protease*" OR telapr))) AND (TITLE-ABS-KEY(cohort* OR "meta analysis" OR "randomized controlled trial*" OR "systematic review*" OR "controlled clinical trial*" OR "placebo" OR "clinical trial*" OR randomized OR randomly))
- TITLE-ABS-KEY(cohort* OR "meta analysis" OR "randomized controlled trial*" OR "systematic review*" OR "controlled clinical trial*" OR "placebo" OR "clinical trial*" OR randomized OR randomly)
- (TITLE-ABS-KEY("antiviral agent*" OR interferon* OR interferon-alpha OR "interferon alfa-2a" OR "interferon alpha-2b" OR ifnalpha2a OR ifnalpha2b OR "interferon alpha 2a" OR "interferon alpha 2b" OR "polyethylene glycols" OR pegasys OR peg-intron) OR TITLE-ABS-KEY("peginterferon alpha-2a" OR "peginterferon alpha-2b" OR "peginterferon alpha 2a" OR "peginterferon alpha-2b" OR "peginterferon alpha 2a" OR "peginterferon alpha 2b" OR "pegylated interferon*" OR ifn* OR peg ifn* OR ribavirin OR rbv OR "protease inhibitor*" OR "polymerase inhibitor*" OR "hcv protease*" OR telaprevir))
- 1 TITLE-ABS-KEY("hepatitis c" OR hepacivirus OR hcv)

OvidSP PSYCINFO Search Strategy 1806 to February Week 4 2011 Searched April 12, 2011; Update Search April 4, 2012

1 hepatitis/ or (Hepatitis C or hepacivirus\$ or HCV).mp.

2	[exp treatment/ or exp intervention/ or exp psychotherapy/ or exp alcohol rehabilitation/ or exp counseling/ or exp support groups/ or exp rehabilitation/ or exp mental health services/ or exp community services/ or exp outreach programs/ or exp drug rehabilitation/ or exp sobriety/ or exp detoxification/ or exp drug rehabilitation/ or exp treatment outcomes/ or exp alcoholics anonymous/]
3	alcohol*.mp.
4	1 and 2 and 3

Clinicaltrials.gov

Searched April 12, 2011; Update Search April 4, 2012

interferon alfa OR peginterferon OR ribavirin OR telaprevir OR boceprevir | Closed Studies | Studies With Results | hepatitis c | Adult, Senior

Updated Search: 8/28/2012

	Ovid MEDLINE Search Strategy 1947 to February Week 3 2011 Searched February 28, 2011; Update Search April 04, 2012; Update Search August 28, 2012		
1	Hepatitis C/ or Hepatitis C, Chronic/ or Hepacivirus/ or Hepatitis C.mp. or hepacivirus\$.mp. or HCV.mp.		
2	Antiviral agents/ or Interferons/ or Interferon-alpha/ or Interferon Alfa-2a/ or Interferon Alpha-2b/ or Interferon\$.mp. or interferon alpha-2a.mp. or interferon alpha-2b.mp. or IFNalpha2a.mp. or IFNalpha2b.mp. or interferon alpha 2a.mp. or interferon alpha 2b.mp. or exp Polyethylene Glycols/ or pegasys.mp. or Peg-intron.mp. or peginterferon alpha-2a.mp. or peginterferon alpha-2b.mp. or peginterferon alpha-2b.mp. or peginterferon alpha-2b.mp. or pegylated interferon\$.mp. or IFN\$.mp. or PEG IFN\$.mp. or Ribavirin/ or ribavirin.mp. or RBV.mp. or exp Protease Inhibitors/ or protease inhibitor\$.mp. or polymerase inhibit\$.mp. or HCV protease\$.mp. or telaprevir.mp. or boceprevir.mp.		
3	1 and 2		
4	(randomized controlled trial or controlled clinical trial or meta analysis or review).pt. or clinical trials as topic/ or cohort studies/ or randomized.ab. or randomly.ab. or placebo.ab. or (systematic adj1 review).ti,ab.		
5	3 and 4		
6	limit 5 to (yr="2002 -Current" and ("adult (19 to 44 years)" or "middle age (45 to 64 years)" or "all aged (65 and over)"))		

7	(unsafe or safety or harm\$ or complication\$ or poison\$ or risk\$).mp. or AE.fs. or MO.fs. or PO.fs. or TO.fs. or CT.fs. or side-effect\$.mp. or (undesirable adj1 effect\$).mp. or (treatment adj1 emergent).mp. or tolerab\$.mp. or toxic\$.mp. or adrs.mp. or (adverse adj2 (effect or effects or reaction or reactions or event or events or outcome or outcomes)).mp.
8	1 and 2 and 7
9	4 and 8
10	limit 9 to (yr="2002 -Current" and ("adult (19 to 44 years)" or "middle age (45 to 64 years)" or "all aged (65 and over)"))
11	Counseling/ or Sex Counseling/ or Health Education/ or Patient Education as Topic/ or Psychotherapy/ or Behavior Therapy/ or Cognitive Therapy/ or Immunization/ or Immunotherapy/ or Psychotherapy, Brief/ or Socioenvironmental Therapy/
12	1 and 11

EMBASE Search Strategy 1976 – 2011

An updated search for August 28, 2012 was not conducted as Oregon Health and Sciences University no longer subscribes to this database.

Cochrane Library:

Cochrane Database of Systematic Reviews & Database of Abstracts of Reviews of Effects 2002-2011

Searched April 11, 2011, Update Search April 4, 2012; Update Search August 28, 2012

"Hepatitis C" OR Hepacivirus OR HCV (Title, Abstract, Keyword)

Limit to reviews, published 2002-2011

Cochrane Library:

Cochrane Central Register of Controlled Trials 2002-2011

Searched April 11, 2011; Update Search April 04, 2012; Update Search August 28, 2012

"Interferon-alpha" OR "Interferon Alfa-2a" OR "Interferon Alpha-2b" OR "IFNalpha2a" OR "IFNalpha2b" OR "Interferon alpha 2a" OR "interferon alpha 2b" OR "Polyethylene Glycol*" OR pegasys OR Peg-intron OR "peginterferon alpha-2a" OR "peginterferon alpha-2b" OR "peginterferon alpha-2b" OR "peginterferon alpha 2a" OR "peginterferon alpha 2b" OR "pegylated interferon*" OR IFN* OR "PEG IFN*" OR Ribavirin OR RBV OR "protease inhibitor*" OR "polymerase inhibit*" OR "HCV protease*" OR telaprevir OR boceprevir (Title, Abstract, Keyword)

SCOPUS Search Strategy 1960-2011

Searched April 11, 2011; Update Search April 04, 2012; Update Search August 28, 2012

(TITLE-ABS-KEY("hepatitis c" OR hepacivirus OR hcv)) AND (TITLE-ABS-KEY(cohort* OR "meta analysis" OR "randomized controlled trial*" OR "systematic review*" OR "controlled clinical trial*" OR "placebo" OR "clinical trial*" OR randomized OR randomly)) AND (TITLE-ABS-KEY(counseling OR "health education" OR "patient

- education" OR psychotherapy OR "behavior therapy" OR "cognitive therapy" OR immuniz* OR immunotherapy OR "socioenvironmental therapy" OR "cognitive behavior* therapy" OR vaccine*))
- TITLE-ABS-KEY(counseling OR "health education" OR "patient education" OR psychotherapy OR "behavior therapy" OR "cognitive therapy" OR immuniz* OR immunotherapy OR "socioenvironmental therapy" OR "cognitive behavior* therapy" OR vaccine*)
- (TITLE-ABS-KEY("hepatitis c" OR hepacivirus OR hcv)) AND ((TITLE-ABS-9 KEY("antiviral agent*" OR interferon* OR interferon-alpha OR "interferon alfa-2a" OR "interferon alpha-2b" OR ifnalpha2a OR ifnalpha2b OR "interferon alpha 2a" OR "interferon alpha 2b" OR "polyethylene glycols" OR pegasys OR peg-intron) OR TITLE-ABS-KEY("peginterferon alpha-2a" OR "peginterferon alpha-2b" OR "peginterferon alpha 2a" OR "peginterferon alpha 2b" OR "pegylated interferon*" OR ifn* OR peg ifn* OR ribavirin OR rbv OR "protease inhibitor*" OR "polymerase inhibitor*" OR "hcv protease*" OR telapr))) AND (TITLE-ABS-KEY(cohort* OR "meta analysis" OR "randomized controlled trial*" OR "systematic review*" OR "controlled clinical trial*" OR "placebo" OR "clinical trial*" OR randomized OR randomly)) AND (TITLE-ABS-KEY(unsafe OR safety OR harm* OR complication* OR poison* OR risk* OR side-effect* OR "side effect*" OR "undesirable effect* OR "treatment emergent" OR tolerab* OR toxic* OR "adverse effect*" OR "adverse reaction*" OR "adverse event*" OR "adverse outcome*")) AND (LIMIT-TO(PUBYEAR, 2011) OR LIMIT-TO(PUBYEAR, 2010) OR LIMIT-TO(PUBYEAR, 2009) OR LIMIT-TO(PUBYEAR, 2008) OR LIMIT-TO(PUBYEAR, 2007) OR LIMIT-TO(PUBYEAR, 2006) OR LIMIT-TO(PUBYEAR, 2005) OR LIMIT-TO(PUBYEAR, 2004) OR LIMIT-TO(PUBYEAR, 2003))
- (TITLE-ABS-KEY("hepatitis c" OR hepacivirus OR hcv)) AND ((TITLE-ABS-KEY("antiviral agent*" OR interferon* OR interferon-alpha OR "interferon alfa-2a" OR "interferon alpha-2b" OR ifnalpha2a OR ifnalpha2b OR "interferon alpha 2a" OR "interferon alpha 2b" OR "polyethylene glycols" OR pegasys OR peg-intron) OR TITLE-ABS-KEY("peginterferon alpha-2a" OR "peginterferon alpha-2b" OR "peginterferon alpha 2a" OR "peginterferon alpha 2b" OR "pegylated interferon*" OR ifn* OR peg ifn* OR ribavirin OR rbv OR "protease inhibitor*" OR "polymerase inhibitor*" OR "hcv protease*" OR telapr))) AND (TITLE-ABS-KEY(cohort* OR "meta analysis" OR "randomized controlled trial*" OR "systematic review*" OR "controlled clinical trial*" OR "placebo" OR "clinical trial*" OR randomized OR randomly)) AND (TITLE-ABS-KEY(unsafe OR safety OR harm* OR complication* OR poison* OR risk* OR side-effect* OR "side effect*" OR "undesirable effect* OR "treatment emergent" OR tolerab* OR toxic* OR "adverse effect*" OR "adverse reaction*" OR "adverse event*" OR "adverse outcome*"))
- 7 TITLE-ABS-KEY(unsafe OR safety OR harm* OR complication* OR poison* OR risk* OR side-effect* OR "side effect*" OR "undesirable effect* OR "treatment emergent" OR tolerab* OR toxic* OR "adverse effect*" OR "adverse reaction*" OR "adverse event*" OR "adverse outcome*")
- 6 (TITLE-ABS-KEY("hepatitis c" OR hepacivirus OR hcv)) AND ((TITLE-ABS-KEY("antiviral agent*" OR interferon* OR interferon-alpha OR "interferon alfa-2a" OR "interferon alpha-2b" OR ifnalpha2a OR ifnalpha2b OR "interferon alpha 2a" OR "interferon alpha 2b" OR "polyethylene glycols" OR pegasys OR peg-intron) OR TITLE-ABS-KEY("peginterferon alpha-2a" OR "peginterferon alpha-2b" OR "peginterferon alpha

2a" OR "peginterferon alpha 2b" OR "pegylated interferon*" OR ifn* OR peg ifn* OR ribavirin OR rbv OR "protease inhibitor*" OR "polymerase inhibitor*" OR "hcv protease*" OR telapr))) AND (TITLE-ABS-KEY(cohort* OR "meta analysis" OR "randomized controlled trial*" OR "systematic review*" OR "controlled clinical trial*" OR "placebo" OR "clinical trial*" OR randomized OR randomly)) AND (LIMIT-TO(PUBYEAR, 2011) OR LIMIT-TO(PUBYEAR, 2009) OR LIMIT-TO(PUBYEAR, 2008) OR LIMIT-TO(PUBYEAR, 2007) OR LIMIT-TO(PUBYEAR, 2006) OR LIMIT-TO(PUBYEAR, 2005) OR LIMIT-TO(PUBYEAR, 2004) OR LIMIT-TO(PUBYEAR, 2003) OR LIMIT-TO(PUBYEAR, 2002))

- (TITLE-ABS-KEY("hepatitis c" OR hepacivirus OR hcv)) AND ((TITLE-ABS-KEY("antiviral agent*" OR interferon* OR interferon-alpha OR "interferon alfa-2a" OR "interferon alpha-2b" OR ifnalpha2a OR ifnalpha2b OR "interferon alpha 2a" OR "interferon alpha 2b" OR "polyethylene glycols" OR pegasys OR peg-intron) OR TITLE-ABS-KEY("peginterferon alpha-2a" OR "peginterferon alpha-2b" OR "peginterferon alpha 2a" OR "peginterferon alpha 2b" OR "pegylated interferon*" OR ifn* OR peg ifn* OR ribavirin OR rbv OR "protease inhibitor*" OR "polymerase inhibitor*" OR "hcv protease*" OR telapr))) AND (TITLE-ABS-KEY(cohort* OR "meta analysis" OR "randomized controlled trial*" OR "systematic review*" OR "controlled clinical trial*" OR "placebo" OR "clinical trial*" OR randomized OR randomly))
- (TITLE-ABS-KEY("hepatitis c" OR hepacivirus OR hcv)) AND ((TITLE-ABS-KEY("antiviral agent*" OR interferon* OR interferon-alpha OR "interferon alfa-2a" OR "interferon alpha-2b" OR ifnalpha2a OR ifnalpha2b OR "interferon alpha 2a" OR "interferon alpha 2b" OR "polyethylene glycols" OR pegasys OR peg-intron) OR TITLE-ABS-KEY("peginterferon alpha-2a" OR "peginterferon alpha-2b" OR "peginterferon alpha 2a" OR "peginterferon alpha 2b" OR "pegylated interferon*" OR ifn* OR peg ifn* OR ribavirin OR rbv OR "protease inhibitor*" OR "polymerase inhibitor*" OR "hcv protease*" OR telapr))) AND (TITLE-ABS-KEY(cohort* OR "meta analysis" OR "randomized controlled trial*" OR "systematic review*" OR "controlled clinical trial*" OR "placebo" OR "clinical trial*" OR randomized OR randomly))
- TITLE-ABS-KEY(cohort* OR "meta analysis" OR "randomized controlled trial*" OR "systematic review*" OR "controlled clinical trial*" OR "placebo" OR "clinical trial*" OR randomized OR randomly)
- (TITLE-ABS-KEY("antiviral agent*" OR interferon* OR interferon-alpha OR "interferon alfa-2a" OR "interferon alpha-2b" OR ifnalpha2a OR ifnalpha2b OR "interferon alpha 2a" OR "interferon alpha 2b" OR "polyethylene glycols" OR pegasys OR peg-intron) OR TITLE-ABS-KEY("peginterferon alpha-2a" OR "peginterferon alpha-2b" OR "peginterferon alpha 2a" OR "peginterferon alpha 2b" OR "pegylated interferon*" OR ifn* OR peg ifn* OR ribavirin OR rbv OR "protease inhibitor*" OR "polymerase inhibitor*" OR "hcv protease*" OR telaprevir))
- 1 TITLE-ABS-KEY("hepatitis c" OR hepacivirus OR hcv)

OvidSP PSYCINFO Search Strategy 1806 to February Week 4 2011 Searched April 12, 2011; Update Search April 4, 2012; Update Search August 28, 2012

1 hepatitis/ or (Hepatitis C or hepacivirus\$ or HCV).mp.

[exp treatment/ or exp intervention/ or exp psychotherapy/ or exp alcohol rehabilitation/ or exp counseling/ or exp support groups/ or exp rehabilitation/ or exp mental health services/ or exp community services/ or exp outreach programs/ or exp drug rehabilitation/ or exp sobriety/ or exp detoxification/ or exp drug rehabilitation/ or exp treatment outcomes/ or exp alcoholics anonymous/]

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Clinicaltrials.gov

Searched April 12, 2011; Update Search April 4, 2012, Update Search August 28, 2012

interferon alfa OR peginterferon OR ribavirin OR telaprevir OR boceprevir | Closed Studies | Studies With Results | hepatitis c | Adult, Senior

Appendix B. Hepatitis C Treatment: Inclusion Criteria by Key Question

	Inclusion Criteria
Populations	Asymptomatic adults with chronic hepatitis C virus infection who have not received antiviral drug treatment previously
	Subgroups include: HCV genotype, race, sex, stage of disease, viral load, weight, and others (e.g. genetic markers)
	 Excluded: Pregnant women, HIV co-infected, transplant recipients, patients with renal failure
Interventions	KQ 1a and b:
	1a. What is the comparative effectiveness of antiviral treatment in improving health outcomes in patients with HCV infection?
	1b. How does the comparative effectiveness of antiviral treatment for health outcomes
	vary according to patient subgroup characteristics, including but not limited to HCV
	genotype, race, sex, disease severity or genetic markers?
	KQ 2a and b:
	2. What is the comparative effectiveness of antiviral treatments in improving intermediate outcomes, such as the rate of viremia, aminotransaminase levels, and histologic changes?
	2a. How does the comparative effectiveness of antiviral treatment for intermediate outcomes vary according to patient subgroup characteristics, including but not limited to HCV genotype, race, sex, disease severity or genetic markers?
	KQ 3a and b:
	3a. What are the comparative harms (including intolerance to treatment) associated with antiviral treatment?
	3b. Do these harms differ according to patient subgroup characteristics, including HCV
	genotype, race, sex, disease severity or genetic markers?
	KQ 4:
	Have improvements in intermediate outcomes (viremia, liver function tests, histologic
	changes) been shown to reduce the risk or rates of health outcomes from HCV infection?

	Inclusion Criteria
Comparisons	KQ 1a and b:
	1a. What is the comparative effectiveness of antiviral treatment in improving health
	outcomes in patients with HCV infection?
	1b. How does the comparative effectiveness of antiviral treatment for health outcomes
	vary according to patient subgroup characteristics, including but not limited to HCV
	genotype, race, sex, disease severity or genetic markers?
	8. 13,1.4, 11
	KQ 2a and b:
	2a. What is the comparative effectiveness of antiviral treatments in improving
	intermediate outcomes, such as the rate of viremia, aminotransaminase levels,
	and histologic changes?
	2b. How does the comparative effectiveness of antiviral treatment for intermediate
	outcomes vary according to patient subgroup characteristics, including but not limited to
	HCV genotype, race, sex, disease severity or genetic markers?
	KQ 3a and b:
	3. What are the comparative harms (including intolerance to treatment) associated with
	antiviral treatment?
	3a. Do these harms differ according to patient subgroup characteristics, including HCV
	genotype, race, sex, disease severity or genetic markers?
	KQ 4:
	Have improvements in intermediate outcomes (viremia, liver function tests, histologic
	changes) been shown to reduce the risk or rates of health outcomes from HCV infection?
Outcomes	Clinical outcomes
	Mortality (all-cause or hepatic)
	Cirrhosis
	Hepatic decompensation
	Hepatocellular carcinoma
	Need for liver transplantation
	Quality of life
	 Harms from antiviral treatments (including withdrawals due to adverse
	events, neutropenia, anemia, psychological adverse events, flu-like
	symptoms, rash)
	Intermediate outcomes
	Sustained virological response
	Improvement in liver histology
Settings	All settings (including primary care and specialty settings) and locales, though focus on
	studies conducted in the U.S. and other developed countries.
Study designs	KQ 3a and b:
	3a. What are the comparative harms (including intolerance to treatment) associated with
	antiviral treatment?
	3b. Do these harms differ according to patient subgroup characteristics, including HCV
	genotype, race, sex, disease severity or genetic markers?
	KQ 4:
	Have improvements in intermediate outcomes (viremia, liver function tests, histologic
	changes) been shown to reduce the risk or rates of health outcomes from HCV infection?

Appendix C. Included Studies List

Key Question 1: Not Applicable Key Questions 2 and 3:

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. Journal of Viral Hepatitis. 2006 Dec;13(12):811-20. PMID: 17109680

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Escudero A, Rodriguez F, Serra MA, et al. Pegylated alpha-interferon-2a plus ribavirin compared with pegylated alpha-interferon-2b plus ribavirin for initial treatment of chronic hepatitis C virus: prospective, non-randomized study. Journal of Gastroenterology & Hepatology. 2008 Jun;23(6):861-6. PMID: 18422960

Ferenci P, Laferl H, Scherzer T-M, et al. Peginterferon alfa-2a/ribavirin for 48 or 72 weeks in hepatitis C genotypes 1 and 4 patients with slow virologic response.[Reprint in Korean J Hepatol. 2010 Jun;16(2):201-5; PMID: 20606507]. Gastroenterology. 2010 Feb;138(2):503-12. PMID: 19909752

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Key Question 4:

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Innes HA, Hutchinson SJ, Allen S, et al. Excess liver-related morbidity of chronic hepatitis C patients, who achieve a sustained viral response, and are discharged from care. Hepatology. 2011; 54(5):1547-58. PMID:22045672

Izumi N, Yasuhiro A, Kurosaki M, et al. Development of hepatocellular carcinoma after interferon therapy in chronic hepatitis C. Is it possible to reduce the incidence by ribanirin and IFN combination therapy? Intervirology. 2005;48(1):59-63. PMID: 15785091

Kasahara A, Tanaka H, Okanoue T, et al. Interferon treatment improves survival in chronic hepatitis C patients showing biochemical as well as virological responses by preventing liver-related death. Journal of Viral Hepatitis. 2004 Mar;11(2):148-56. PMID: 14996350

Maruoka D, Imazeki F, Arai M, et al. Long-term cohort study of chronic hepatitis C according to interferon efficacy. Journal of Gastroenterology and Hepatology; 2012;27(2):291-99. PMID: 21793911

McHutchison J, Manns M, Harvey J, et al. Adherence to therapy enhanges sustained response in chronic hepatitis C patients receiving PEG-Interferon alfa-2b plus Ribavirin[abstract]. Journal of Hepatology. 2001;34(1):2-3.

McHutchison J, Sulkowski M. Scientific rationale and study design of the individualized dosing efficacy vs flat dosing to assess optimal pegylated interferon therapy (IDEAL) trial: determining optimal dosing in patients with genotype 1 chronic hepatitis C. Journal of **Viral Hepatitis**. 2008;1

Morgan TR, Ghany MG, Kim H-Y, et al. Outcome of sustained virological responders with histologically advanced chronic hepatitis C. Hepatology. 2010 Sep;52(3):833-44. PMID: 20564351

Neary MP, Cort S, Bayliss MS, et al. Sustained virologic response is associated with improved health-related quality of life in relapsed chronic hepatitis C patients. Seminars in Liver Disease. 1999;19(1):77-85. PMID: 10349695

Rasenack J, Zeuzem S, Feinman SV, et al. Peginterferon alpha-2a (40kD) [Pegasys] improves HR-QOL outcomes compared with unmodified interferon alpha-2a [Roferon-A]: in patients with chronic hepatitis C.[Erratum appears in Pharmacoeconomics. 2003;21(17):1290]. Pharmacoeconomics. 2003;21(5):341-9. PMID: 12627987

Shiratori Y, Ito Y, Yokosuka O, et al. Antiviral therapy for cirrhotic hepatitis C: association with reduced hepatocellular carcinoma development and improved survival. Annals of Internal Medicine. 2005 Jan 18:142(2):105-14. PMID: 15657158

Veldt BJ, Heathcote EJ, Wedemeyer H, et al. Sustained virologic response and clinical outcomes in patients with chronic hepatitis C and advanced fibrosis. Annals of Internal Medicine. 2007;147(10):677-84. PMID: 18025443

Ware JE, Bayliss MS, Mannocchia M, et al. Health-related quality of life in chronic hepatitis C: impact of disease and treatment response. The Interventional Therapy Group. Hepatology. 1999;30(2):550-5. PMID: 10421667

Yoshida H, Arakawa Y, Sata M, et al. Interferon therapy prolonged life expectancy among chronic hepatitis C patients. Gastroenterology. 2002 Aug;123(2):483-91. PMID: 12145802

Yu M-L, Lin S-M, Chuang W-L, et al. A sustained virological response to interferon or interferon/ribavirin reduces hepatocellular carcinoma and improves survival in chronic hepatitis C: a nationwide, multicentre study in Taiwan. Antiviral Therapy. 2006;11(8):985-94. PMID: 17302368

Appendix D. Excluded Studies List

Abergel A, Achim A, Kain I, et al. Efficacy of interferon (standard or pegylated) plus ribavirin in naive patients with hepatitis C virus genotype 5. A french multicenter study. Hepatology; 2011: 817A. **Exclusion Reason -**Wrong Population

Adherence in Patients Receiving PegIntron Pen/Rebetol for Hepatitis C in Conjunction With a Patient Assistance Program (Study P04281)(COMPLETED). **Exclusion Reason** -Wrong Population

Adherence in Patients Receiving PegIntron/Rebetol for Hepatitis C in Conjunction With a Psychotherapy Support Program (Study P04252)(COMPLETED). **Exclusion Reason -** Wrong Population

Adiwijaya BS, Hare B, Caron PR, et al. Rapid decrease of wild-type hepatitis C virus on telaprevir treatment. Antiviral Therapy. 2009;14(4):591-5. PMID: 19578245. Exclusion Reason -Wrong Study Design

Adiwijaya B, Herrmann E, Hare B, et al. A Multi-Variant, Viral Dynamic Model of Genotype 1 HCV to Assess the *in vivo* Evolution of Protease-Inhibitor Resistant Variants. PLoS Comput Biol. 2010;6(4):e1000745. PMID: 20419154. Exclusion Reason -Not Relevant

Adiwijaya BS, Kieffer TL, Henshaw J, et al. A Viral Dynamic Model for Treatment Regimens with Directacting Antivirals for Chronic Hepatitis C Infection. PLoS Comput Biol. 2012;8(1):e1002339-e1002339. PMID: 22241977. Exclusion Reason -Not Relevant

Adiwijaya BS, Kieffer TL, Henshaw J, et al. A viral dynamic model for treatment regimens with direct-acting antivirals for chronic hepatitis c infection. PLoS Computational Biology. 2012;8(1). **Exclusion Reason** - Background

Afdhal NH, Dieterich DT, Pockros PJ, et al., Epoetin alfa maintains ribavirin dose in HCV-infected patients: a prospective, double-blind, randomized controlled study. Gastroenterology. 2004 May;126(5):1302-11. PMID: 15131791. Exclusion Reason -Not Relevant

Ahn & Flamm..Boceprevir versus telaprevir: A new era of directly acting antiviral therapy. Current Hepatitis Reports. 2012;11(1):23-33. **Exclusion Reason** – Background

Akuta N, Suzuki F, Hirakawa M, et al. Amino acid substitution in hepatitis C virus core region and genetic variation near the interleukin 28B gene predict viral response to telaprevir with peginterferon and ribavirin. Hepatology. 2010;52(2):421-429. PMID: 20648473. Exclusion Reason - Wrong Study Design

Akuta N, Suzuki F, Hirakawa M, et al. Amino acid substitutions in the hepatitis C virus core region of genotype 1b affect very early viral dynamics during treatment with telaprevir, peginterferon, and ribavirin. Journal of Medical Virology. 2010;82(4):575-582. PMID: 20166188. Exclusion Reason - Wrong Study Design

Akuta N, Suzuki F, Suzuki Y, et al. Long-term followup of interferon monotherapy in 454 consecutive naive patients infected with hepatitis C virus: multi-course interferon therapy may reduce the risk of hepatocellular carcinoma and increase survival. Scandinavian Journal of Gastroenterology. 2005 Jun;40(6):688-96. PMID: 16036529. Exclusion Reason -Wrong Drug

Alavian SM, Behnava B, Tabatabaei SV., et al. Comparative efficacy and overall safety of different doses of consensus interferon for treatment of chronic HCV infection: a systematic review and meta-analysis. Pharmacol Clin. 2010;66(11):1071-1071. PMID: 20857094. Exclusion Reason -Wrong Drug

Alavian SM, Jabbari, H, Daryani, NE, Hepatitis C virus the rising concerns and growing hopes, report from the HCV symposium, fourth Tehran Hepatitis Congress, November 2011, Tehran, Iran. Hepatitis Monthly. 2012;12(7):107-113. Exclusion Reason - Background

Alsiö A, Rembeck K, Askarieh G, et al.Impact of obesity on the bioavailability of peginterferon-α2a and ribavirin and treatment outcome for chronic hepatitis c genotype 2 or 3. PLoS ONE [Electronic Resource]. 2012;7(5). **Exclusion Reason** – Background

Alvarez-Uria G, Day JN, Nasir AJ, et al. Factors associated with treatment failure of patients with psychiatric diseases and injecting drug users in the treatment of genotype 2 or 3 hepatitis C chronic infection. Liver International. 2009 Aug;29(7):1051-5. PMID: 19580634. Exclusion Reason -Not Relevant

Amarapurkar DN, Patel ND, Rane P, et al. Do different hepatitis C virus genotypes behave differently? Tropical Gastroenterology. 2007 Jul-Sep;28(3):99-104. PMID: 18383996. Exclusion Reason -Not Relevant

Andersen ES, Moessner BK, Christensen PB, et al. Lower liver stiffness in patients with sustained virological response 4 years after treatment for chronic hepatitis C. European Journal of Gastroenterology & Hepatology. 2011 Jan;23(1):41-4. PMID: 21079513. Exclusion Reason -Not Relevant

Angelico M, Koehler-Horst B, Piccolo P, et al. Peginterferon alpha-2a and ribavirin versus peginterferon alpha-2a monotherapy in early virological responders and peginterferon alpha-2a and ribavirin versus peginterferon alpha-2a, ribavirin and amantadine triple therapy in early virological nonresponders: the SMIEC II trial in naive patients with chronic hepatitis C. European Journal of Gastroenterology & Hepatology. 2008 Jul;20(7):680-7. PMID: 18679072. Exclusion Reason -Not Relevant

Angelico M, Petrolati A, Lionetti R, et al. A randomized study on Peg-interferon alfa-2a with or without ribavirin in liver transplant recipients with recurrent hepatitis C. Journal of Hepatology. 2007 Jun;46(6):1009-17. PMID: 17328985. Exclusion Reason -Not Relevant

Arase Y, Suzuki F, Akuta N, et al. Combination therapy of peginterferon and ribavirin for chronic hepatitis C patients with genotype 1b and low-virus load. Internal Medicine. 2009;48(5):253-8. PMID: 19252344. Exclusion Reason -Not Relevant

Arase Y, Suzuki F, Sezaki H, et al. Efficacy in patients with dose reduction in combination therapy of peginterferon and ribavirin for chronic hepatitis C. Intervirology. 2008;51(1):1-6. PMID: 18309242. Exclusion Reason -Not Relevant

Arase Y, Suzuki F, Suzuki Y, et al. Side effects of combination therapy of peginterferon and ribavirin for chronic hepatitis-C. Internal Medicine. 2007;46(22):1827-32. PMID: 18025763. Exclusion Reason -Not Relevant

Asahina Y, Tsuchiya K, Tamaki N, et al. Effect of aging on risk for hepatocellular carcinoma in chronic hepatitis C virus infection. Hepatology. 2010 Aug;52(2):518-27. PMID: 20683951. Exclusion Reason -Not Relevant

Asselah T. A revolution in HCV treatment with directacting antivirals: From non-response to eradication. Journal of Hepatology. 2012;57(2):455-457. **Exclusion Reason -** Background

Awad T, Brok J, Thorlund K, et al. Pegylated interferon plus ribavirin versus non-pegylated interferon plus ribavirin for chronic hepatitis C. Cochrane Database of Systematic Reviews. 2009(1). **Exclusion Reason** – Background

Awad T, Thorlund K, Hauser G, et al. Pegylated interferon alpha 2a versus pegylated interferon alpha 2b for chronic hepatitis C. Cochrane Database of Systematic Reviews. 2009(1). **Exclusion Reason** - Background

Awad T, Thorlund K, Hauser G, et al. Peginterferon alpha-2a is associated with higher sustained virological response than peginterferon alfa-2b in chronic hepatitis C: Systematic review of randomized trials. Hepatology. 2010;51(4):1176-1184. PMID: 20187106. Exclusion Reason -Not Relevant

Ayaz C, Celen MK, Yuce UN, et al. Efficacy and safety of pegylated-interferon alpha-2a in hemodialysis patients with chronic hepatitis C. World Journal of Gastroenterology. 2008 Jan 14;14(2):255-9. PMID: 18186564. Exclusion Reason -Wrong Population

Backus LI, Boothroyd DB, Phillips BR, et al. Pretreatment assessment and predictors of hepatitis C virus treatment in US veterans coinfected with HIV and hepatitis C virus. Journal of Viral Hepatitis. 2006 Dec;13(12):799-810. PMID: 17109679. Exclusion Reason -Wrong Outcome

Bacon BR, Khalid O. Triple therapy with boceprevir for HCV genotype 1 infection: Phase III results in relapsers and nonresponders. Liver International. 2012;32(SUPPL. 1):51-53. **Exclusion Reason -** Background

Bain VG, Lee SS, Peltekian K, et al. Clinical trial: exposure to ribavirin predicts EVR and SVR in patients with HCV genotype 1 infection treated with peginterferon alpha-2a plus ribavirin. Alimentary Pharmacology & Therapeutics. 2008 Jul;28(1):43-50. PMID: 18397386. Exclusion Reason -Not Relevant

Balon R. Clinical factor 2011. Psychotherapy and Psychosomatics. 2012;81(4):199-205. **Exclusion Reason** - Background

Barbotte L, Ahmed-Belkacem A, Chevaliez S, et al. Characterization of V36C, a Novel Amino Acid Substitution Conferring Hepatitis C Virus (HCV) Resistance to Telaprevir, a Potent Peptidomimetic Inhibitor of HCV Protease. Antimicrobial Agents and Chemotherapy. 2010 June 2010;54(6):2681-2683. PMID: 20368394. Exclusion Reason - Wrong Outcomes

Barritt Iv AS, Fried MW. Maximizing opportunities and avoiding mistakes in triple therapy for hepatitis C virus. Gastroenterology. 2012;142(6):1314-1323.e1. **Exclusion Reason** - Background

Bartels DJ, Zhou Y, Zhang EZ, et al. Natural Prevalence of Hepatitis C Virus Variants with Decreased Sensitivity to NS3·4A Protease Inhibitors in Treatment-Naive Subjects. Journal of Infectious Diseases. 2008 September 15, 2008;198(6):800-807. PMID: 18637752. Exclusion Reason -Wrong Study Design

Berak. Randomized, open label trial comparing efficacy and safety of pegylated interferon alfa 2a vs alfa 2b treatment of patients with chronic hepatitis C infected with non 2/3 genotypes – 12 week virological response analysis. Hepatology. 2005;42(Suppl 1):1. **Exclusion Reason -**Wrong Drug

Boceprevir (SCH 503034) Plus Peg-Intron, With and Without Added Ribavirin, in Patients With Chronic Hepatitis C, Genotype 1, Who Did Not Respond to Previous Treatment With Peginterferon Alfa Plus Ribavirin.. **Exclusion Reason** -Wrong Population

Bognar F. Boceprevir in addition to standard of care enhanced SVR in hepatitis C virus (HCV) genotype-1 with advanced fibrosis/cirrhosis: Subgroup analysis of SPRINT-2 and RESPOND-2 Studies. Journal of Gastroenterology and Hepatology. 2011; 26:93. **Exclusion Reason -**Wrong Population

Bognar F. IL28B polymorphism predicts virologic response in patients with hepatitis C genotype 1 treated with boceprevir (BOC) combination therapy. Journal of Gastroenterology and Hepatology. 2011; 19-20. Exclusion Reason -Wrong Population

Bognar F. Projecting the clinical impact of therapeutic regimens including boceprevir in previously untreated adult subjects with chronic hepatitis C genotype 1. Journal of Gastroenterology and Hepatology. 2011; 19. **Exclusion Reason -**Wrong Population

Bonner, JB, Barret AS, Fried FW, et al. Tangible resources for preparing patients for antiviral therapy for chronic hepatitis C. Digestive Diseases & Sciences. 2012 Jun;57(6):1439-44. PMID: 22488633. **Exclusion Reason** - Background

Bonkovsky HL, Snow KK, Malet PF, et al. Healthrelated quality of life in patients with chronic hepatitis C and advanced fibrosis. Journal of Hepatology. 2007 Mar;46(3):420-31. PMID: 17196293. Exclusion Reason -Not Relevant

Bonkovsky HL, Tice AD, Yapp RG, et al. Efficacy and safety of peginterferon alfa-2a/ribavirin in methadone maintenance patients: randomized comparison of direct observed therapy and self-administration. American Journal of Gastroenterology. 2008 Nov;103(11):2757-65. PMID: 18684176. Exclusion Reason -Not Relevant

Borroni G, Andreoletti M, Casiraghi MA, et al. Effectiveness of pegylated interferon/ribavirin combination in 'real world' patients with chronic hepatitis C virus infection. Alimentary Pharmacology & Therapeutics. 2008 May;27(9):790-7. PMID: 18298638. Exclusion Reason -Not Relevant

Bourlière, M, Ouzan D, Rosenheim M, et al. Pegylated interferon-α2a plus ribavirin for chronic hepatitis C in a real-life setting: The Hepatys French cohort (2003-2007). Antiviral Therapy. 2012;17(1):101-110. **Exclusion Reason** – Background

Brandman, D, Bacchetti P, Ayala CE, et al. Impact of insulin resistance on HCV treatment response and impact of HCV treatment on insulin sensitivity using direct measurements of insulin action. Diabetes Care. 2012;35(5):1090-1094. **Exclusion Reason** – Background

Brennan, BJ, Morcos PN, Wang K, et al. The pharmacokinetics of peginterferon alfa-2a and ribavirin in African American, Hispanic and Caucasian patients with chronic hepatitis C. Alimentary Pharmacology & Therapeutics. 2012 May;35(10):1209-20. PMID: 22469033. Exclusion Reason - Wrong Outcomes

Brok J, Gluud LL, Gluud C, et al. Ribavirin monotherapy for chronic hepatitis C. Cochrane Database of Systematic Reviews. 2009(1). **Exclusion Reason** – Background

Brok J, Gluud LL, Gluud C, et al. Ribavirin plus interferon versus interferon for chronic hepatitis C. Cochrane Database of Systematic Reviews. 2010(5). **Exclusion Reason** - Background

Bruno R, Sacchi P, Ciappina V, et al. Viral dynamics and pharmacokinetics of peginterferon alpha-2a and peginterferon alpha-2b in naive patients with chronic hepatitis c: a randomized, controlled study. Antiviral Therapy. 2004;9(4):491-497. PMID: 15456079. **Exclusion Reason** -Wrong Outcome

Bruno R, Sacchi P, Cima S, et al. Comparison of peginterferon pharmacokinetic and pharmacodynamic profiles. Journal of Viral Hepatitis. 2012;19(1):33-36. **Exclusion Reason -** Wrong Outcomes

Bühler S, Bartenschlager R. New targets for antiviral therapy of chronic hepatitis C. Liver International. 2012;32(1):9-16. **Exclusion Reason** – Background

Burton, MJ, Passarella MJ, McGuire BM. Telaprevir and boceprevir in African Americans with genotype 1 chronic hepatitis C: Implications for patients and providers. Southern Medical Journal. 2012;105(8):431-436. **Exclusion Reason -** Wrong Study Design

Calès P, Zarski JP, Marc Chapplain J, et al. Fibrosis progression under maintenance interferon in hepatitis C is better detected by blood test than liver morphometry. Journal of Viral Hepatitis. 2012;19(2):e143-e153. **Exclusion Reason** – Background

Carruthers SJ. Hepatitis C treatment and injecting drug users in Perth, Western Australia: Knowledge of personal status and eligibility criteria for treatment. Journal of Substance Use. 2012 Feb;17(1):32-40. PMID: Peer Reviewed Journal: 2012-00263-003. Exclusion Reason – Background

Casey LC, Lee W.M. Hepatitis C therapy update. Current Opinion in Gastroenterology. 2012;28(3):188-192. Exclusion Reason - Background

Chak E, Talal AH, Sherman KE, et al. Hepatitis C virus infection in USA: an estimate of true prevalence. Liver Int; 2011. 31(8):1090-101 Exclusion Reason – Background

Chang CH, Chen KY, Lai MY, et al. Meta-analysis: ribavirin-induced haemolytic anaemia in patients with chronic hepatitis C. Alimentary Pharmacology & Therapeutics. 2002 Sep;16(9):1623-32. PMID: 12197841. Exclusion Reason -Not Relevant

Charlebois A, Lee L, Cooper E, et al. Factors associated with HCV antiviral treatment uptake among participants of a community-based HCV programme for marginalized patients. Journal of Viral Hepatitis. 2012. **Exclusion Reason -** Background

Chavalitdhamrong D, Tanwandee T. Long-term outcomes of chronic hepatitis C patients with sustained virological response at 6 months after the end of treatment. World Journal of Gastroenterology. 2006 Sep 14;12(34):5532-5. PMID: 17006994. **Exclusion Reason -**Wrong Drug

Chayama K, Hayes CN, Abe H, et al. IL28B But Not ITPA Polymorphism Is Predictive of Response to Pegylated Interferon, Ribavirin, and Telaprevir Triple Therapy in Patients With Genotype 1 Hepatitis C. Journal of Infectious Diseases. 2011 July 1, 2011;204(1):84-93. PMID: 21628662. Exclusion Reason -Wrong Outcomes

Chen L-j, Li M-h, Xie Y, et al. [Effect of hepatitis C virus serotype on the response of patients with chronic hepatitis C to interferon treatment]. Chinese Journal of Experimental & Clinical Virology. 2007
Jun;21(2):117-9. PMID: 17653309. Exclusion
Reason -Not Relevant

Chen T-M, Huang P-T, Lin C-H, et al. Feasibility of individualized treatment for hepatitis C patients in the real world. Journal of Gastroenterology & Hepatology. 2010 Jan;25(1):61-9. PMID: 19780879. Exclusion Reason -Not Relevant

Chen W-l, Chen X-p, Chen X-f, et al. [Individualized respond guidance treatment of chronic hepatitis C with combination of peginterferon -2a and ribavirin]. Chung Hua Kan Tsang Ping Tsa Chih. 2010 Aug;18(8):585-9. PMID: 20825712. Exclusion Reason -Not Relevant

Cheng WSC, Roberts SK, McCaughan G, et al. Low virological response and high relapse rates in hepatitis C genotype 1 patients with advanced fibrosis despite adequate therapeutic dosing. Journal of Hepatology. 2010 Oct;53(4):616-23. PMID: 20619475. Exclusion Reason -Not Relevant

Chevaliez, S, Hézode C., Pawlotsky, JM. Antiviral strategies in hepatitis C infection. Stratégies antivirales dans l'hépatite chronique C. 2012;14(2):78-88. **Exclusion Reason** – Background

Chu TW, Kulkarni R, Gane EJ,,et al. Effect of IL28B genotype on early viral kinetics during interferon-free treatment of patients with chronic hepatitis C. Gastroenterology. 2012;142(4):790-795. **Exclusion Reason** - Background

Ciancio A, Picciotto A, Giordanino C, et al. A randomized trial of pegylated-interferon-alpha2a plus ribavirin with or without amantadine in the retreatment of patients with chronic hepatitis C not responding to standard interferon and ribavirin. Alimentary Pharmacology & Therapeutics. 2006 Oct 1;24(7):1079-86. PMID: 16984502. Exclusion Reason -Not Relevant

Comparison of Safety and Resulting Blood Level Profiles After Administration of a New Boceprevir Tablet Versus Its Current Capsule Formulation for Treatment of Chronic Hepatitis C. 2010. **Exclusion Reason** - Background

Dalgard O, Bjoro K, Hellum KB, et al. Treatment with pegylated interferon and ribavarin in HCV infection with genotype 2 or 3 for 14 weeks: a pilot study. Hepatology. 2004 Dec;40(6):1260-5. PMID: 15558712. Exclusion Reason -Not Relevant

Dalgard O, Bjoro K, Ring-Larsen H, et al. In patients with HCV genotype 2 or 3 infection and RVR 14 weeks treatment is noninferior to 24 weeks. Pooled analysis of two Scandinavian trials. European Journal of Gastroenterology & Hepatology. 2010 May;22(5):552-6. PMID: 20154627. Exclusion Reason -Not Relevant

Dan AA, Crone C, Wise TN, et al. Anger experiences among hepatitis C patients: relationship to depressive symptoms and health-related quality of life. Psychosomatics. 2007 May-Jun;48(3):223-9. PMID: 17478591. Exclusion Reason -Not Relevant

Daw MA, Dau AA. Hepatitis C virus in Arab world: A state of concern. The Scientific World Journal. 2012;2012. **Exclusion Reason** – Background

De Azevedo FKSF, de Azevedo CCSF, Souto FJD. Souto. Assessment of the treatment of chronic hepatitis C in the state of mato grosso, central Brazil. Memorias do Instituto Oswaldo Cruz. 2012;107(2):117-123. Exclusion Reason - Background

de Bruijne J, Bergmann JF, Reesink HW, et al. Antiviral activity of narlaprevir combined with ritonavir and pegylated interferon in chronic hepatitis C patients. Hepatology. 2010 Nov;52(5):1590-9. PMID: 20938912. Exclusion Reason -Not Relevant

De Bruijne J, Van Vliet A, Weegink CJ, et al. Rapid decline of viral RNA in chronic hepatitis C patients treated once daily with IDX320: A novel macrocyclic HCV protease inhibitor. Antiviral Therapy. 2012;17(4):633-642. **Exclusion Reason -** Background

De Rosa FG, Bargiacchi O, Audagnotto S, et al.. Dose-dependent and genotype-independent sustained virological response of a 12 week pegylated interferon alpha-2b treatment for acute hepatitis C. Journal of Antimicrobial Chemotherapy. 2006 Feb;57(2):360-3. PMID: 16396921. Exclusion Reason -Not Relevant

De-Rueda PM, Ruiz-Extremera A, Candel JM, et al. Plasma Ribavirin trough concentrations during treatment of chronic hepatitis C in genotype-1 patients. Journal of Clinical Gastroenterology. 2012;46(4):328-333. **Exclusion Reason** - Background

Derbala MF, Al Kaabi SR, El Dweik NZ, et al. Treatment of hepatitis C virus genotype 4 with peginterferon alfa-2a: impact of bilharziasis and fibrosis stage. World Journal of Gastroenterology. 2006 Sep 21;12(35):5692-8. PMID: 17007024. **Exclusion Reason** -Not Relevant

Desai, TK, Bortman J, Al-Sibae R, et al. The role of iron in hepatitis C infection. Current Hepatitis Reports. 2012;11(1):41-47. **Exclusion Reason** - Background

Di Bisceglie AM, Ghalib RH, Hamzeh FM, et al. Early virologic response after peginterferon alpha-2a plus ribavirin or peginterferon alpha-2b plus ribavirin treatment in patients with chronic hepatitis C. Journal of Viral Hepatitis. 2007 Oct;14(10):721-9. PMID: 17875007. Exclusion Reason -Not Relevant

Di Bisceglie AM, Shiffman ML, Everson GT, et al. Prolonged therapy of advanced chronic hepatitis C with low-dose peginterferon. New England Journal of Medicine. 2008 Dec 4;359(23):2429-41. PMID: 19052125. Exclusion Reason - Wrong Outcomes

Di Bisceglie AM, Dusheiko GM, Muir AJ, et al. Telaprevir in combination with peginterferon alfa-2a and ribavirin: Analyses of pre-defined subpopulations in the phase 3 advance trial. Gastroenterology; 2011a. p. S908. **Exclusion Reason -**Wrong Population

Diago M, Olveira A, Sola R, et al. Treatment of chronic helpatitis C genotype 1 with peginterferonalpha2a (40 kDa) plus ribavirin under routine clinical practice in Spain: early prediction of sustained virological response rate. Alimentary Pharmacology & Therapeutics. 2007 Apr 15;25(8):899-906. PMID: 17402993. Exclusion Reason -Not Relevant

Diago M, Shiffman ML, Bronowicki J-P, et al. Identifying hepatitis C virus genotype 2/3 patients who can receive a 16-week abbreviated course of peginterferon alfa-2a (40KD) plus ribavirin. Hepatology. 2010 Jun;51(6):1897-903. PMID: 20196118. Exclusion Reason -Not Relevant

Dieterich DT, Wasserman R, Brau N, et al. Onceweekly epoetin alfa improves anemia and facilitates maintenance of ribavirin dosing in hepatitis C virusinfected patients receiving ribavirin plus interferon alfa. American Journal of Gastroenterology. 2003 Nov;98(11):2491-9. PMID: 14638354. Exclusion Reason -Not Relevant

Druyts E, Mills EJ, Nachega J, et al. Differences in clinical outcomes among hepatitis C genotype 1-infected patients treated with peginterferon alpha-2a or peginterferon alpha-2b plus ribavirin: A meta-analysis. Clinical and Experimental Gastroenterology. 2012; 5(1): 11-21. PMID:22427726 Exclusion Reason - Wrong Population

Ebner N, Wanner C, Winklbaur B, et al. Retention rate and side effects in a prospective trial on hepatitis C treatment with pegylated interferon alpha-2a and ribavirin in opioid-dependent patients. Addiction Biology. 2009 Apr;14(2):227-37. PMID: 19291011. **Exclusion Reason -**Not Relevant

Effects of 48 Weeks Versus 24 Weeks of Therapy With Peg-Intron/Ribavirin in Patients With Chronic Hepatitis C, Genotype 3 (Study P04143)(TERMINATED). **Exclusion Reason -**Wrong Population

Efficacy and Safety of 24 vs 48 Weeks of Pegetron® (Peginterferon Alfa-2b + Ribavirin) in Naïve Genotype 1 Hepatitis C (Study P05016)(TERMINATED). Exclusion Reason -Wrong Population

Efficacy and Safety of Peginterferon Alfa-2b and Ribavirin Therapy in Subjects With Type C Compensated Liver Cirrhosis (Study P05116). **Exclusion Reason -**Wrong Population

El-Serag HB, Mason AC. Rising Incidence of Hepatocellular Carcinoma in the United States. New England Journal of Medicine. 1999; 340(10): 745-50. **Exclusion Reason** – Background

El-Serag HB, Mason AC. Risk factors for the rising rates of primary liver cancer in the United States. Archives of Internal Medicine. 2000; 160(21):3227-30. **Exclusion Reason -** Background

El-Serag HB, Davila JA, Petersen NJ, et al. The continuing increase in the incidence of hepatocellular carcinoma in the United States: an update. Annals of Internal Medecine. 2003; 139(10): 817-23. **Exclusion Reason** – Background

El-Zayadi AR, Attia M, Barakat EM, et al. Response of hepatitis C genotype-4 naive patients to 24 weeks of Peg-interferon-alpha2b/ribavirin or induction-dose interferon-alpha2b/ribavirin/amantadine: a non-randomized controlled study. American Journal of Gastroenterology. 2005 Nov;100(11):2447-52. PMID: 16279899. Exclusion Reason -Not Relevant

Everhart JE, Lok AS, Kim H-Y, et al. Weight-related effects on disease progression in the hepatitis C antiviral long-term treatment against cirrhosis trial. Gastroenterology. 2009 Aug;137(2):549-57. PMID: 19445938. Exclusion Reason -Not Relevant

Everson GT, Balart L, Lee SS, et al. Histological benefits of virological response to peginterferon alfa-2a monotherapy in patients with hepatitis C and advanced fibrosis or compensated cirrhosis.

Alimentary Pharmacology & Therapeutics. 2008 Apr 1;27(7):542-51. PMID: 18208570. Exclusion Reason -Wrong Outcomes

Everson GT, Shiffman ML, Hoefs JC, et al. Quantitative tests of liver function measure hepatic improvement after sustained virological response: results from the HALT-C trial. Alimentary Pharmacology & Therapeutics. 2009 Mar 1;29(5):589-601. PMID: 19053983. Exclusion Reason -Not Relevant

Extended Treatment With PEG-Intron® and Rebetol® in Patients With Genotype 1 Chronic Hepatitis C and Slow Virologic Response (Study P03685AM3)(COMPLETED). Exclusion Reason - Wrong Population

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Appendix E. Quality Assessment Methods

Individual studies were rated as "good," "fair" or "poor" as defined below¹:

For Controlled Trials:

Each criterion was give an assessment of yes, no, or unclear.

1. Was the assignment to the treatment groups really random?

Adequate approaches to sequence generation:

Computer-generated random numbers

Random numbers tables

Inferior approaches to sequence generation:

Use of alternation, case record numbers, birth dates or week days

Randomization reported, but method not stated

Not clear or not reported

Not randomized

2. Was the treatment allocation concealed?

Adequate approaches to concealment of randomization:

- Centralized or pharmacy-controlled randomization (randomization performed without knowledge of patient characteristics).
- Serially-numbered identical containers
- On-site computer based system with a randomization sequence that is not readable until allocation
- Sealed opaque envelopes

Inferior approaches to concealment of randomization:

- Use of alternation, case record numbers, birth dates or week days
- Open random numbers lists
- Serially numbered non- opaque envelopes
- Not clear or not reported
- 3. Were the groups similar at baseline in terms of prognostic factors?
- 4. Were the eligibility criteria specified?
- 5. Were outcome assessors and/or data analysts blinded to the treatment allocation?
- 6. Was the care provider blinded?
- 7. Was the patient kept unaware of the treatment received?
- 8. Did the article include an intention-to-treat analysis, or provide the data needed to calculate it (i.e., number assigned to each group, number of subjects who finished in each group, and their results)?
- 9. Did the study maintain comparable groups?
- 10. Did the article report attrition, crossovers, adherence, and contamination?
- 11. Is there important differential loss to followup or overall high loss to followup?

For Cohort Studies:

Each criterion was give an assessment of yes, no, or unclear.

- 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 use accurate methods for ascertaining exposures, potential confounders, and outcomes?
- 4. Were outcome assessors and/or data analysts blinded to treatment?
- 5. Did the article report attrition?
- 6. Did the study perform appropriate statistical analyses on potential confounders?
- 7. Is there important differential loss to followup or overall high loss to followup?
- 8. Were outcomes pre-specified and defined, and ascertained using accurate methods?

Appendix F. Sustained Virologic Response and Quality of Life

Author, Year Country Quality	Comparison Definition of Sustained Virologic Response	Population Characteristics	Treatments	Results (by clinical outcome)
Arora , 2006 ¹ Australia, Europe, New Zealand, North America, and South America Quality: Poor	SVR vs. no SVR SVR=No detectable HCV RNA at end of followup (72 weeks)	Not reported by SVR status Mean age: 43 years Female: 60% Race: Non-white: 14% Advanced fibrosis: 10% Genotype 1: 68% Viral load: 1.1-1.2 x 10 ⁶ copies/ml IVDU: 30% HIV positive: excluded	Pegylated interferon alfa-2a (24 or 48 weeks)	SVR vs. no SVR, mean difference in change from baseline SF-36 physical function: +4.7 (p<0.05) SF-36 role limitations-physical: +13 (p<0.05) SF-36 bodily pain: +11 (p<0.0001) SF-36 general health: +10 (p<0.0001) SF-36 vitality: +9.3 (p<0.0001) SF-36 social function: +5.1 (p>0.05) SF-36 role limitations-emotional: +7.3 (p>0.05) SF-36 mental health: +3.1 (p>0.05) SF-36 physical component summary: +4.9 (p<0.0001) SF-36 mental component summary: +2.0 (p>0.05) Fatigue Severity Scale, total score: -4.4 (p<0.01)
				Fatigue Severity Scale, VAS: -10 (p<0.01)

Author, Year Country Quality	Comparison Definition of Sustained Virologic Response	Population Characteristics	Treatments	Results (by clinical outcome)
Bernstein , 2002 ² Australia, North America, Europe, Taiwan, New Zealand Quality: Poor	SVR vs. no SVR SVR=No detectable HCV RNA 24 weeks after completion of antiviral therapy	Not reported by SVR status Mean age <=40 years: 41% Female: 32% Race: Non-white: 14% Cirrhosis: 32% Genotype, viral load, HIV infection, IV drug use not reported	Pegylated interferon alfa-2a or interferon alfa-2a	SVR vs. no SVR, mean difference in change from baseline SF-36 physical function: +4.6 (p<0.001) SF-36 role limitations-physical: +9.8 (p<0.001) SF-36 bodily pain: +2.9 (p<0.01) SF-36 general health: +9.1 (p<0.001) SF-36 vitality: +9.6 (p<0.001) SF-36 social function: +6.2 (p<0.001) SF-36 role limitations-emotional: +8.4 (p<0.01) SF-36 mental health: +4.6 (p<0.001) SF-36 physical component summary: +2.8 (p<0.001) SF-36 mental component summary: +3.0 (p<0.001) Fatigue Severity Scale, total score: -0.5 (p<0.001) Fatigue Severity Scale, VAS: -11.5 (p<0.001)

Author, Year	Comparison			
Country	Definition of Sustained			
Quality	Virologic Response	Population Characteristics	Treatments	Results (by clinical outcome)
Bini, 2006 ³ USA	SVR vs. no SVR SVR=No detectable HCV RNA 24 weeks after	Normal ALT and elevated ALT groups, respectively (not reported by SVR status)	Interferon alfa-2b + ribavirin	SVR vs. no SVR, mean difference in change from baseline (normal ALT and elevated ALT subgroups, respectively;
Quality: Poor	completion of antiviral therapy	Mean age: 50 and 49 years Female: 11% and 8%		p values not reported) SF-36 physical function: +18 and +15 SF-36 role limitations-physical: +22 and +27
		Race: Non-white: 59% and 66%		SF-36 bodily pain: +3.4 and +9.3 SF-36 general health: +3.0 and +9.9 SF-36 vitality: +12 and +12
		Normal ALT and elevated ALT groups, respectively (not reported by SVR status) Cirrhosis: 11% and 11%		SF-36 social function: +9.5 and +11 SF-36 role limitations-emotional: +20 and +18 SF-36 mental health: +14 and +18 SF-36 physical component summary:
		Genotype 1: 78% and 78% Viral load >2 x 10 ⁶ copies/ml:		+3.8 and +7.1 SF-36 mental component summary: +6.0 and +2.1 Positive well being: +14 and -3.1
		44% and 44% IVDU: 67% and 65%		Sleep somnolence: +11 and +5.4 Health distress: +9.3 and +11 Hepatitis-specific health distress: +5.4
		HIV positive: excluded		and +2.6 Hepatitis-specific limitations: +13 and +3.8
Bonkovsky , 1999 ⁴ USA and Canada	SVR vs. no SVR SVR=No detectable HCV RNA 24 weeks after	Not reported by SVR status Mean age: 43 years	Consensus interferon or interferon alfa-2b	SVR vs. no SVR, mean difference in change from baseline (values estimated from graph)
Quality: Poor	completion of antiviral therapy	Female: 27%		SF-36 physical function: +6.0 (p<0.05) SF-36 role limitations-physical: +22 (p<0.01)
		Race: Non-white: 23%		SF-36 bodily pain: -0.5 (p>0.05) SF-36 general health: +7.5 (p<0.01) SF-36 vitality: +9.5 (p<0.05)
		Cirrhosis: 16%		SF-36 social function: +10 (p<0.05) SF-36 role limitations-emotional: +11
		Genotype 1: 68%		(p>0.05) SF-36 mental health: +4.0 (p>0.05)
		Viral load: Not reported IVDU: 41%		
		HIV positive: excluded		

Author, Year Country Quality	Comparison Definition of Sustained Virologic Response	Population Characteristics	Treatments	Results (by clinical outcome)
Hassanein, 2004 ⁵	SVR vs. no SVR	Not reported by SVR status	Pegylated interferon	SVR vs. no SVR, mean difference in
Australia, North America,	SVR=No detectable HCV		alfa-2a, pegylated	change from baseline
Europe, Taiwan, Brazil,	RNA 24 weeks after	Mean age: 43 years	interferon alf-2a	SF-36 physical function: +5.5 (p<0.01)
Mexico	completion of antiviral therapy	Female: 29%	+ribavirin, or interferon alfa-2b +	SF-36 role limitations-physical: +5.7 (p<0.05)
Quality: Poor		Female. 29%	ribavirin	SF-36 bodily pain: +4.1 (p<0.05)
Quality. 1 001		Race:	IIDAVIIII	SF-36 general health: +8.6 (p<0.01)
		Non-white: 16%		SF-36 vitality: +6.3 (p >0.05)
				SF-36 social function: +5.8 (p<0.01)
		Cirrhosis: 13%		SF-36 role limitations-emotional: +9.3
				(p<0.01)
		Genotype 1: 63%		SF-36 mental health: +5.0 (p<0.01)
		\" \\ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \		SF-36 physical component summary:
		Viral load: 5.9 to 6.0 x 10 ⁶		+2.2 (p<0.01)
		copies/ml IVDU: Not reported		SF-36 mental component summary: +2.6 (p<0.01)
		1VDO. Not reported		Total fatigue: +3.3 (p<0.01)
		HIV positive: excluded		Fatigue severity: +7.4 (p<0.01)
McHutchison, 2001 ⁶	SVR vs. relapse vs. non-	Mean age: 43 vs. 44 years	Interferon alfa-2a for	SVR and relapse, mean difference in
USA	responder		24 or 48 weeks, with	change from baseline vs. non-
	SVR=No detectable HCV	Female: 42% vs. 32%	or without ribavirin	responder (p not reported, values
Quality: Poor	RNA 24 weeks after			estimated from graph)
	completion of antiviral therapy	Race:		SF-36 physical function: +2.4 and +0.8
	Relapse: Not defined	Non-white: 8% vs. 12%		SF-36 role limitations-physical: +5.2 and +3.2
		Cirrhosis: Not reported		SF-36 bodily pain: +1.6 and +1.7
		Cirriosis. Not reported		SF-36 general health: +5.2 and +1.5
		Genotype 1: 43% vs. 81%		SF-36 vitality: +4.7 and +2.0
				SF-36 social function: +3.1 and +0.4
		Viral load >2 million copies/ml:		SF-36 role limitations-emotional: +3.0
		58% vs. 74%		and +1.2
		IVDU: Not reported		SF-36 mental health: +2.0 and 0.0
		11007 110		Sleep somnolence: +3.4 and +2.3
		HIV positive: excluded		Health distress: +5.4 and +1.2
				Hepatitis-related health distress: +5.7 and +1.1
				Hepatitis-related limitations: +4.6 and
				+2.1

Author, Year	Comparison			
Country	Definition of Sustained			
Quality	Virologic Response	Population Characteristics	Treatments	Results (by clinical outcome)
Neary , 1999 ⁷ USA, Europe, Australia Quality: Poor	SVR vs. no SVR and overall response versus no overall response SVR=No detectable HCV RNA 24 weeks after completion of antiviral therapy Overall response=SVR plus >=2-point improvement in Knodell HAI score	Not reported by SVR or overall response status Mean age: 43 years Female: 35% Race: Non-white: 6.4% Not reported by SVR or overall response status Bridging fibrosis or cirrhosis: 17% Genotype 1: 56% Viral load >2 million copies/ml: 75% IVDU: 40% HIV positive: excluded	Interferon alfa-2b with or without ribavirin	SVR and relapse. mean difference in change from baseline vs. non-responder (estimated from graph) (p values not reported) SF-36 physical function: +8.0 and +3.8 SF-36 role limitations-physical: +7.6 and +4.9 SF-36 bodily pain: +2.4 and +2.7 SF-36 general health: +9.4 and +5.6 SF-36 vitality: +7.8 and +5.6 SF-36 social function: +9.4 and +4.1 SF-36 role limitations-emotional: +6.0 and +12 SF-36 mental health: +2.8 and +1.8 Sleep somnolence: +2.1 and +3.8 Health distress: +8.9 and +1.6 Hepatitis-related health distress: +11 and -0.8 Hepatitis-related limitations: +6.7 and +2.6 Mental health-18: +3.4 and +2.3 Overall response vs. no response (estimated from graph) SF-36 physical function: +8.3 (p<0.05) SF-36 social function: +8.3 (p<0.05) SF-36 social function: +9.2 (p<0.05) SF-36 social function: +9.2 (p<0.05) SF-36 role limitations-emotional: +3.6 (p>0.05) SF-36 mental health: +1.3 (p>0.05) SF-36 mental health: +1.3 (p>0.05) Hepatitis-related health distress: +12 (p<0.05) Hepatitis-related limitations: +7.8 (p<0.05) Mental health-18: +1.5 (p>0.05) Mental health-18: +1.5 (p>0.05)

Author, Year Country Quality	Comparison Definition of Sustained Virologic Response	Population Characteristics	Treatments	Results (by clinical outcome)
Rasenack , 2003 ⁸ Germany, Canada, New Zealand, Spain Quality: Poor	SVR vs. no SVR SVR=No detectable HCV RNA 24 weeks after completion of antiviral therapy	Not reported by SVR status Mean age: 41 years Female: 33% Race: Non-white: 15% Bridging fibrosis/cirrhosis: 13% Injection drug use: 37% Viral load: 7.4 to 8.2 x 10 ⁶ copies/ml HIV positive: Not reported Genotype: Not reported	Pegylated interferon alfa-2a or interferon alfa-2a	SVR vs. no SVR, mean difference in change from baseline SF-36 physical function: +5.0 (p=0.001) SF-36 role limitations-physical: +14 (p<0.001) SF-36 bodily pain: +5.2 (p=0.014) SF-36 general health: 12 (p<0.001) SF-36 vitality: +9.4 (p<0.001) SF-36 social function: +5.8 (p=0.005) SF-36 role limitations-emotional: +8.4 (p=0.02) SF-36 mental health: +5.3 (p=0.001) SF-36 physical component summary: +3.2 (p<0.001) SF-36 mental component summary: +2.9 (p=0.005) Fatigue Severity Scale, total score: -0.5 (p=0.001) Fatigue Severity Scale, VAS: -8.4 (p<0.001)
Ware , 1999 ⁹ Australia, North America, and Europe Quality: Poor	SVR vs. no SVR SVR=No detectable HCV RNA 24 weeks after completion of antiviral therapy Overall response vs. no overall response Overall response=SVR + Knodell histology activity index inflammation score improved by 2 U or more	Not reported by response status Mean age: 43 years Female: 35% Race: Non-white: 6.4% Bridging fibrosis/cirrhosis: 18% Injection drug use: 40% Viral load: 4.8 to 5.2 x 10 ⁶ copies/ml HIV positive: Excluded Genotype 1: 56%	Interferon alfa-2b or interferon alfa-2b + ribavirin	SVR vs. no SVR and overall response vs. no overall response, mean difference in change from baseline (p values not reported) SF-36 physical function: +2.6 and +3.5 SF-36 role limitations-physical: +1.5 and +3.1 SF-36 bodily pain: +0.45 and +1.6 SF-36 general health: +3.3 and +3.5 SF-36 vitality: +2.2 and +2.8 SF-36 social function: +3.4 and +4.3 SF-36 role limitations-emotional: -0.02 and +1.1 SF-36 mental health: +1.3 and +0.62 Sleep: +0.02 and +1.2 Health distress: +7.6 and +6.2 Chronic hepatitis C health distress: +11.5 and +11.3 Chronic hepatitis C limitations: +5.3 and +7.5

Abbreviations: ALT, alanine aminotransferase; HCV, hepatitis C virus; SVR, sustained virologic response.

Appendix F References

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Appendix G. Overall Strength of Evidence

Key Question	Number of Studies	Quality (Good, Fair, Poor)	Consistency (High, Moderate, Low)	Directness (Direct or Indirect)	Precision (High, Moderate, Low)	Number of Subjects	Strength of Evidence
1a. What is the comparative effectiveness of antiviral treatment in improving health outcomes in patients with HCV infection?	Number of Studies	1 001)	Low	murrecty	Liow)	Subjects	Evidence
Long-term clinical outcomes	No studies	No studies	Unknown (no studies)	No studies	No studies	No subjects	Insufficient
Short-term mortality	3 randomized trials	Fair	High	Direct	Low	N = 5,255	Low
Short-term quality of life	1 randomized trial	Fair	Unknown (one study)	Direct	Low	N = 516	Low
effectiveness of antiviral treatment for health outcomes vary according to patient subgroup characteristics, including but not limited to HCV genotype, race, sex, disease severity or genetic markers?							
Any clinical outcome 2a. What is the comparative effectiveness of antiviral treatments in improving intermediate outcomes, such as the rate of viremia, aminotransaminase levels, and histologic changes?	No studies	No studies	No studies	No studies	No studies	No subjects	Insufficient
SVR: Dual therapy with pegylated interferon alfa-2a plus ribavirin vs. pegylated interferon alfa-2b plus ribavirin	7 randomized trials	Fair	High	Direct	High	N = 4,660	Moderate

Key Question	Number of Studies	Quality (Good, Fair, Poor)	Consistency (High, Moderate, Low)	Directness (Direct or Indirect)	Precision (High, Moderate, Low)	Number of Subjects	Strength of Evidence
Duration effects, dual therapy with pegylated interferon plus ribavirin (genotype 2 or 3)			,				
SVR: 48 vs. 24 weeks	2 randomized trials	Fair	High	Direct	Moderate	N = 609	Moderate
SVR: 24 vs. 12-16 weeks	4 randomized trials	Fair	High	Direct	Moderate	N = 2,599	Moderate
SVR: 24 vs. 12-16 weeks in patients with rapid virological response	3 randomized trials	Fair	High	Direct	Moderate	N = 583	Moderate
Dose effects, dual therapy with pegylated interferon plus ribavirin (genotype 2 or 3)							
SVR: Lower vs. higher dose pegylated interferon	6 randomized trials	Fair	High	Direct	Moderate	N = 865	Moderate
SVR: Lower vs. higher dose ribavirin	3 randomized trials	Fair	Moderate	Direct	Moderate	N = 2,605	Moderate
SVR: Lower vs. higher dose ribavirin, patients with advanced fibrosis or cirrhosis	1 randomized trial	Fair	Unknown (one study)	Direct	Low	N = 60	Low
Triple therapy with boceprevir SVR: Triple therapy with boceprevir vs. dual therapy	2 randomized trials	Fair	High	Direct	Moderate	N = 1608	Moderate
SVR: Lower vs. higher dose ribavirin	1 randomized trial	Fair	Unknown (one study)	Direct	Low	N = 75	Low
Triple therapy with telaprevir							
SVR: 24 weeks fixed duration triple therapy with telaprevir vs. 48 weeks dual therapy	3 randomized trials	Fair	High	Direct	Moderate	N= 506	Moderate
SVR: 12 weeks fixed duration triple therapy with telaprevir vs. 48 weeks dual therapy	1 randomized trial	Fair	Unknown (one study)	Direct	Low	N = 209	Low
SVR: 48 weeks fixed duration triple therapy with telaprevir vs. 24 weeks triple therapy	1 randomized trial	Fair	Unknown (one study)	Direct	Low	N = 189	Low

Key Question	Number of Studies	Quality (Good, Fair, Poor)	Consistency (High, Moderate, Low)	Directness (Direct or Indirect)	Precision (High, Moderate, Low)	Number of Subjects	Strength of Evidence
SVR: Response-guided triple therapy with telaprevir vs. dual therapy	1 randomized trial	Fair	Unknown (one study)	Direct	Low	N = 1,088	Low
SVR: Triple therapy with telaprevir, lower versus higher telaprevir dose and pegylated interferon alfa-2a vs. alfa-2b	1 randomized trial	Fair	Unknown (one study)	Direct	Low	N = 161	Low
SVR: 48 vs. 24 weeks in patients with an extended rapid virological response	1 randomized trial	Fair	Unknown (one study)	Direct	Low	N = 540	Low
treatment for intermediate outcomes vary according to patient subgroup characteristics, including but not limited to HCV genotype, race, sex, disease severity or genetic markers?							
SVR: Dual therapy with pegylated interferon alfa-2a plus ribavirin vs. dual therapy with pegylated interferon alfa-2b plus ribavirin: effects of race, sex, age, baseline fibrosis stage, or baseline viral load	1 randomized trial	Fair	Unknown (one study)	Direct	Moderate	N = 3070	Low
SVR: Dual therapy with pegylated interferon alfa-2a plus ribavirin vs. dual therapy with pegylated interferon alfa-2b plus ribavirin: effects of genotype	4 randomized trials	Fair	High	Direct	High	N = 1,152	Moderate
SVR: Triple therapy with boceprevir vs. dual therapy: effects of sex and race	2 randomized trials	Fair	High	Direct	Moderate	N = 1,617	Moderate
SVR: Triple therapy with boceprevir vs. dual therapy: effects of baseline viral load	2 randomized trials	Fair	High	Direct	Moderate	N = 1,617	Moderate
SVR: Triple therapy with telaprevir vs. dual therapy: effects of age, sex, race, baseline fibrosis, and body weight	1 randomized trial	Fair	Unknown (1 study)	Direct	Moderate	N = 1,088	Moderate (for age and sex) to low (for other factors)

Key Question SVR: Triple therapy with telaprevir vs. dual therapy: effects of baseline viral load	Number of Studies 2 randomized trials	Quality (Good, Fair, Poor) Fair	Consistency (High, Moderate, Low)	Directness (Direct or Indirect) Direct	Precision (High, Moderate, Low) Moderate	Number of Subjects N = 729	Strength of Evidence Insufficient
3a. What are the comparative harms (including intolerance to treatment) associated with antiviral treatment?							
Harms: Dual therapy with pegylated interferon alfa-2b plus ribavirin vs. pegylated interferon alfa-2a plus ribavirin	5 randomized trials, depending on specific harm	Fair	High	Direct	Moderate	N = 4,047	Moderate
Harms: Triple therapy with boceprevir	2 randomized trials	Fair	High	Direct	Moderate	N = 3,501	Moderate
Harms: 24 weeks fixed duration triple therapy with telaprevir vs. 48 weeks dual therapy	3 randomized trials	Fair	High	Direct	Moderate	N = 3,591	Moderate
Harms: 12 weeks fixed duration triple therapy with telaprevir vs. 48 weeks dual therapy	2 randomized trials	Fair	High	Direct	Moderate	N = 573	Moderate
Harms: Response-guided triple therapy with telaprevir vs. dual therapy	1 randomized trial	Fair	Unknown (one study)	Direct	Low	N = 189	Low
3b. Do these harms differ according to patient subgroup characteristics, including HCV genotype, race, sex, disease severity or genetic markers?							
Dual therapy with pegylated interferon alfa-2b plus ribavirin vs. pegylated interferon alfa-2a plus ribavirin	3 randomized trials	Fair	High	Indirect (no study stratified harms by patient subgroups, 3 trials evaluated only genotype 1 patients)	Moderate	N = 3,305	Insufficient
Triple therapy with pegylated interferon, ribavirin, and telaprevir or boceprevir	No studies	No studies	Unknown (no studies)	No studies	No studies	No subjects	Insufficient

Key Question	Number of Studies	Quality (Good, Fair, Poor)	Consistency (High, Moderate, Low)	Directness (Direct or Indirect)	Precision (High, Moderate, Low)	Number of Subjects	Strength of Evidence
4. Have improvements in							
intermediate outcomes (viremia,							
liver function tests, histologic							
changes) been shown to reduce							
the risk or rates of health							
outcomes from HCV infection?							
Mortality and long-term hepatic	19 cohort studies	Fair	High	Direct	High	N = 27,992	Moderate
complications							
Short-term quality of life	9 cohort studies	Poor	High	Direct	High	N = 4,981	Low

Note: HCV=hepatitis C virus, SVR=sustained virologic response.

Appendix H. Evidence Tables and Quality Ratings Key Questions 2a - 3b

Evidence Table 1. Trials of dual therapy with pegylated interferon alpha-2a plus ribavirin compared with pegylated interferon alfa-2b plus ribavirin

Author, Year Country Study Name Ouality	Interferon Regimen	Ribavirin Regimen	Protease Inhibitor Regimen	Eligibility	Exclusion	Number Screened/ Eligible/ Enrolled/ Analyzed	Age Sex Race
Ascione, 2010¹ Liver Unit of Cardarelli Hospital - Napoli, Italy Pegylated Interferon alfa-2a plus Ribavirin is more effective than Pegylated Interferon alfa-2b plus Ribavirin for treating chronic HCV Infection Overall Quality: Fair	A: Pegylated interferon alpha-2a 180 µg/week for 24 or 48 weeks (genotype 2/3 and 1/4 respectively) B: Genotype 2/3: Pegylated interferon alpha-2b 1.5 µg/kg/week for 24 or 48 weeks (genotype 2/3 and 1/4 respectively)	A: 800-1200 mg daily for 24 or 48 weeks (genotype 2/3 and 1/4 respectively) B: 800-1200 mg daily for 24 or 48 weeks (genotype 2/3 and 1/4 respectively)	None	Detectable serum HCV RNA level ALT level 1.5x the upper limit of normal for 6 months Liver biopsy within 12 months of starting treatment graded according to Scheuer's criteria (2002) Negative pregnancy test result/using Contraceptive methods during therapy and for 6 months after the end of treatment No alcohol use 6 months pre-enrollment Cirrhosis on basis of clinical/lab testing liver-spleen ultrasonography Upper gastrointestinal endoscopy for patients who did not have a biopsy	Hemoglobin level <120 g/L Neutrophil count <1.5x10 ⁹ /L or a platelet count <70x10 ⁹ /L Abnormal serum creatinine level; Hepatitis B surface antigen positive HIV+ Any other cause of liver disease History of liver decompensation Clinically relevant depression or any other Psychiatric disease Cancer Severe cardiac/pulmonary/renal disease Uncontrolled diabetes or severe hypertension with vascular complications including Retinopathy	408/322/320/320	A vs. B Age (mean): 51 vs. 49 years Female: 49% vs. 61% Race: Not reported Cirrhosis: 21% vs. 16%4% (overall) Minimal or no fibrosis: Not reported Elevated transaminases: 100% (mean ALT 2.4 vs. 2.4 upper limit of normal)

Author, Year Country Study Name Quality	Naïve	Duration of Followup	Outcome	Subgroup Analyses	Subgroup Analyses	Histologic Response	Adverse Events	Funding Source
Ascione, 2010 ¹	A vs. B	Followup at 3 and		NR	A vs. B	NR	A vs. B	Carderelli
Liver Unit of Cardarelli Hospital – Napoli, Italy	Genotype 1/4 - 93/160(58%) vs. 93/160(58%)	6 months post- treatment (12 and 24 weeks)	ETR: 134/160(83.8%) vs. 103/160(64.4%), p≤0.0001		Genotype 1/4 - 51/93(54.8%) vs. 37/93(39.8%), p=0.04		Overall Withdrawals: 4/160(3%) vs. 22/160(14%) Withdrawals due to adverse	Hospital, Napoli, Italy
Continued	Genotype 2/3 - 67/160(42%) vs. 67/160(42%) Severity by liver biopsy graded via "simple system" (Scheuer et al 2002): Chronic Hepatitis: 127/160(79.4%) vs. 134/160(83.7%) Cirrhosis (with biopsy): 33/160(20.6%) vs. 26/160(16.3%) Cirrhosis (without biopsy): 12/160(7.5%) vs. 7/160(4.4%) Treatment-naïve: 100%	24 weeks)	SVR: 110/160(68.8%) vs. 87/160(54.4%), p=0.008		Genotype 2/3 - 59/67(88.1%) vs. 50/67(74.6%), p=0.046 Genotype 2 - 45/49(91.8%) vs. 38/50(76.0%), p=0.062 Genotype 3 - 14/18(77.8%) vs. 12/17(70.6%), p=0.92 Chronic hepatitis - 96/127(75.6%) vs. 75/134(55.9%), p=0.005 Cirrhosis - 14/33(42.4%) vs. 12/26(46.1%), p=0.774 SVR by baseline Genotype RNA level in serum, no./total (%): <500,000 IU/mL - 52/76(68.4%) vs. 44/67(65.7%), p=0.727 >500,000 IU/mL - 58/84(69.0%) vs. 43/93(46.2%), p=0.002		events; 4/160 (3%) vs. 17/160 (11%) Deaths: none Severe Adverse Events: none Fatigue - 93/160(58%) vs. 86/160(54%) Arthralgia - 48/160(30%) vs. 66/160(41%) Irritability - 53/160(33%) vs. 49/160(31%) Decreased appetite - 30/160(19%) vs. 34/160(21%) Fever - 30/160(19%) vs. 75/160(47%) Pruritus - 27/160(17%) vs. 24/160(15%) Headache - 25/160(16%) vs. 28/160(18%) Cough - 20/160(13%) vs. 20/160(13%) Myalgia - 23/160(14%) vs. 30/160(19%) Dermatitis - 19/160(12%) vs. 9/160(6%) Nausea - 14/160(9%) vs. 15/160(9%)	

Author, Year Country Study Name Quality	Genotype Severity of Liver Disease Proportion Treatment- Naïve	Duration of Followup	Outcome	Subgroup Analyses	Subgroup Analyses	Histologic Response	Adverse Events	Funding Source
Ascione, 2010 ¹ Liver Unit of Cardarelli Hospital - Napoli, Italy Continued							Dyspnea - 13/160(8%) vs. 19/160(12%) Thyroid - 12/160(8%) vs. 9/160(6%) Insomnia - 11/160(7%) vs. 17/160(11%) Alopecia - 9/160(6%) vs. 22/160(14%) Depression - 11/160(7%) vs. 9/160(6%) Dose modification due to: Anemia - 30/160(19%) vs. 30/160(19%) Neutropenia - 4/160(3%) vs. 4/160(3%) Thrombocytopenia - 7/160(4%) vs. 6/160(4%)	

Author, Year Country			Protease			Number Screened/	Age
Study Name	Interferon	Ribavirin	Inhibitor			Eligible/ Enrolled/	Sex
Quality	Regimen	Regimen	Regimen	Eligibility	Exclusion	Analyzed	Race
Bruno, 2004 ²	A. Pegylated	A. 1000-1200mg	None	Treatment-naïve	Neutrophils <1500/ mL3	NR/NR/22/22	A vs. B
Italy	interferon alpha-	mg/day		HCV-RNA ≥ 2000 /	Platelet count < 90K mL3		Age mean: 47 vs. 40
	2a 180 mcg/week	depending of		mL	Hemoglobin <12 g/dL in		Female: 30% vs.
Viral dynamics and	for 12 weeks	body weight for		ALT > upper limit of	women and <13 g/dL in men		25%
pharmacokinetics of	B. Pegylated	12 weeks (<u><</u> 75 kg		normal within 6	Creatinine level >1.5 times		Non White: 10% vs.
Pegylated interferon	interferon alpha-	/ >75 kg)		months of study	upper limit of normal		0%
alpha-2a and Pegylated	2b 1.0 mcg/week	B. 1000-1200mg		Liver biopsy consistent	Co infection with HIV		
interferon alpha-2b in	for 12 weeks	mg/day		with chronic hepatitis	Decompensated liver disease		
naïve patients with		depending of			Poorly controlled psychiatric		
chronic hepatitis C; a		body weight for			disease		
randomized, controlled		12 weeks (<75 kg			Alcohol or drug abuse within		
study		/ >75 kg)			year		
					Substantial coexisting medical		
Overall Quality: Poor					conditions		

Author, Year Country Study Name Quality	Genotype Severity of Liver Disease Proportion Treatment- Naïve	Duration of Followup	Outcome	Subgroup Analyses	Subgroup Analyses	Histologic Response	Adverse Events	Funding Source
Bruno, 2004 ²	A vs. B	12 weeks	NA	NA	NA	NA	NR	Hoffman-
Italy	Genotype 1: 70% vs.							LaRoche
	50%							
Continued	Cirrhosis/transition to							
	cirrhosis: 20% vs. 16%							
	HCV-RNA mean (log):							
	5.8 vs. 5.6							
	Treatment-naïve: 100%							

Author, Year Country Study Name Quality	Interferon Regimen	Ribavirin Regimen	Protease Inhibitor Regimen	Eligibility	Exclusion	Number Screened/ Eligible/ Enrolled/ Analyzed	Age Sex Race
DiBisceglie, 2007 ³ United States Early virologic response after Pegylated interferon alpha-2a plus ribavirin or Pegylated interferon alpha-2b plus ribavirin treatment in patients with chronic hepatitis C Overall Quality: Fair	A. Pegylated interferon alpha- 2a 180 mcg weekly for 12 weeks B. Pegylated interferon alpha- 2b 1.5 mcg/kg weekly for 12 weeks	A. 1000-1200mg mg/day depending of body weight for 12 weeks (<75 kg / >75 kg) B. 1000-1200mg mg/day depending of body weight for 12 weeks (<75 kg / >75 kg)	None	Treatment-naïve patients Chronic HCV genotype 1 infection Age 18 years or older HCV RNA >800K IU/mL	HBV HIV co infection History of other chronic liver disease Decompensated liver disease or Child-Pugh score >6 Alcohol or drug abuse within year Pregnant or breastfeeding women and male partners Neutrophils <1500/mL3 Platelet count <90K /mL3 Hemoglobin <12 g/dL in women and <13 g/dL in men Creatinine >1.5 times upper limit of normal History of server psychiatric, immunologically mediated, cardiac, or chronic pulmonary disease	NR/NR/385/380	A vs. B Age mean: 47 vs. 48 Female: 36% vs. 29% Non White: 31% vs. 28%

Author, Year Country Study Name Quality	Genotype Severity of Liver Disease Proportion Treatment- Naïve	Duration of Followup	Outcome	Subgroup Analyses	Subgroup Analyses	Histologic Response	Adverse Events	Funding Source
DiBisceglie, 2007 ³ United States Continued	A vs. B Genotype 1: all Cirrhotic: 14.8% vs. 15.2% HCV RNA mean (log): 6.5 vs. 6.5 Treatment-naïve: 100%	12 weeks	NA	NA	NA	NA	A vs. B Overall withdrawals: 18/189 (10%) vs. 27/191 (14%); p=NS Withdrawals for adverse events: 2/189 (1%) vs. 11/191 (6%); p=NS Serious adverse events: NR Deaths: NR Fatigue: 132/187 (71%) vs. 137/190 (72%); p=NS Headache: 105/187 (56%) vs. 112/190 (59%); p=NS Nausea: 77/187 (41%) vs. 85/190 (45%); p=NS Chills: 46/187 (25%) vs. 79/190 (30%); p=NS Chills: 46/187 (20%) vs. 57/190 (30%); p=NS Fever: 38/187 (20%) vs. 62/190 (33%); p=NS Depression: 46/187 (25%) vs. 46/190 (24%); p=NS Arthralgia: 45/187 (24%) vs. 44/190 (23%); p=NS Dizziness: 39/187 (21%) vs. 48/190 (25%); p=NS Influenza-like illness: 34/187 (18%) vs. 44/190 (23%); p=NS Diarrhea: 33/18 (18%) vs. 39/190 (21%); p=NS Decreased appetite: 28/187 (15%) vs. 40/190 (21%); p=NS	Roche

Author, Year Country Study Name Quality	Genotype Severity of Liver Disease Proportion Treatment- Naïve	Duration of Followup	Outcome	Subgroup Analyses	Subgroup Analyses	Histologic Response	Adverse Events	Funding Source
DiBisceglie, 2007 ³							Rash: 27/187 (14%) vs.	
United States							39/190 (21%); p=NS Myalgia: 31/187 (17%) vs.	
Continued							34/190 (18%); p=NS	
							Vomiting: 26/187 (14%) vs. 38/190 (20%); p=NS	
							Injection-site erythema:	
							25/187 (13%) vs. 38/190	
							(20%); p=NS	
							Anemia: 20/187 (11%) vs. 22/190 (12%); p=NS	
							Dysgeusia: 17/187 (9%) vs.	
							21/190 (11%); p=NS	

Author, Year Country Study Name Quality	Interferon Regimen	Ribavirin Regimen	Protease Inhibitor Regimen	Eligibility	Exclusion	Number Screened/ Eligible/ Enrolled/ Analyzed	Age Sex Race
Escudero, 2008 ⁴	A: Pegylated	A: 800-1200 mg	None	Treatment naïve	HIV infection, Hepatitis B	NR/NR/183/183	A vs. B
Valencia, Spain	interferon alpha-	daily for 24 or 48		patients 18 years and	infection		Age: mean (SD):
(outpatient clinic -	2a 180 μg/week	weeks (genotype		older	Autoimmune disease		44.4(9.34) vs.
Service of Hepatology	for 24 or 48	2/3 and 1/4		Sero-positive	Autoimmune hepatitis,		43.6(9.62) years
of University Hospital	weeks (genotype	respectively)		Genotype-RNA	decompensated Liver disease		
Clinic)	2/3 and 1/4			Evidence of Genotype	hematological conditions		Male - 64/91(70%)
	respectively)	B: 800-1200 mg		1,2,3 or 4 infection	Decompensated diabetes		vs. 56/92 (61%)
Pegylated alpha-		daily for 24 or 48		Serum Genotype RNA	Thyroid disease (poorly		
interferon-2a plus	B: Genotype 2/3:	weeks (genotype		concentration > 30	controlled) History of Severe		Race: NR
ribavirin compared with	Pegylated	2/3 and 1/4		IU/mL	Psychiatric Disease, Alcohol		
pegylated alpha-	interferon alpha-	respectively)		ALT above upper limit	or Drug dependence within 1		
interferon-2b plus	2b 1.5			of normal	year prior to entry into study		
ribavirin for initial	μg/kg/week for			Diagnostic liver biopsy			
treatment of chronic	24 or 48 weeks			done within 6 months	Subjects recruited in actual		
HCV: prospective,	(genotype 2/3 and			prior to enrollment	conditions of daily practice in		
nonrandomized study	1/4 respectively)				outpatient clinic		
Overall Quality: Poor							

Author, Year Country Study Name Quality	Naïve	Duration of Followup	Outcome	Subgroup Analyses	Subgroup Analyses	Histologic Response	Adverse Events	Funding Source
Escudero, 2008 ⁴ Valencia, Spain (outpatient clinic - Service of Hepatology of University Hospital Clinic) Continued	A vs. B Genotype 1- 59/91(65%) vs. 58/92(64%) Genotype 2- 5/91(6%) vs. 4/92(4%) Genotype 3- 13/91(14%) vs. 23/92(25%) Genotype 4- 12/91(13%) vs. 6/92(7%) Genotype 5- 2/91(2%) vs. 1/92(1%) Scale by Batts & Ludwig, 1995: Grade - mean (SD): 2.1(.81) vs. 2.1(.91) Stage - mean (SD): 2.1(.98) vs. 2.0(1.07) Steatosis - 30/91(34%) vs. 43/92(46.7%) HCVRNA mean(log IU/mL): 5.9 vs. 5.8 Treatment-naïve: 100%	Followup at 24 weeks post-treatment	A vs. B ETR: NR SVR: 60/91(65.9%) vs. 57/92(62%)	NR	A vs. B Variables significantly associated with response to antiviral therapy: Genotype (odds ratio [OR] = 0.076, 95% confidence interval [CI] 0.029 – 0.198, P = 0.000) Presence of steatosis in the liver biopsy (OR = 2.799, 95% CI 1.362–5.755, p=0.005). Genotype 1: steatosis was the only variable significantly associated with response to antiviral treatment: (OR = 2.450, 95% CI 1.126–5.332, p=0.024) SVR: Genotype 1 - 30/59 (50.8%) vs. 27/58(46.6%) Genotype 2/3 - 17/18 (95%) vs. 24/27(89.3%) Genotype 4 - 11/12 (91.7%) vs. 5/6(83.3%)	NR	A vs. B Overall withdrawals - 22/91(24%) vs. 28/92(30%) Deaths - NR Dermatological symptoms: 5/183(3%) Severe neutropenia (<0.5 x 109 cells/L): 3/183(2%) Depression-related events: 2/183(1%) Anemia (hemoglobin, <10.0 g/dL): 2/183(1%) Thrombocytopenia (<50 x 109 cells/L): 2/183(2%) Hypothyroidism: 2/183(1%) Tachyarrhythmia: 1/183(0.5%) Poor tolerability with various adverse events: 5/183(3%) Dose modifications because of neutropenia: 8/91(8%) vs. 7/92(8%)	Internal funding

Author, Year Country Study Name Quality	Interferon Regimen	Ribavirin Regimen	Protease Inhibitor Regimen	Eligibility	Exclusion	Number Screened/ Eligible/ Enrolled/ Analyzed	Age Sex Race
Kamal, 2011 ⁵ Egypt Enhanced efficacy of pegylated interferon alpha-2a over pegylated interferon and ribavirin in chronic hepatitis C genotype 4A randomized trial and quality of life analysis Overall Quality: Fair	A. Pegylated interferon alfa-2a 180 mcg/week for 48 weeks B. Pegylated interferon alfa-2b 1.5 mcg/kg/week for 48 weeks	A. Ribavirin 1000-1200 mg daily (<75 kg / >75 kg) for 48 weeks B. Ribavirin 1000-1200 mg daily(<75 kg / >75 kg) for 48 weeks	None	Treatment naïve Age 18-60 years HCV genotype 4 ALT at least twice the upper limit of normal during the 6 months prior Detectable anti-HCV antibodies Detectable HCV RNA Histologic evidence of chronic hepatitis C in liver biopsy within preceding year	Evidence of other liver disease Co-infection with HIV, hepatitis A, B, or schistosomiasis Leucocytes <3000/mm3 Neutrophils <1500/mm3 Hemoglobin <12 g/dl for women or <13 g/dl for men Thrombocytopenia <90K/mm3 Creatinine >1.5x upper limit of normal Organ transplantation Cancer Severe cardiac or pulmonary disease Unstable thyroid dysfunction Severe depression or psychiatric disorder Active substance abuse Pregnancy Breast feeding BMI>30Kg/m2 Known sensitivity to drugs tested Determined by investigators to be unreliable or noncompliant	226/217/217	A vs. B Age: 42 vs. 41 Female: 46% vs. 56% Race: NR (Egyptian centers)

Author, Year Country Study Name Quality	Genotype Severity of Liver Disease Proportion Treatment- Naïve	Duration of Followup	Outcome	Subgroup Analyses	Subgroup Analyses	Histologic Response	Adverse Events	Funding Source
Kamal, 2011 ⁵	A vs. B	24 weeks after	A vs. B	NR	NR	NR	A vs. B	Ain Shams
Egypt	Genotype 1: 0% Genotype 4: 100%	treatment completion	SVR: 77/109 (70.6%) vs. 59/108 (54.6%);				Overall withdrawals: 2/109 (2%) vs. 1/108 (1%); p=NS	University
Continued	Grade 3 Steatosis: 38%		p=0.0172				Withdrawals for adverse events: 1/109 (1%) vs. 1/108	
	vs. 37%		SF-6D (During Treatment): 0.735 vs.				(1%); p=NS	
	Treatment-naïve: NR		0.730; p=0.8067 SF-6D (after treatment): 0.769 vs. 0.737; =0.04				Mild adverse events: 54/109 (50%) vs. 40/108 (37%); p=NS Moderate adverse events; 18/109 (17%) vs. 12/108	
			Chronic Liver Disease Health Survey Questionnaire (CLDQ) (during treatment): 5.3 vs. 5.0; p=0.16 CLDQ (after				(11%); p=NS Severe adverse events; 4/109 (4%) vs. 3/108 (3%); p=NS	
			treatment): 5.9 vs. 5.5; p=0.02					

Author, Year Country Study Name Quality	Interferon Regimen	Ribavirin Regimen	Protease Inhibitor Regimen	Eligibility	Exclusion	Number Screened/ Eligible/ Enrolled/ Analyzed	Age Sex Race
Khan, 2007 ⁶	A: Pegylated	A: 800 mg/day	None	NR	NR	NR/NR/NR/66	NR
Pakistan	interferon alfa-2a 180 mcg/week for	for 24 weeks B: 800 mg/day					
Pegylated interferon	24 weeks	for 24 weeks					
alfa-2a ribavirin vs.	B: Pegylated						
Pegylated interferon alfa-2b/ribavirin	interferon alfa-2b 1.0 mcg/week for						
combination therapy in	24 weeks						
chronic hepatitis C	21 WOORS						
genotype 3							
Overall Quality: Not Assessed							

Author, Year Country Study Name Quality	Genotype Severity of Liver Disease Proportion Treatment- Naïve	Duration of Followup	Outcome	Subgroup Analyses	Subgroup Analyses	Histologic Response	Adverse Events	Funding Source
Khan, 2007 ⁶	J 1	24 weeks after	A vs. B	NR	NR	NR	A vs. B	NR
Pakistan	Genotype 4: 100%	end of treatment	SVR: 26/33 (79%) vs. 27/33 (82%), p=NS				Overall withdrawals: 1/33 (3%) vs. 1/33 (3%)	
Continued			((6,1) 121 5,00 (6,1)	

Author, Year Country			Protease			Number Screened/	Age
Study Name	Interferon	Ribavirin	Inhibitor			Eligible/ Enrolled/	Sex
Quality	Regimen	Regimen	Regimen	Eligibility	Exclusion	Analyzed	Race
Mach 2011 ⁷	A:	A:	None	Patients with	Patients with decompensated	NR/NR/260/260	A vs. B
Poland	Pegylated	Ribavirin 1.0–1.2		anti-HCV and	liver cirrhosis, autoimmune		Age: 44 vs. 45.2
	interferon alfa-2a	g oral daily		HCV-RNA in serum	liver disease, alcohol abuse,		years
Efficacy of pegylated	- 180 μg			and elevated alanine	liver cancer, hepatitis B virus		Female: 37.7% vs.
interferon alfa-2a	subcutaneously	B:		aminotransferase	or HIV coinfection, any		42%
or alfa-2b in	once	Ribavirin 1–1.2 g		(ALT) levels at least 6	severe chronic disease,		Race: NR (Polish
combination with	a week	oral daily		months before the	diabetes, dyslipidemia,		centers)
ribavirin in				inclusion, chronic	metabolic syndrome,		
the treatment of chronic	B:			hepatitis confirmed by	hemochromatosis,		
hepatitis caused by	Pegylated			histological	and immunosuppressive		
hepatitis C virus	interferon alfa-2b			examination, body	therapy.		
genotype 1b	- 1.5 mg/kg of			mass index (BMI)			
	body weight			below 30 kg/m ² .			
Overall Quality: Fair	once a week						

Author, Year Country Study Name Quality	Genotype Severity of Liver Disease Proportion Treatment- Naïve	Duration of Followup	Outcome	Subgroup Analyses	Subgroup Analyses	Histologic Response	Adverse Events	Funding Source
Mach 2011 ⁷	A vs. B	24 weeks after	A vs. B:	NR	NR	NR	NR	Polish
Poland	Genotype 1b: 100%	end of treatment	ETR:71.7% vs. 60.7%,					National
			p=NR					Health Fund
Continued	Liver fibrosis:							
	F0-2 – 78.1% vs. 72.9%		SVR: 49.3% vs.					
	F3-4 – 21.95% vs. 27.1%		44.3%, p=NS					
	Treatment-naïve: NR							

Author, Year Country Study Name Quality	Interferon Regimen	Ribavirin Regimen	Protease Inhibitor Regimen	Eligibility	Exclusion	Number Screened/ Eligible/ Enrolled/ Analyzed	Age Sex Race
Magni, 2009 ⁸ Italy	A: Pegylated interferon alfa-2a 180 mcg/week for	A: 10.5 mg/kg for 24-48 weeks based on	None	NR	NR	NR/NR/NR/218	NR
Antiviral activity and tolerability between pegylated interferon alfa-2a and alfa-2b in naïve patients with chronic hepatitis C: results of a prospective monocentric randomized trial	24-48 weeks based on genotype B: Pegylated interferon alfa-2b 1.0 mcg/week for 24-48 weeks based on genotype	genotype B: 10.5 mg/kg for 24-48 weeks based on genotype					
Overall Quality: Not Assessed							

Author, Year Country Study Name Quality	Genotype Severity of Liver Disease Proportion Treatment- Naïve	Duration of Followup	Outcome	Subgroup Analyses	Subgroup Analyses	Histologic Response	Adverse Events	Funding Source
Magni, 2009 ⁸	A vs. B	24 weeks after	A vs. B	NR	A vs. B	NR	A vs. B	NR
Italy	Genotype 1/4: 61% vs.	end of treatment	SVR: 68/100 (68%)		Genotype 1/4: 36/58		Withdrawals due to adverse	
	51%		vs. 79/118 (67%);		(62%) vs. 34/55 (62%);		events: 5% vs. 6.8%; p=NS	
Continued	Genotype 2/3: 39% vs.		p=NS		p=NS			
	49%				Genotype 2/3: 32/37			
					(87%) vs. 45/52 (87%);			
	Treatment-naïve: NR				p=NS			

Author, Year							
Country			Protease			Number Screened/	Age
Study Name	Interferon	Ribavirin	Inhibitor			Eligible/ Enrolled/	Sex
Quality	Regimen	Regimen	Regimen	Eligibility	Exclusion	Analyzed	Race
McHutchison, 2008 ⁹	A. Pegylated	A. Weight-based	None	Treatment-naïve	HIV	4469/3431/3083/3070	A vs. B vs. C
US	interferon alfa-2b	800-1400 mg		Ages 18 years or older	HBV		Age mean: 48 vs. 48
	1.0 mcg/kg/week	daily for 48		Chronic HCV genotype	Other liver disease		vs. 48
Individualized Dosing	for 48 weeks.	weeks		1 infection	Poorly controlled diabetes		Female: 40% vs.
Efficacy vs. Flat Dosing	B. Pegylated	B. Weight-based		Detectable HCV RNA	Weight >125 kg		40% vs. 41%
to Assess Optimal	interferon alfa-2b	800-1400 mg		level	Severe depression		Non White: 29% vs.
Pegylated Interferon	1.5 cg/kg/week	daily for 48		Neutrophil count >	Severe psychiatric disorder		28% vs. 29%
Therapy (IDEAL)	for 48 weeks.	weeks		1500 /mm3	Active substance abuse		
	C. Pegylated	C. 1000 mg (<75		Platelets \geq 80,000			
Overall Quality: Fair	interferon alfa-2a	kg) - 1200 mg		/mm3			
	180 mcg/week for	$(\geq 75 \text{ kg})$ daily for		Hemoglobin $\geq 12 \text{ g/dL}$			
	48 weeks.	48 weeks		for women or 13 g/dL			
				for men			
	Discontinued if	Weight-based					
	HCV RNA	dosing					
	detectable and not	\leq 65 kg: 800 mg					
	decreased by 2	daily					
	log IU from	66 - 85kg: 1000					
	baseline at 12	mg daily					
	weeks or HCV	86-105kg: 1200					
	RNA detectable	mg daily					
	at 24 weeks	106 -125kg: 1400					
		mg daily					
		T. 110					
		Discontinued if					
		HCV RNA					
		detectable and					
		not decreased by					
		2 log IU from baseline at 12					
		weeks or HCV					
		RNA detectable					
	ĺ	at 24 weeks	l	ĺ			

Author, Year Country Study Name Ouality	Genotype Severity of Liver Disease Proportion Treatment- Naïve	Duration of Followup	Outcome	Subgroup Analyses	Subgroup Analyses	Histologic Response	Adverse Events	Funding Source
McHutchison, 2008 ⁹	A vs. B vs. C	24 weeks after	A vs. B vs. C	A vs. B vs. C	A vs. B vs. C	NR	A vs. B vs. C	Schering-
US Continued	Genotype 1: 100% Metavir fibrosis score 3 or 4: 11% vs. 11% vs. 11% HCV-RNA≥600K: 82% vs. 82% vs. 82% Treatment-naïve: 100%	24 weeks after treatment completion	ETR: 500/1016 (49%) vs. 542/1016 (53%) vs. 667/1035 (64%); (p=0.04 for A vs. B, p<0.001 for B vs. C) SVR: 386/1016 (38%) vs. 406/1019 (40%) vs. 423/1035 (41%); (p=0.20 for A vs. B, p=0.57 for B vs. C)	A vs. B vs. C (p-values from interaction models given) Black: 31/187 (17%) vs. 42/183 (23%) vs. 52/200 (26%); White: 316/362 (36%) vs. 319/732 (44%) vs. 324/733 (44%); (p=0.18 for A vs. B, p=0.62 for B vs. C) Female: 147/409 (36%) vs. 180/406 (44%) vs. 177/422 (42%) Male: 239/607 (39%) vs. 226/613 (37%) vs. 246/613 (40%); (p=0.01 for A vs. B, p=0.20 for B vs. C)	Metavir fibrosis score F3 or F4: 32/107 (30%) vs. 23/111 (21%) vs. 26/110 (24%) Metavir fibrosis score F0-F2: 335/864 (39%) vs. 366/869 (42%) vs. 376/862 (44%); (p=0.06 for A vs. B, p=0.75 for B vs. C) Baseline HCV RNA >600K IU/mL: 277/830 (33%) vs. 295/836 (35%) vs. 303/852 (36%) Baseline HCV RNA <600K IU/mL: 109/186 (59%) vs. 111/183 (61%) vs. 120/183 (66%); (p=0.99 for A vs. B, p=0.41 for B vs. C) Weight <75 kg: 211/555 (38%) vs. 264/605 (44%) Weight >75 kg: 175/461 (38%) vs. 187/455 (41%) vs. 159/430 (37%); (weight in kg as continuous variable p=0.94 for A vs. B; p=0.39 for B vs. C)	INK	Overall withdrawals: 523/1016 (52%) vs. 479/1019 (47%) vs. 414/1035 (40%); (p=0.04 for A vs. B, p=0.001 for B vs. C, p<0.001 for A vs. C) Withdrawals for adverse events: 98/1016 (10%) vs. 129/1019 (13%) vs. 135/1035 (13%); (p=0.03 for A vs. B, p=0.80 for B vs. C, p<0.001 for A vs. C) Deaths: 1/1016 (<1%) vs. 5/1019 (<1%) vs. 6/1035 (<1%); (p=NS) Serious adverse event: 94/1016 (9%) vs. 88/1019 (9%) vs. 121/1035 (12%); (p=0.63 for A vs. B, p=0.02 for B vs. C, p=0.07 for A vs. C) Fatigue: 676/1016 (67%) vs. 672/1016 (66%) vs. 656/1035 (63%); (p=NS) Headache: 486/1016 (48%) vs. 508/1019 (50%) vs. 438/1035 (42%); (p=0.36 for A vs. B, p=0.001 for B vs. C, p=0.01 for A vs. C) Nausea: 377/1016 (37%) vs. 433/1019 (43%) vs. 377/1035 (36%); (p=0.01 for A vs. B, p=0.005 for B vs. C, p=0.75 for A vs. C)	Plough

Author, Year Country Study Name Quality	Genotype Severity of Liver Disease Proportion Treatment- Naïve	Duration of Followup	Outcome	Subgroup Analyses	Subgroup Analyses	Histologic Response	Adverse Events	Funding Source
McHutchison, 2008 ⁹ US Continued				Age ≤40: 72/154 (47%) vs. 74/140 (53%) vs. 91/163 (56%) Age >40: 314/862 (36%) vs. 332/879 (38%) vs. 332/872 (38%); (p=0.46 for A vs. B, p=0.67 for B vs. C)			Pyrexia: 311/1016 (33%) vs. 356/1019 (35%) vs. 237/1035 (23%); (p=0.26 for A vs. B, p<0.001 for B vs. C, p<0.001 for A vs. C) Myalgia: 270/1016 (27%) vs. 274/1019 (27%) vs. 233/1035 (23%); (p=0.87 for A vs. B, p=0.02 for B vs. C, p=0.03 for A vs. C) Depression: 197/1016 (19%) vs. 260/1019 (26%) vs. 217/1035 (21%); (p=0.001 for A vs. B, p=0.02 for B vs. C, p=0.37 for A vs. C) Neutropenia: 188/1016 (19%) vs. 263/1019 (26%) vs. 326/1035 (32%); (p<0.001 for A vs. B, p=0.02 for B vs. C, p=0.37 for A vs. C) Neutropenia: 188/1016 (19%) vs. 263/1019 (26%) vs. 326/1035 (32%); (p<0.001 for A vs. B, p=0.004 for B vs. C, p<0.001 for A vs. C) Anemia: 293/1016 (29%) vs. 345/1016 (34%) vs. 348/1035 (34%); (p=0.02 for A vs. B, p=0.91 for B vs. C, p=0.02 for A vs. C) Neutrophils <750/mm3: 147/1008 (15%) vs. 222/1000 (22%) vs. 279/1034 (27%); (p<0.001 for A vs. B, p=0.01 for B vs. C, p<0.001 for A vs. C) Hemoglobin <10 g/dl: 255/1008 (25%) vs. 307/1000 (31%) vs. 306/1034 (30%); (p=0.007 for A vs. B, p=0.59 for B vs.	

Author, Year Country Study Name Quality	Interferon Regimen	Ribavirin Regimen	Protease Inhibitor Regimen	Eligibility	Exclusion	Number Screened/ Eligible/ Enrolled/ Analyzed	Age Sex Race
Miyase, 2012 ¹⁰ Japan Randomized trial of peginterferon alpha-2a plus ribavirin versus peginterferon alpha-2b plus ribavirin for chronic hepatitis C in Japanese patients Overall Quality: Good	A: Pegylated interferon alpha-2a at a dosage of 180 mcg once weekly. B: Pegylated interferon alpha-2b at a dosage of 60-150 mcg/kg (weight-based) once weekly. 35–45 kg - 60 mcg 46–60 kg, - 80 mcg 61–75 kg - 100 mcg 76–90 kg, - 120 mcg 91–120 kg - 150 mcg	A: RBV(weight-based) 600 mg/day ≤60 kg - 800 mg/day 60–80 kg - 1000 mg/day B: RBV(weight-based) 600 mg/day ≤60 kg - 800 mg/day 60–80 kg - 1000 mg/day	None	Consecutive PEG IFN- naïve adults (C18 years of age) who were infected with HCV genotype 1 were eligible for enrollment. The inclusion criteria were a serum HCV RNA level [5.0 log IU/mL, a liver biopsy performed within 6 months of starting treatment, and use of contraceptive methods during therapy and for 6 months after the end of treatment.	hemoglobin level \10 g/dL; white blood cell count\1.8x 10 ³ /mm3 or platelet count<\7.0 x 10 ⁴ /mm3; abnormal serum creatinine level; hepatitis B surface antigen positivity; human immune deficiency virus positivity; other cause of liver disease; history of liver decompensation; clinically relevant depression or any other psychiatric disease; cancer; severe cardiac, pulmonary, or renal disease; uncontrolled diabetes; or severe hypertension with vascular complications, including retinopathy.	N/NR/206/201	A vs. B Age mean: 59.2 vs. 58.9 years Female: 61.4% vs. 60% Non White: NR

	Genotype		-	1				1
Author, Year	Severity of Liver							
,	•							
Country	Disease	D4		C1		TT:-4-1		F 12
Study Name	1	Duration of		Subgroup		Histologic		Funding
Quality	Naïve	Followup	Outcome	Analyses	Subgroup Analyses	Response	Adverse Events	Source
Miyase, 2012 ¹⁰	A vs. B	24 weeks after	A vs. B	A vs B	A vs. B		Overall withdrawals -	NR
Japan	Genotype 1: 100%	end of treatment	ETR: NR	Age:	Non cirrhosis - 55/81		17(16.8%) vs. 26(26.0%)	
				<60 years-	(67.9%) vs. 46/83		0.124	
Continued	HCV RNA (log IU/mL) -		SVR: 66/101(65.3%)	33/52 (63.5%)	(55.4%), p=0.100			
	6.3 ± 0.6 vs. 6.2 ± 0.7 , p=		vs. 51/100(51%),	vs. 31/49	Cirrhosis - 11/20 (55.0%)		Neutropenia - 43(42.6%) vs.	
	0.151		p=0.039	(63.3%),	vs. 5/17 (29.4%),		29(29.0%), p=0.056	
				p=0.984	p=0.117		Anemia - 62(61.4%) vs.	
	Cirrhosis: 20% vs. 17%			≥60 years -			63(63.0%), p=0.885	
				33/49 (67.3%)	HCV RNA:		Thrombocytopenia -	
	Treatment-naïve: 100%			vs. 20/51	≤6 log IU/mL - 22/28		30(29.7%) vs. 27(27.0%),	
				(39.2%),p=0.00	(78.6%) vs. 28/39		p=0.755	
				5	(71.8%), p=0.530		Dose modification -	
					>6 log IU/mL - 44/73		13(12.9%) vs. 19(19.0%),	
				Female: 38/62	(60.3%) vs. 23/61		p=0.253	
				(61.3) vs. 26/60	(37.7%), p=0.009			
				(43.3), p=0.047			Fever - 41(40.6%) vs.	
							76(76.0%), p<0.001	
				Weight (kg)			Dermatitis, itching -	
				≥60 kg - 39/61			71(70.3%) vs. 56(56.0%),	
				(63.9%) vs.			p=0.041	
				28/64 (43.8%),			Fatigue - 47(46.5%) vs.	
				p=0.024			42(42.0%), p=0.571	
				≤60 kg - 27/40			Decreased appetite -	
				(67.5%) 23/36			43(42.6%) vs. 56(56.0%),	
				vs. (63.9%), p=			p=0.067	
				0.740			Insomnia - 34(33.7%) vs.	
							39(39.0%), p=0.465	
							Headache - 28(27.7%) vs.	
							24(24.0%), p=0.630	
							Stomatitis - 15(14.9%) vs.	
							22(22.0%), p=0.207	
							Nausea - 13(12.9%) vs.	
							19(19.0%), p=0.253	
							Arthralgia -15(14.9%) vs.	
							9(9.0%), p=0.277	
						1	Irritability - 12(11.9%) vs.	
						1	8(8.0%), p=0.481	
							Depression - 9(8.9%) vs.	
							8(8.0%), p=1.000	
							Cough - 6(5.9%) vs. 3(3.0%),	
ĺ							p=0.498	

Author, Year Country Study Name Quality	Interferon Regimen	Ribavirin Regimen	Protease Inhibitor Regimen	Eligibility	Exclusion	Number Screened/ Eligible/ Enrolled/ Analyzed	Age Sex Race
Rumi, 2010 ¹¹ University of Milan, Italy	Genotype 1/4: A. Pegylated interferon alfa-2a 180 mcg/week for	Genotype 1/4: A. 1000-1200 mg/day for 48 weeks	None	Treatment naïve patients 18-70 years old with serum HCV-RNA	Persistently normal ALT Hemoglobin ≤ 12g/dL in women and ≤13g/dL in men White Blood Cell count <=	473/447/447/431	A vs. B Age: Mean (SD): 51.6(12.0) vs. 52.8(12.0) years
Clinical Advances in Liver, Pancreas, and Biliary Tract (MIST Study) - Randomized Study of Pegylated	48 weeks B. Pegylated interferon alfa-2b 1.5 mcg/kg/week for 48 weeks	B. 800-1200 mg/day for 48 weeks Genotype 2/3:		Higher than normal ALT activity, and Diagnostic Liver Biopsy done within 24 months prior to	2.5x103 /mm3 Neutrophil <= 1.5x103 /mm3 Platelet count<= 75x103 /mm3 Serum creatinine level >1.5x		Male - 128/212 (60.4%) vs. 120/219 (54.8%)
interferon-alpha-2a Plus Ribavirin vs. Pegylated interferon-alpha-2b plus Ribavirin in Chronic Hepatitis C	Genotype 2/3: A. Pegylated interferon alfa-2a 180 mcg/week for 24 weeks	A. 800 mg/day for 24 weeks B. 800-1200 mg/day for 24 weeks		enrollment	upper limit of normal Liver disease (any other) HIV co infection Autoimmune diseases Contraindications to Interferon and Ribavirin		Race: NR
Overall Quality: Fair	B. Pegylated interferon alfa-2b 1.5 mcg/kg/week for 24 weeks						

Author, Year Country Study Name Quality	Genotype Severity of Liver Disease Proportion Treatment- Naïve	Duration of Followup	Outcome	Subgroup Analyses	Subgroup Analyses	Histologic Response	Adverse Events	Funding Source
Rumi, 2010 ¹¹ University of Milan, Italy Continued	A vs. B Genotype 1- 91/212 (42.9%) vs. 87/219 (39.7%) Genotype 2- 69/212 (32.5%) vs. 74/219 (33.8%) Genotype 3- 34/212 (16.0%) vs. 32/219 (14.6%) Genotype 4- 18/212 (08.5%) vs. 26/219 (11.9%) Ishak score of S5, 6: Overall: 81/212(38%) vs. 39/219(18%) HCV-RNA >600K IU/L: 53% vs. 55% Treatment-naïve: 100%	Followup at 24 weeks post-treatment	A vs. B ETR: 166/212 (78%) vs. 146/219 (67%), p=0.009 SVR: 140/212 (66%) vs. 119/219 (54%), p=0.02	NR	A vs. B ETR: Genotype 1: 59/91 (65%) vs. 38/87 (44%), p=0.007 Genotype 2: 66/69 (96%) vs. 69/74 (93%), p=0.09 Genotype 3: 32/34 (94%) vs. 29/32 (91%), p=0.09 Genotype 4: NR ("sound comparison of treatment efficacy compromised by small sample size") SVR: Genotype 1: 44/91 (48%) vs. 28/87 (32%), p=0.04 Genotype 2: 66/69 (96%) vs. 61/74 (82%), p=0.01 Genotype 3: 22/34 (65%) vs. 22/32 (69%), p=0.09 Genotype 4: NR ("sound comparison of treatment efficacy compromised by small sample size")	NR	A vs. B Discontinuation due to adverse events: 16/212(8%) vs. 17/219(8%) Overall Withdrawals (including loss to followup and "other"): 46/212(22%) vs. 73/219(33%) Deaths: NR Serious Adverse Events: 2/212 (1%) vs. 2/219(1%) Adverse Events: Grade 2 anemia: 35/212(16%) vs. 50/219(23%) Grade 3 anemia: 2/212(1%) vs. 2/219(1%) Grade 3 neutropenia: 46/212(22%) vs. 34/219(16%) Grades 2 or 3/thrombocytopenia: 5/212 (2%) vs. 3/219(1%) Treated with GCSF: 21/212(10%) vs. 15/219(7%) Treated with erythropoietin: 30/212(14%) vs. 27/219(12%) Depression: 19/212(9%) vs. 15/219 (7%)	Schering- Plough (now Merck), Roche, Novartis, Vertex

Author, Year Country	Genotype Severity of Liver Disease							
Study Name	Proportion Treatment-	Duration of		Subgroup		Histologic		Funding
Quality	Naïve	Followup	Outcome	Analyses	Subgroup Analyses	Response	Adverse Events	Source
Rumi, 2008 ¹¹							Influenza-like syndrome:	
University of Milan, Italy							134/212(63%) vs.	
							136/219(62%)	
Continued							Gastrointestinal symptoms:	
							8/212(4%) vs. 12/219(5%)	
							Psychiatric symptoms:	
							79/212(37%) vs.	
							70/219(32%)	
							Coughing and dyspnea:	
							22/212(10%) vs.	
							25/219(11%)	
							Dermatologic symptoms:	
							99/212(47%) vs.	
							91/219(42%)	

Author, Year Country Study Name	Interferon	Ribavirin	Protease Inhibitor	El 1 II.		Number Screened/ Eligible/ Enrolled/	Age Sex
Quality	Regimen	Regimen	Regimen	Eligibility	Exclusion	Analyzed	Race
Silva, 2006 ¹²	A. Pegylated	A. 13 mcg/kg in	None	Treatment-naïve	Liver disease of other cause	NR/NR/32/32	A vs. B
Argentina, Mexico,	interferon alfa-2a	divided dose		patients	HIV		Age mean: 46 vs. 48
Germany	180 mcg/week for	(bid) after 4th		Genotype 1a or 1b	Hemoglobinopathy		Female: 50% vs.
	8 weeks	week		Ages 18-65 years	Hemophilia		44%
A randomized trial to	B. Pegylated	B. 13 mcg/kg in		$HCV-RNA > 6x10^5$	Severe psychiatric disease		Non White: 11% vs.
compare the	interferon alfa-2b	divided dose		IU/mL	Poorly controlled diabetes		22%
pharmacokinetics,	1.5 mcg/kg/week	(bid) after 4th		$ALT/AST \le 10x$ the	mellitus		
pharmacodynamic, and	for 8 weeks	week		upper limit of normal	Significant ischemic heart		
antiviral effects of				Normal hemoglobin	disease		
pegylated interferon	After study			White-blood cells >	Chronic obstructive		
alfa-2b and Pegylated	patients were			cells/mcg L,	pulmonary disease		
interferon alfa-2b in	offered full			Neutrophils >1500	Active immune disease		
patients with chronic	course of weight-			/mcg L			
hepatitis C	based pegylated			Platelets >100K/mcg L			
· * · · · · ·	interferon alfa-2b						
Overall Quality: Poor	and ribavirin						

Author, Year Country Study Name Quality	Genotype Severity of Liver Disease Proportion Treatment- Naïve	Duration of Followup	Outcome	Subgroup Analyses	Subgroup Analyses	Histologic Response	Adverse Events	Funding Source
Silva, 2006 ¹² Argentina, Mexico, Germany Continued	A vs. B Genotype 1: all Fibrosis stage: NR HCV-RNA mean (x10 ⁶ IU/mL): 1.8 vs. 1.8 Treatment-naïve: 100%	8 weeks	NA	NA	NA	NA	A vs. B Overall withdrawals: NR Withdrawals for adverse events: 2/18 (11.1%) vs. 4/18 (22.2%); p=NS Serious adverse events: NR Deaths: NR Fatigue: 4/18 (22%) vs. 6/18 (33%); p=NS Fever: 1/18 (6%) vs. 10/18 (56%); p=0.001 Headache: 16/18 (89%) vs. 16/18 (89%); p=NS Influenza-like symptoms: 3/18 (17%) vs. 5/18 (28%); p=NS Anemia: 9/18 (50%) vs. 10/18 (56%); p=NS Hematocrit decrease: 9/18 (50%) vs. 5/18 (28%); p=NS Hemoglobin decrease: 12/18 (67%) vs. 6/18 (33%); p=0.05 Leukopenia: 14/18 (78%) vs. 9/18 (50%); p=NS Neutropenia: 12/18 (67%) vs. 10/18 (56%); p=NS Neutropenia: 12/18 (67%) vs. 10/18 (56%); p=NS Neutropenia: 12/18 (67%) vs. 11/18 (61%); p=NS Platelet count decrease: 5/18 (28%) vs. 5/18 (28%); p=NS Thrombocytopenia: 5/18 (28%) vs. 3/18 (17%); p=NS	Schering Plough

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	Author, Year							
	Country			Protease				Age
	Study Name	Interferon	Ribavirin	Inhibitor			Number Screened/ Eligible/	Sex
	Quality	Regimen	Regimen	Regimen	Eligibility	Exclusion	Enrolled/ Analyzed	Race

Author, Year Country Study Name Quality	Interferon Regimen	Ribavirin Regimen	Protease Inhibitor Regimen	Eligibility	Exclusion	Number Screened/ Eligible/ Enrolled/ Analyzed	Age Sex Race
Yenice, 2006 ¹³	A: Pegylated	A: 800-1200 mg	None	Anti HCV+, normal	Abdominal ascites	NR/80/80/74	A vs. B
Okmeydani Research &	interferon alpha-2a	daily for 48 weeks		and/or elevated serum	History of bleeding from		Age - Mean: 48.2 vs.
Training Hospital	180µg/week for 48	B: 800-1200 mg		transaminase levels	esophageal varicosities		50.8
(Istanbul, Turkey)	weeks	daily for 48 weeks		HCV+ RNA	Hepatocellular carcinoma		
				At least stage 1 fibrosis	(HCC) or other malignant		Male - 24/37(65%)
The efficacy of pegylated	B: Pegylated-			according to Knodell	disorders		vs. 27/37(73%)
interferon alpha 2a or 2b	interferon alpha-			Scoring System on liver	Use of antidepressants or		
plus ribavirin in chronic	2b1.5μg/kg for 48			biopsy	tranquilizing agents for more		Race: NR
hepatitis C patients	weeks			Hemoglobin 12 g/dl for	than 3 months		
				women and 13 g/dl for	History of depression, psychosis		
Overall Quality: Poor				men	or suicide attempt		
				Leukocyte 3x10 ³ /mm ³	Significant cardiac or		
				Neutrophils	pulmonary problems		
				$1.5 \times 10^3 / \text{mm}^3$	Hepatitis B or D		
				Platelets 100x10 ³ /mm ³	Human Immunodeficiency		
				Normal range: bilirubin,	Virus or antibodies (HIV)		
				albumin, and creatinine			
				No positive test results			
				for hepatitis B, hepatitis			
				D, or human			
				immunodeficiency virus			
				antibodies or antigens.			

Author, Year Country Study Name Quality	Genotype Severity of Liver Disease Proportion Treatment- Naïve	Duration of Followup	Outcome	Subgroup Analyses	Subgroup Analyses	Histologic Response	Adverse Events	Funding Source
Yenice, 2006 ¹³ Okmeydani Research & Training Hospital (Istanbul, Turkey) Continued	vs. 0/37(0%) 100% of subjects included had at least Stage 1 fibrosis (Knodell scale)	Followup at 24 weeks post-treatment Most patients refused a followup biopsy at the end of treatment; therefore, histological improvement was not assessed in this study due to the low number of followup biopsies.	A vs. B ETR: 28/37(75.7%) vs. 27/37(73%), p=0.79 SVR: 18/37(48.6%) vs. 13/37(35.1%), p=0.239	NR	NR	NR	A vs. B Discontinuation: 3/37(8%) vs. 3/37(8%) Overall Withdrawals: 3/37(8%) vs. 3/37(8%) Deaths: NR Serious Adverse Events: NR	Okmeydani Research and Training Hospital
	Treatment-naïve: 100%							

Evidence Table 2. Quality rating: Trials of dual therapy with pegylated interferon alpha-2a plus ribavirin compared with pegylated interferon alfa-2b plus ribavirin

			<u> </u>									
Author, Year	Randomization adequate?	Allocation concealment adequate?	Groups similar at baseline?	Eligibility criteria specified?	Outcome assessors masked?	Care provider masked?	Patient masked?	Attrition and withdrawals reported?	Loss to followup: differential/high?	Intention- to-treat analysis	Quality	Funding
Ascione, 2010 ¹	Yes	Yes	Yes	Yes	No	No	No	Yes	Yes	Yes	Fair	Carderelli Hospital, Napoli, Italy
Escudero, 2008 ⁴	No	No	Yes	Yes	No	No	No	No	Unclear	Yes	Poor	Hoffman- LaRoche
Kamal, 2011 ⁵	Yes	Yes	Yes	Yes	No - open label	No - open label	No - open label	Yes	No	Yes	Fair	NR
Mach 2011 ⁷	Unclear	Unclear	Yes	Yes	No - open label	No - open label	No - open label	No	Unclear	No	Fair	Polish National Health Fund
McHutchison 2008 ⁹	Yes	Yes	Unclear	Unclear	Unclear	Yes	Unclear	Yes	No	Yes	Fair	Vertex Pharmaceuticals
Miyase, 2012 ¹⁰	Unclear	Unclear	Yes	Yes	No - open label	No - open label	No - open label	Yes	Unclear	Yes	Fair	NR
Rumi, 2010 ¹¹	Yes	Unclear	Yes	Yes	No	No	No	Yes	Unclear	Yes	Fair	Roche
Yenice, 2006 ¹³	Unclear	Unclear	Yes	Yes	Unclear	Unclear	Unclear	Yes	No	No	Poor	Okmeydani Research and Training Hospital

Evidence Table 3. Trials of protease inhibitors plus pegylated interferon and ribavirin

						Number		Genotype
Author, Year						Screened/		Severity of Liver
Country			Protease			Eligible/	Age	Disease
Study Name	Interferon		Inhibitor			Enrolled/	Sex	Proportion
Quality	Regimen	Ribavirin Regimen	Regimen	Eligibility	Exclusion	Analyzed	Race	Treatment-Naïve
Hezode,	A. Pegylated	A. Ribavirin1000-1200	A. Telaprevir	Treatment naïve	histologic evidence of cirrhosis	388/ 334/ 334/	A vs. B vs. C	A vs. B vs. C vs. D
2009^{14}	interferon alfa-2a	mg daily for 24 weeks	750 mg tid for 12	patients ages 18-65	within 2 years of enrollment	323	vs. D	Genotype 1: all
Europe	180 mcg weekly	B. Ribavirin1000-1200	weeks	years			Age median:	Cirrhosis: 0% vs.
	for 24 weeks	mg daily for 12 weeks	B. Telaprevir	Genotype 1 with			46 vs. 44 vs.	0% vs. 1% vs. 0%
Protease Inhibition	B. Pegylated	C. Placebo	750 mg tid for 12	detectable HCV RNA			45 vs. 45	Minimal or no
for Viral	interferon alfa-2a	D. Ribavirin1000-1200	weeks				Female: 33%	Fibrosis: 43% vs.
Evaluation 2	180 mcg weekly	mg daily for 48 weeks	C. Telaprevir				vs. 40% vs.	37% vs. 40% vs.
(PROVE2)	for 12 weeks		750 mg tid for 12				45% vs. 44%	34%
	C. Pegylated	1000 mg daily for	weeks				Non White:	Treatment-naïve:
Overall Quality:	interferon alfa-2a	patients <75 kg	D. placebo				7% vs. 7% vs.	100%
Fair	180 mcg weekly	1200 mg daily for					1% vs. 7%	
	for 12 weeks	patients $\geq 75 \text{ kg}$	On day 1,					
	D. Pegylated		patients received					
	interferon alfa-2a		telaprevir 1250					
	180 mcg weekly		mg					
	for 48 weeks							

Author, Year							1
Country							
Study Name	Duration of				Histologic		Funding
Quality	Followup	Outcome	Subgroup Analyses	Subgroup Analyses	Response	Adverse Events	Source
Hezode, 2009 ¹⁴	Up to 48	A vs. B vs. C vs. D	Not reported	NR	NR	A vs. B vs. C vs. D	Vertex
Europe	weeks	ETR: 57/81 (70%)	multivariate predictors of	TVK	1110	Overall withdrawals: 20/81 (25%) vs. 10/82 (12%) vs. 8/78	Pharmaceut
Larope	following	\ /	SVR presented in			(10%) vs. 32/82 (39%); (p=0.05 for A vs. D, p<0.01 for B,	icals
Continued	treatment	48/78 (62%) vs.	supplementary table			C vs. D)	icais
Continued	completion		(variables included treatment			Withdrawals due to adverse events: 11/81 (14%) vs. 9/82	
	completion	vs. D p<0.05)	arm HCV geno-subtype,			(11%) vs. 7/78 (9%) vs. 6/82 (7%); (p=NS for A, B, C vs.	
		vs. <i>B</i> p (0.05)	baseline HCV RNA, age):			D)	
		SVR: 56/81 (69%)	Baseline HCV RNA <800K			Serious adverse event: 13/81 (16%) vs. 17/82 (21%) vs.	
		` /	IU/ml adjusted odds ratio			10/78 (13%) vs13/82 (16%); (p=NS for A, B, C vs. D)	
		28/78 (36%) vs.	4.69 (95% 2.22-9.88)			Asthenia: 37/81 (46%) vs. 43/82 (52%) vs. 30/78 (38%) vs.	
		38/82 (46%); (A vs.	Age ≤45 years adjusted odds			26/82 (32%); (p<0.05 A, B vs. D, p=0.37 for C vs. D)	
		, ,, ,	ratio 1.59 (0.99-2.57)			Influenza-like illness: 32/81 (40%) vs. 32/82 (39%) vs.	
		p=NS)	,			28/78 (36%) vs. 43/82 (52%); (p=NS for A, B vs. D, p=0.04	
						for C vs. D)	
						Fatigue: 21/81 (26%) vs. 23/82 (28%) vs. 26/78 (33%) vs.	
						30/82 (37%); (p=NS for A, B, C vs. D)	
						Pyrexia: 14/81 (17%) vs. 15/82 (18%) vs. 15/78 (19%) vs.	
						19/82 (23%); (p=NS for A, B, C vs. D)	
						Pruritus: 41/81 (51%) vs. 52/82 (63%) vs. 46/78 (59%) vs.	
						29/82 (35%); (p<0.05 for A, B, C vs. D)	
						Any rash: 40/81 (49%) vs. 36/82 (44%) vs. 37/78 (47%) vs.	
						29/82 (35%); (p=NS for A, B, C vs. D)	
						Nausea: 39/81 (48%) vs. 39/82 (48%) vs. 24/78 (31%) vs.	
						33/82 (40%); (p=NS for A, B, C vs. D)	
						Headache: 36/81 (44%) vs. 32/82 (39%) vs. 37/78 (47%) vs.	
						37/82 (45%); (p=NS for A, B, C vs. D)	
						Depression: 16/81 (20%) vs. 18/82 (22%) vs. 17/78 (22%)	
						vs. 19/82 (23%); (p=NS for A, B, C vs. D)	
						Myalgia: 11/81 (14%) vs. 12/82 (15%) vs. 12/78 (15%) vs.	
						17/82 (21%); (p=NS for A, B, C vs. D)	
						Arthralgia: 8/81 (10%) vs. 8/82 (10%) vs. 20/78 (26%) vs.	
						14/82 (17%); (p=NS for A, B, C vs. D)	
						Anemia: 22/81 (27%) vs. 15/82 (18%) vs. 7/78 (9%) vs.	
						14/82 (17%); (p=NS for A, B, C vs. D)	

Author, Year Country Study Name Ouality	Interferon Regimen	Ribavirin Regimen	Protease Inhibitor Regimen	Eligibility	Exclusion	Number Screened/ Eligible/ Enrolled/ Analyzed	Age Sex Race	Genotype Severity of Liver Disease Proportion Treatment-Naïve
Jacobson, 2011 ¹⁵ International Telaprevir for previously untreated chronic hepatitis C virus infection Overall Quality: Good	A. Pegylated interferon alfa-2a 180 mcg/week for 24 or 48 weeks (Response guided: if HCV RNA undetectable at weeks 4 and 12 then 24 total weeks, 48 weeks otherwise) B. Pegylated interferon alfa-2a 180 mcg/week for 24 or 48 weeks (Response guided: if HCV RNA undetectable at weeks 4 and 12 then 24 total weeks, 48 weeks otherwise) C. Pegylated interferon alfa-2a 180 mcg/week for 24 or 48 weeks otherwise) C. Pegylated interferon alfa-2a 180 mcg/week for 48 weeks	A. 1000-1200 mg/day for 24 or 48 weeks (Response guided: if HCV RNA undetectable at weeks 4 and 12 then 24 total weeks, 48 weeks otherwise) B. 1000-1200 mg/day for 24 or 48 weeks (Response guided: if HCV RNA undetectable at weeks 4 and 12 then 24 total weeks, 48 weeks otherwise) C. 1000-1200 mg/day for 48 weeks	A. Telaprevir 750 mg tid for 12 weeks B. Telaprevir 750 mg tid for 8 weeks C. Placebo for 12 weeks	Treatment naïve Ages 18-70 years of age HCV genotype 1 infection HCV virus confirmed with liver biopsy in the previous year Neutrophil count ≥ 1500 /mm³ Platelets ≥ 90,000 / mm³ Hemoglobin ≥ 12 g/dL in women and ≥ 13 g/dL in men	Decompensated liver disease Hepatocellular carcinoma HBV HIV	NR/ NR/ 1095/ 1088	A vs. B vs. C Age median: 49 vs. 49 vs. 49 Female: 41% vs. 42% vs. 42% Non White: 10% vs. 13% vs. 12%	A vs. B vs. C Genotype 1: all Proportion treatment-naïve: 100% Cirrhosis: 6% overall Minimal or no fibrosis: 28% Elevated transaminases: NR HCV RNA ≥ 800,000: 77% vs. 77% vs. 77%

Author, Year Country Study Name Quality	Duration of Followup	Outcome	Subgroup Analyses	Subgroup Analyses	Histologic Response	Adverse Events	Funding Source
Jacobson, 2011 ¹⁵ International	Up to week	A vs. B vs. C ETR: 314/363 (87%)	A vs. B vs. C Male: 159/214 (74%) vs.	A vs. B vs. C HCV genotype 1a: 138/210	NR	A vs. B vs. C Overall withdrawals: 95/363 (26%) vs. 104/364 (29%) vs.	Vertex, Tibotec
		vs. 295/364 (81%)	147/211 (70%) vs. 94/211	(66%) vs. 152/213 (71%)		159/361 (44%); A or B vs. C p<0.001	
Continued		vs. 229/361 (63%);	(45%); A or B vs. C p<0.001	vs. 85/208 (41%); A or B		Withdrawals for adverse events: 36/363 (10%) vs. 37/364	
		p<0.001 for A or B	Female: 112/149 (75%) vs.	vs. C p<0.001		(10%) vs. 26/361 (7%); p=NS	
		vs. C	103/153 (67%) vs. 64/150	HCV genotype 1b: 111/151		Serious adverse events: 33/363 (9%) vs. 31/364 (9%) vs.	
		GT TD . 054 (0 50 (554))	(43%): A or B vs. C p<0.001	(74%) vs. 118/149 (79%)		24/361 (7%); p=NS	
		SVR: 271/363 (75%)	A 445 110/140	vs. 73/151 (48%); A or B		Deaths: 0 vs. 0 vs. 1 (<1%); p=NS	
		vs. 250/364 (69%) vs. 158/361 (44%);	Age <45 years: 118/142 (83%) vs. 102/139 (73%) vs.	vs. C p<0.001		Fatigue: 207/363 (57%) vs. 211/364 (58%) vs. 206/361	
		p<0.001 for A or B	74/143 (52%); A or B vs. C	Baseline HCV RNA <800K		(57%); p=NS	
		vs. C	p<0.001	IU/ml: 67/85 (79%) vs.		Influenza-like illness 102/363 (28%) vs. 105/364 (29%) vs.	
			Age >45 to <65 years:	64/82 (78%) vs. 57/82		101/361 (28%); p=NS	
			150/214 (70%) vs. 145/222	(70%); A vs. C p=0.16; B		Pyrexia: 95/363 (26%) vs. 108/364 (30%) vs. 87/361 (24%);	
			(65%) vs. 82/216 (38%); A	vs. C p=0.19		p=NS	
			or B vs. C p<0.001	Baseline HCV RNA >800K IU/ml: 183/279 (66%) vs.		Pruritus: 181/363 (50%) vs. 165/364 (45%) vs. 131/361 (36%); p=NS	
			White: 244/325 (75%) vs.	207/281 (74%) vs. 101/279		Rash: 133/363 (37%) vs. 129/364 (35%) vs. 88/361 (24%);	
			220/315 (70%) vs. 147/318	(36%); A or B vs. C		A or B vs. C p<0.01	
			(46%); A or B vs. C p<0.001	p<0.001		Anemia: 135/363 (37%) vs. 141/364 (39%) vs. 70/361	
			Black: 16/26 (62%) vs. 23/40			(19%); A or B vs. C p<0.001	
			(58%) vs. 7/28 (25%); A vs.	No or minimal fibrosis:		Neutropenia: 51/363 (14%) vs. 62/364 (17%) vs. 68/361	
			C p=0.05; B vs. C p=0.04	101/128 (79%) vs. 109/134 (81%) vs. 67/147 (46%); A		(19%); p=NS Depression: 66/363 (18%) vs. 61/364 (17%) vs. 79/361	
			BMI <25: 129/155 (83%) vs.	or B vs. C p<0.001		(22%); p=NS	
			104/145 (72%) vs. 57/130	01 B vs. C p vs.001		Myalgia: 54/363 (15%) vs. 76/364 (21%) vs. 77/361 (21%);	
			(44%); A or B vs. C p<0.001			p=NS	
			*			Arthralgia: 49/363 (13%) vs. 56/364 (15%) vs. 68/361	
						(19%); p=NS	

Author, Year Country Study Name Quality	Duration of Followup	Outcome	Subgroup Analyses	Subgroup Analyses	Histologic Response	Adverse Events	Funding Source
Jacobson, 2011 ¹⁵	•		BMI >25 and <30: 87/129	Portal fibrosis: 104/151			
International			(67%) vs. 92/131 (70%) vs.	(69%) vs. 117/156 (75%)			
			65/144 (45%); A or B vs. C	vs. 67/141 (48%); A or B			
Continued			p<0.001	vs. C p<0.001			
			BMI >30: 55/77 (71%) vs.	Bridging fibrosis: 34/59			
			53/86 (62%) vs. 36/87	(58%) vs. 32/52 (62%) vs.			
			(41%); A vs. C p<0.001, B	17/52 (33%); A vs. C			
			vs. C p=0.02	p=0.02, B vs. C p=0.01			
				Cirrhosis: 11/26 (42%) vs.			
				13/21 (62%) vs. 7/21			
				(33%); A vs. C p=0.04; B			
				vs. C p=0.15			

A 41						Number		Genotype
Author, Year Country			Protease			Screened/ Eligible/	1 4 70	Severity of Liver Disease
Study Name	Interferon		Inhibitor			Engible/ Enrolled/	Age Sex	Proportion Proportion
		Dibayinin Dagiman		Eliaibility	Exclusion		Race	Treatment-Naïve
Quality		Ribavirin Regimen	Regimen	Eligibility		Analyzed		
Kumada 2011 ¹⁶	A:	A:	A:	Diagnosis with chronic	Patients with decompensated	NR/ NR/ 220/	A vs B:	A vs. B:
Japan	Pegylated interferon		1 0	hepatitis C,	liver cirrhosis, hepatitis B surface	189	Age (mean):	Genotype 1a: 1.6 %
T. 1			three times day at	and had not received	antigen, hepatocellular carcinoma		53 vs 55	vs. 0%
				antiviral treatments	or other malignancy, or its		years	Genotype 1b: 98.4%
peginterferon and	for 12 weeks,	_	(q8h) one time a	before, infected with	history, autoimmune hepatitis,		Female: 48%	vs. 100%
ribavirin for	followed by an	12 weeks (24 weeks)	week	HCV-1 confirmed	alcoholic liver disease,		vs 48%	Proportion
treatment-naive	additional 12 weeks		simultaneously	by the sequence analysis	hemochromatosis or chronic liver		Non White:	treatment-naïve:
patients chronically	(24 weeks)	<60 kg – 800 mg	with interferon	in the NS5B region, had	disease other than chronic		Not reported	100%
infected with HCV		\ge 60 - \le 80kg - 800 mg		HCV RNA levels	hepatitis C,		(conducted in	Cirrhosis: NR
of genotype 1 in	B:	>80kg - 1000 mg	B: None	P5.0 log10 IU/ml	depression or schizophrenia, or its		Japan)	
Japan	Pegylated interferon			determined by the	history, or history of suicide			Elevated
	alpha 2b 1.5 mcg/kg	B:		COBAS TaqMan HCV	attempts,			transaminases: NR
Overall Quality:	one time per week	Ribavirin 200 – 600		test, Japanese aged from	chronic renal disease or creatinine			HCV RNA (log 10
Fair	for 12 weeks,	mg/kg (weight-based)		20 to 65 years at the	clearance 650 ml/min at the			<u>IU/ml)</u> 6.7 vs. 6.9
	followed by an	twice a day for 12 weeks,		entry, had the body	baseline, hemoglobin <12 g/dl,			
	additional 12 weeks	followed by an additional		weight between >40 and	neutrophil counts <1500/mm3 or			
	(24 weeks)	12 weeks (24 weeks)		6120 kg, were not	platelet counts			
				pregnant and capable of	<100,000/mm3 at the baseline;			
		<60 kg – 800 mg		contraception until 24	and (h) pregnancy in progress or			
		\geq 60 - \leq 80kg - 800 mg		weeks after the treatment.	planned during the study period of			
		>80kg - 1000 mg		and agreed on the	either partner.			
				admission for	• • • • • • • • • • • • • • • • • • • •			
				15 days since the				
				treatment start				

NR	Author, Year Country Study Name Quality	Duration of Followup	Outcome	Subgroup Analyses	Subgroup Analyses	Histologic Response	Adverse Events	Funding Source
Stomatitis - 24/126(19.0%) vs. 12/63(19.0%) Abdominal discomfort - 23/126(18.3%) vs.12/63(19.0%) Pruritus - 23/126(18.3%) vs.13/63(20.6%)	Kumada 2011 ¹⁶ Japan	>_24 weeks after	A vs. B: ETR: NR SVR: 73% vs.	A vs. B: ETR; NR SVR: Male: 50/66 (75.8%) vs. 18/33 (54.5%), p=0.0400 Female: 42/60 (70.0%) 13/30 (43.3%), p0.0214 Age: <49 years - 35/41 (85.4%) vs. 13/21 (61.9%), p= 0.0543 >50 years - 57/85 (67.1%) vs. 18/42 (42.9%), p= 0.0125 HCV RNA (log10 IU/ml): >7 - 18/26 (69.2%) vs. 5/18 (27.8%), p= 0.0132 <7 - 74/100 (74.0%) vs.	NR		A vs. B: Overall withdrawals: NR Withdrawals for adverse events: NR Serious adverse events: NR Deaths: NR Anemia - 115/126(91.3%) vs. 46/63(73.0%) Pyrexia - 98/126(77.8%) vs. 46/63(73.0%) Leukocytopenia - 86/126(68.3%) vs. 46/63(73.0%) Thrombocytopenia - 81/126(64.3%) vs. 23/63(36.5%) Malaise - 73/126(57.9%) vs. 30/63(47.6%) Serum uric acid increased - 65/126(51.6%) vs. 5/63(7.9%) Serum hyaluronic acid increased - 64/126(50.8%) vs. 25/63(39.7%) Alopecia - 51/126(40.5%) vs. 29/63(46.0%) Headache - 48/126(38.1%) vs. 32/63(50.8%) Skin rashes - 48/126(38.1%) vs. 18/63(28.6%) Anorexia - 42/126(33.3%) vs. 17/63(27.0%) Usmiting - 37/126(29.4%) vs. 9/63(14.3%) Drug eruption - 37/126(29.4%) vs. 2/63(3.2%) Arthralgia - 36/126(28.6%) vs. 15/63(23.8%) Serum triglycerides increased - 36/126(28.6%) vs. 11/63(17.5%) Dysgeusia - 34/126(27.0%) vs. 19/63(30.2%) Nausea - 32/126(25.4%) vs. 7/63(11.1%) Serum creatinine increased - 32/126(25.4%) vs. 0 Erythema at the injection site - 33/26.2%) vs. 21/63(33.3%) Reactions at the injection site - 29/126(23.0%) vs. 16/63(25.4%) Stomatitis - 24/126(19.0%) vs. 12/63(19.0%) Abdominal discomfort - 23/126(18.3%) vs.12/63(19.0%)	

Author, Year Country Study Name Quality	Duration of Followup	Outcome	Subgroup Analyses	Subgroup Analyses	Histologic Response	Adverse Events	Funding Source
Kumada 2011 ¹⁶						Serum bilirubin increased - 22/126(17.5%) vs.	
Japan						13/63(20.6%)	
						Back pain - 21/126(16.7%) vs. 12/63(19.0%)	
Continued						Hyperuricemia - 20/126(15.9%) vs. 2/63(3.2%)	
						Serum phosphorus decreased - 16/126(12.7%) vs.	
						13/63(20.6%)	
						Constipation - 14/126(11.1%) vs. 13/63(20.6%)	
						Erythema - 9/126(7.1%) vs. 13/63(20.6%)	

Author, Year Country Study Name	Interferon		Protease Inhibitor			Number Screened/ Eligible/ Enrolled/	Age Sex	Genotype Severity of Liver Disease Proportion
Quality	Regimen	Ribavirin Regimen	Regimen	Eligibility	Exclusion	Analyzed	Race	Treatment-Naïve
Kwo, 2010 ¹⁷	A. Pegylated	A. 800-1400 mg daily	A. Boceprevir	Treatment naïve	History of decompensated	765/642/520/	A vs. B vs. C	A vs. B vs. C vs. D
US, Canada,	interferon alfa-2b	for 48 weeks	800 mg tid for 48	patients with genotype 1	cirrhosis	520	vs. D vs. E	vs. E
Europe	1.5 mcg/kg weekly	B. 800-1400 mg daily	weeks	Ages 18-60 years	HIV infection		Age: mean 47	Genotype 1: 100%
	for 48 weeks	for 28 weeks	B. Boceprevir	Liver biopsy consistent	Previous organ transplantation		vs. 46 vs. 48	Cirrhosis: 9% vs.
Efficacy of	B. Pegylated	C. 800-1400 mg daily	800 mg tid for 28	with chronic HCV	Other causes of liver disease		vs. 48 vs. 48	7% vs. 6% vs. 7%
boceprevir, an Ns3	interferon alfa-2b	for 48 weeks	weeks	infection within 5 years	Pre-existing psychiatric disease		Female: 39%	vs. 8%
protease inhibitor,	1.5 mcg/kg weekly	D. 800-1400 mg daily	C. Boceprevir	of enrollment	Seizure disorder		vs.41% vs.	Minimal or no
in combination	for 28 weeks	for 28 weeks	800 mg tid for	Hemoglobin ≥ 130 g/L	Cardiovascular disease		44% vs. 50%	fibrosis: NR
with Pegylated	C. Pegylated	E. 800-1400 mg daily	weeks 5 through	in men ≥ 120 g/L in	Hemoglobinopathies		vs. 33%	Elevated
interferon alfa-2b	interferon alfa-2b	for 48 weeks	48 (44 weeks	women	Hemophilia		Non White:	transaminases: NR
and ribavirin in	1.5 mcg/kg weekly		total)	Neutrophils ≥	Poorly controlled diabetes		16% vs. 20%	Treatment-naïve:
treatment-naïve	for 48 weeks	≤ 65 kg: 400 mg bid	D. Boceprevir	1500/mm ³	Autoimmune disease		vs. 17% vs.	100%
patients with	D. Pegylated	66-80 kg: 400 mg every	800 mg tid for	Platelets $\geq 100 \text{K} / \text{mm}^3$			17% vs. 20%	HCV-RNA ≥600K
genotype 1	interferon alfa-2b	morning, 600 mg every	weeks 5 through	Normal bilirubin,				IU/mL: 91% vs.
hepatitis C	1.5 mcg/kg weekly	evening	weeks 28 (24	albumin, and creatinine				92% vs. 90% vs.
infection	for 28 weeks	81-105 kg: 600 mg bid	weeks total)					87% vs. 90%
(SPRINT-1): an	E. Pegylated	>105 kg: 600 mg every	E. Placebo					
open-label,	interferon alfa-2b	morning, 800 mg every						
randomized,	1.5 mcg/kg weekly	evening						
multicentre phase	for 48 weeks							
2 trial								
Overall Quality:								
Fair								

Author, Year Country Study Name Quality Durati Follow	rup Outcome	Subgroup Analyses	Subgroup Analyses	Histologic Response	Adverse Events	Funding Source
Kwo, 2010 ¹⁷ US, Canada, Europe Continued 24 wee after entreatme	nd of vs. E	Black: 4/14 (29%) vs. 7/18 (39%) vs. 8/15 (53%) vs. 6/15 (40%) vs. 2/16 (13%); (A, B, D vs. E p=NS, C vs. E p<0.05) Non black 65/89 (73%) vs. 52/88 (59%) vs. 37/88 (42%) (54%) vs. (A, B, C, D vs. E p<0.05) Male: 40/63 (64%) vs. 33/63 (52%) vs. 41/58 (71%) vs. (52%) vs. 41/58 (71%) vs. 33/51 (65%) 28/70 (40%);	p=NS) non Cirrhosis: 62/97 (66%) vs. 54/100 (54%) vs. 74/97 (76%) vs. 54/96 (56%) vs. 37/96 (39%) (A, B, C, D vs. E p<0.05) Baseline HCV-RNA >600K IU/mL: 63/97 (67%) vs. 52/99 (53%) vs. 67/92 (73%) vs. 48/89 (54%) vs. 30/93 (32%) (A, B C, D vs. E p<0.01) Baseline HCV-RNA < 600K IU/mL: 6/9 (67%) vs.	NR	A vs. B vs. C vs. D vs. E Overall Withdrawals: 40/103 (39%) vs. 30/107 (28%) vs. 27/103 (26%) vs. 27/103 (26%) vs. 16/104 (15%); (A, B vs. E p<0.05; C, D vs. E p=0.055) Withdrawals due to adverse events: 20/103 (19%) vs. 12/107 (11%) vs. 9/103 (9%) vs. 15/103 (15%) vs. 8/104 (8%); (A vs. E p=0.01, B vs. E p=0.38, C vs. E p= 0.78, Dives E p=0.12) Influenza-like illness: 19/103 (18%) vs. 24/107 (22%) vs. 15/103 (15%) vs. 21/103 (20%) vs. 25/104 (24%); p=NS for all comparisons Fatigue: 51/103 (50%) vs. 65/107 (61%) vs. 73/103 vs. 70/103 (68%) vs. 57/104 (55%); (A vs. E p = 0.45; B vs. E p=0.38, C vs. E p=0.02, D vs. E p=0.05) Headache: 44/103 (43%) vs. 52/107 (49%) vs. 54/103 (52%) vs. 41/103 (40%) vs. 45/104 (43%); (A, B, C, D vs. E p=NS) Nausea: 56/103 (103%) vs. 41/107 (38%) vs. 48/103 (47%) vs. 42/103 (41%) vs. 45/104 (43%); (A, B, C, D vs. E p=NS) Pyrexia: 41/103 (40%) vs. 28/107 (26%) vs. 35/103 (34%) vs. 27/103 (26%) vs. 35/104 (34%); (A, B, C, D vs. E p=NS) Chills: 33/103 (32%) vs. 31/107 (29%) vs. 35/103 (34%) vs. 31/103 (30%) vs. 35/104 (34%); (A, B, C, D vs. E p=NS) Dysgeusia: 33/103 (32%) vs. 23/107 (21%) vs. 28/103 (27%) vs. 27/103 (26%) vs. 9/104 (9%); (A, B, C, D vs. E p<0.01) Influenza-like illness: 19/103 (18%) vs. 24/107 (22%) vs. 15/103 (15%) vs. 21/103 (20%) vs. 25/104 (24%); (A, B, C, D vs. E p<0.01) Influenza-like illness: 19/103 (18%) vs. 24/107 (22%) vs. 15/103 (15%) vs. 21/103 (20%) vs. 25/104 (24%); (A, B, C, D vs. E p=NS) Arthralgia: 21/103 (20%) vs. 14/107 (13%) vs. 19/103 (18%) vs. 22/103 (21%) vs. 21/104 (20%); (A, B, C, D vs. E	

Author, Year Country Study Name Quality	Duration of Followup	Outcome	Subgroup Analyses	Subgroup Analyses	Histologic Response	Adverse Events	Funding Source
Kwo, 2010 ¹⁷						Neutrophils <750: 38/103 (37%) vs. 36/107	
US, Canada,						(34%) vs. 37/103 (36%) vs. 21/103 (20%) vs.	
Europe						18/104 (17%); (A, B, C vs. E p<0.01, D vs. E	
						p=0.52)	
Continued						Hemoglobin <100 g/L: 48/103 (47%) vs. 57/107 (53%) vs.	
						48/103 (47%) vs. 51/103 (50%) vs. 25/104 (24%); (A, B, C,	
						D vs. E p<0.01)	
						Platelets <50K / mm3: 1/103 (1%) vs. 4/107 (4%) vs. 4/103	
						(4%) vs. 2/103 (2%) vs. 0/104 (0%); (A, B, C, D vs. E	
						p=NS)	

Author, Year						Number Screened/		Genotype Severity of Liver
Country			Protease			Eligible/	Age	Disease
Study Name	Interferon		Inhibitor			Enrolled/	Sex	Proportion
Quality	Regimen	Ribavirin Regimen	Regimen	Eligibility	Exclusion	Analyzed	Race	Treatment-Naïve
Marcellin, 2011 ¹⁸	A. Pegylated	A. 1000-1200 mg/day	A. Telaprevir	Treatment-naïve	Contraindication to pegylated	176/ 170/ 166/	A vs. B vs. C	A vs. B vs. C vs. D
Europe	interferon alfa-2a	for 24 or 48 weeks	750 mg tid for 12	Ages 18-65 years	interferon or ribavirin	161	vs. D	Genotype 1: all
	180 mcg/week for	B. 800-1200 mg/day for		Chronic HCV genotype	History of drug use		Age median:	Cirrhosis: 2.5% vs.
Telaprevir is	24 or 48 weeks	24 or 48 weeks	B. Telaprevir	1 infection	Documented cirrhosis		47 vs. 46 vs.	2.4% vs. 0 vs.
effective given	B. Pegylated	C. 1000-1200 mg/day	750 mg tid for 12	HCV RNA >10,000	Hepatitis B		40 vs. 49	5.1%
every 8 or 12	interferon alfa-2b	for 24 or 48 weeks	weeks	IU/mL	Hepatocellular cancer		Female: 50%	Minimal or no
Hours with	1.5 mcg/kg/week	D. 800-1200 mg/day for		Neutrophil count ≥ 1500			vs. 52% vs.	fibrosis: 39%
ribavirin and	for 24 or 48 weeks	24 or 48 weeks	1125 mg bid for	mm ³	History or suspicion of alcohol		48% vs. 51	overall
Pegylated	C. Pegylated		12 weeks	Platelets $\geq 100,000 \text{ mm}^3$	abuse		Non White:	Elevated
interferon alfa-2a	interferon alfa-2a	Response guided: 24	D. Telaprevir	Liver fibrosis status			10% vs. 10%	transaminases: NR
or 2b to patients	180 mcg/week for	weeks total if HCV	1125 mg bid for	documented within 18			vs. 10% vs.	Proportion
with chronic	24 or 48 weeks	RNA undetectable from	12 weeks	months			8%	treatment-naïve: all
hepatitis C	D. Pegylated	weeks 4 through 20, 48						HCV-RNA ≥ 800K
0 11 0 11	interferon alfa-2b	weeks total otherwise						IU/mL: 75% vs.
Overall Quality:	1.5 mcg/kg/week							81% vs. 83% vs.
Fair	for 24 or 48 weeks							87%
	Dosmonso ovidade							
	Response guided: 24 weeks total if							
	HCV RNA							
	undetectable from							
	weeks 4 through							
	,							
	20, 48 weeks total otherwise							

Author, Year Country							
Study Name	Duration of				Histologic		Funding
Quality	Followup	Outcome	Subgroup Analyses	Subgroup Analyses	Response	Adverse Events	Source
Marcellin,	24 weeks	A vs. B vs. C vs. D	NR	NR	NR	A vs. B vs. C vs. D vs. E	Janssen,
2011 ¹⁸	after end of	ETR: 37/40 (93%)				Overall withdrawals: 10/40 (25%) vs. 8/42 (19%) vs. 11/40	Vertex
Europe	treatment	vs. 37/42 (88%) vs.				(28%) vs. 17/39 (44%);	Pharma-
		37/40 (93%) vs.				Withdrawals due to adverse events: 3/40 (7.5%) vs. 2/42	ceuticals
Continued		34/39 (87%);				(5%) vs. 4/40 (10%) vs. 4/39 (10%)	
						Nausea: 18/40 (45%) vs. 14/42 (33%0 vs. 16/40 (40%) vs.	
		Pooled A+B (TID				23/39 (59%)	
		telaprevir) vs. C+D				Fatigue: 15/40 (38%) vs. 15/42 (36%) vs. 16/40 (40%) vs.	
		(BID telaprevir)				15/39 (39%)	
		p=NS				Influenza-like illness: 16/40 (40%) vs. 19/42 (45%) vs.	
		Pooled A+C (alpha-				11/40 (28%) vs. 20/39 (51%)	
		2a) vs. B + D (alpha-				Pyrexia: 9/40 (23%) vs. 15/42 (36%) vs. 9/40 (23%) vs.	
		2b) p=NS				12/39 (31%)	
		SVR: 34/40 (85%)				Depression: 7/40 (18%) vs. 9/42 (21%) vs. 4/40 (10%) vs. 9/39 (23%)	
		vs. 34/42 (81%) vs.				Pruritus: 19/40 (48%) vs. 23/42 (55%) vs. 20/40 (50%) vs.	
		33/40 (83%) vs.				25/39 (64%)	
		32/39 (82%)				Rash: 29/40 (73%) vs. 23/42 (55%0 vs. 20/40 (50%) vs.	
		32/37 (02/0)				25/39 (64%)	
		Pooled A+B (TID				Anemia: 18/40 (45%) vs. 14/42 (33%) vs. 18/40 (45%) vs.	
		telaprevir) vs. C+D				20/39 (51%)	
		(BID telaprevir)				Leukopenia: 9/40 (23%) vs. 9/42 (21%) vs. 9/40 (23%) vs.	
		p=NS				10/39 (26%)	
		Pooled A+C (alpha-					
		2a) vs. B + D (alpha-				Pooled A+C (alpha-2a) vs. B + D (alpha-2b) - all	
		2b) p=NS				comparisons p=NS	
						Pooled A+B (TID telaprevir) vs. C+D (BID telaprevir) - all	
						comparisons p=NS	

Author, Year Country Study Name Interferon Quality Regimen Ribay	Protease Inhibitor avirin Regimen Regimen	Eligibility	Exclusion	Screened/ Eligible/ Enrolled/ Analyzed		Genotype Severity of Liver Disease Proportion Treatment-Naïve
McHutchison, 2009 ¹⁹ US Protease Inhibition for Viral Evaluation 1 (PROVE1) A. Peg interferon alfa-2a B. Rib mg dai meekly for 24 weeks B. Pegylated interferon alfa-2a mg dai meekly for 48 weeks C. Rib mg dai meekly for 48 weeks C. Pegylated interferon alfa-2a 1000 r 180 mcg weekly for 12 weeks A. Rib mg dai meg dai meekly mg dai meekly for 48 meeks mg dai neekly for 48 weeks mg dai neekly for 12 weeks	Ribavirin1000-1200 daily for 24 weeks Ribavirin 800-1400 daily for 12 weeks Ribavirin 800-1400 daily for 12 weeks Ribavirin1000-1200 daily for 48	Treatment naïve patients ages 18-65 years, neutrophils ≥ 1500 / mm3, platelets ≥ 90K / mm3, normal hemoglobin	decompensated liver disease, hepatocellular carcinoma, cirrhosis (liver biopsy within 2 years)	329/ 263/ 263/ 250	49 vs. 50 vs.	Genotype 1: all Portal or Bridging fibrosis: 70% vs. 57% vs. 76% vs. 75% Treatment-naïve: all

Author, Year Country Study Name Quality	Duration of Followup	Outcome	Subgroup Analyses	Subgroup Analyses	Histologic Response	Adverse Events	Funding Source
McHutchison, 2009 ¹⁹ US Continued	Up to 24 weeks following treatment completion	A vs. B vs. C vs. D ETR: 45/79 (57%) vs. 51/79 (65%) vs. 12/17 (71%) vs. 35/75 (47%) (A, C vs. D p=NS, B vs. D p=0.03) SVR: 48/79 (61%) vs. 53/79 (67%) vs. 6/17 (35%) vs. 31/75 (41%); (A vs. D p=0.02, B vs. D p=0.002, C vs. D p=NS)	NR	NR	NR	A vs. B vs. C vs. D Overall withdrawals: 26/79 (33%) vs. 25/79 (32%) vs. 4/17 (24%) vs. 17/75 (23%) Withdrawals due to adverse events (telaprevir regimens A+B+C vs. D): 37/175 (21%) vs. 8/75 (11%) Fatigue: 70% vs. 73% vs. 82% vs. 76% Nausea: 56% vs. 48% vs. 65% vs. 29% Influenza-like illness: 49% vs. 40% vs. 24% vs. 23% Pruritus: 48% vs. 40% vs. 24% vs. 23% Headache: 47% vs. 43% vs. 53% vs. 60% Rash: 60% vs. 61% vs. 53% vs. 41% Vomiting: 24% vs. 20% vs. 18% vs. 12% Arthralgia: 17% vs. 22% vs. 24% vs. 21% Myalgia: 11% vs. 19% vs. 18% vs. 24% Chills: 10% vs. 23% vs. 18% vs. 19% Anemia: 37% vs. 29% vs. 35% vs. 27% Neutropenia: 14% vs. 24% vs. 0% vs. 24%	Vertex Pharmaceut icals

Author, Year Country			Protease			Number Screened/ Eligible/	Age	Genotype Severity of Liver Disease
Study Name	Interferon		Inhibitor			Enrolled/	Sex	Proportion
Quality	Regimen	Ribavirin Regimen	Regimen	Eligibility	Exclusion	Analyzed	Race	Treatment-Naïve
Poordad, 2011 ²⁰	A: Pegylated	A: 600-1400 mg	A: Boceprevir	No previous treatment	Liver disease of other cause	1472/NR/1099	A vs. B vs. C	A vs. B vs. C
USA and Europe	interferon alfa-2b	(weight-based) daily for	0 3	for HCV infection	Decompensated cirrhosis	/1097		Genotype 1: 100%
	1.5 µg/kg/week for	48 weeks	mouth tid from	Age 18 years or older	Renal insufficiency		vs. 50 vs. 49	Cirrhosis
Serine Protease	48 weeks	B: 600-1400 mg	weeks 5 to 28	Weight 40 to 125 kg	HIV or hepatitis B infection		years	(METAVIR
Inhibitor Therapy		(weight-based) daily for	(24 weeks total)	Chronic infection with	Pregnancy or current breast-		Female: 40%	fibrosis score 3 or
2 (SPRINT-2)	B: Pegylated	48 weeks	B: Boceprevir	HCV genotype 1	feeding		vs. 38% vs.	4): 11% vs. 9% vs.
	interferon alfa-2b	-if HCV RNA	800 mg by	Plasma HCV RNA level	Active cancer		43%	7%
Overall Quality:	1.5 µg/kg 1x/week	undetectable from week	mouth tid from	>=10,000 IU/mL			Non White:	Treatment-naïve:
Fair	for 48 weeks	8 through 24 treatment	weeks 5 to 48				19% vs. 17%	100%
	-if HCV RNA	completed	(44 weeks total)				vs. 18%	
	undetectable from	-if HCV RNA	C: Placebo					
	week 8 through 24	detectable at any point						
	treatment	from week 8 through 23						
	completed	ribavirin continued						
	-if HCV RNA	through week 48						
	detectable at any	C: 600-1400 mg						
	point from week 8	(weight-based) daily for						
	through 23	48 weeks						
	Pegylated							
	interferon	*<51 kg: 600mg/day						
	continued through	51-65 kg: 800mg/day						
	week 48	66 - 75 kg: 1000mg/day						
	C. Dogwlatad	76 - 105 kg:						
	C: Pegylated	1200mg/day						
	interferon alfa-2b	>105 kg: 1400mg/day						
	1.5 µg/kg 1x/week							
	for 48 weeks							

Author, Year							1
Country							
Study Name	Duration of				Histologic		Funding
Quality	Followup	Outcome	Subgroup Analyses	Subgroup Analyses	Response		Source
Poordad, 2011 ²⁰	72 weeks (24	A vs. B vs. C	A vs. B vs. C	A vs. B vs. C	NR	A vs. B vs. C	Schering-
USA and Europe	`	ETR: 277/366 (76%)	Black: 29/55 (53%) vs. 22/52	METAVIR score 0, 1, or 2:	1,11	Overall withdrawals: 152/367 (41%) vs. 139/368 (38%) vs.	Plough
Corruna Europe	treatment	vs. 261/368 (71%)	(42%) vs. 12/52 (23%)	211/313 (67%) vs. 213/319		205/364 (56%) (p<0.001 for A or B vs. C)	(now
Continued	end)	vs. 191/363 (53%)	(p=0.004 for A vs. C, p=0.04	(67%) vs. 123/328 (38%)		Withdrawals due to adverse events: 60/366 (16%) vs.	Merck)
	,	(p<0.001 for A or B	for B vs. C)	(p<0.001 for A or B vs. C)		45/368 (12%) vs. 57/363 (16%) (p>0.05)	,
		vs. C)	Non black: 197/313 (63%)	METAVIR score 3 or 4:		Deaths: 1/366 (<1%) vs. 1/368 (<1%) vs. 4/363 (1%)	
		/	vs. 192/314 (61%) vs.	22/42 (52%) vs. 14/34		(p>0.05)	
		vs. 233/368 (63%)	102/308 (33%) (p<0.001 for	(41%) vs. 9/24 (38%)		Serious adverse event: 45/366 (12%) vs. 42/368 (11%) vs.	
		vs. 137/363 (38%)	A or B vs. C)	(p=0.31 for A vs. C and		31/363 (9%) (p>0.05)	
		(p<0.001 for A or B	Male: 145/221 (66%) vs.	p=1.0 for B vs. C)		Fatigue: 209/366 (57%) vs. 196/368 (53%) vs. 217/363	
		vs. C)	149/229 (65%) vs. 72/206	Low viral load (<=800,000		(60%) (p>0.05)	
		,	(35%) (p<0.001 for A or B	IU/mL): 45/53 (85%) vs.		Headache: 167/366 (46%) vs. 168/368 (46%) vs. 153/363	
			vs. C)	41/54 (76%) vs. 35/55		(42%) (p>0.05)	
			Female: 97/145 (67%) vs.	(64%)		Nausea: 159/366 (43%) vs. 175/368 (48%) vs. 153/363	
			84/139 (60%) vs. 65/157	High viral load: 197/313		(42%) (p>0.05)	
			(41%) (p<0.001 for A or B	(63%) vs. 192/314 (61%)		Pyrexia: 118/366 (32%) vs. 123/368 (33%) vs. 121/363	
			vs. C)	vs. 102/308 (33%) (p<0.001		(33%) (p>0.05)	
			Age <=40 years: 41/59	for A or B vs. C)		Chills: 121/366 (33%) vs. 134/368 (36%) vs. 102/363 (28%)	
			(69%) vs. 37/51 (73%) vs.	Genotype 1a: 118/187		(p=0.15 for A vs. C, p=0.02 for B vs. C)	
			35/67 (52%) (p<0.001 for A	(63%) vs. 106/179 (59%)		Dysgeusia: 156/366 (43%) vs. 137/368 (37%) vs. 64/363	
			or B vs. C)	vs. 62/177 (35%) (p<0.001		(18%) (p<0.001 for A or B vs. C)	
			Age >40 years: 201/307	for A or B vs. C)		Neutrophil count <750 per mm ³ : 119/366 (32%) vs. 108/368	
			(65%) vs. 196/317 (62%) vs.	Genotype 1b: 93/133 (70%)		(29%) vs. 66/363 (18%) (p<0.001 for A or B vs. C)	
			102/296 (34%) (p<0.001 for	vs. 89/134 (66%) vs. 51/128		Neutrophil count <500 per mm ³ : 29/366 (8%) vs. 21/368	
			A or B vs. C)	(40%) (p<0.001 for A or B		(6%) vs. 16/363 (4%) (p>0.05)	
			Weight <75 kg: 83/131	vs. C)		Use of granulocyte stimulating agent: 31/366 (8%) vs.	
			(63%) vs. 82/131 (63%) vs.	Cirrhosis: 10/24 (42%) vs.		43/368 (12%) vs. 21/363 (6%) (p=0.20 for A vs. C, p=0.006	
			67/146 (46%) (p<0.001 for A	5/16 (31%) vs. 6/13 (46%);		for B vs. C)	
			or B vs. C)	p=NS for A or B vs. C		Platelet count <50,000 per mm ³ : 14/366 (4%) vs. 12/368	
			Weight $>=75$ kg): $159/235$	Non cirrhosis: 223/331		(3%) vs. 5/363 (1%) (p=0.99 for A or B vs. C)	
			(68%) vs. 151/237 (64%) vs.	(67%) vs. 222/337 (66%)		Hemoglobin <8.0 g/dl: 13/366 (4%) vs. 9/368 (2%) vs.	
			70/217 (32%) (p<0.001 for A	vs. 126/339 (37%);		6/363 (2%) (p>0.05)	
			or B vs. C)	(p<0.001 for A or B vs. C)		Red-cell transfusion: 9/366 (2%) vs. 11/368 (3%) vs. 2/363	
						(1%) (p=0.06 vs. A vs. C and p=0.02 for B vs. C)	
						Erythropoietin use: 159/366 (43%) vs. 159/368 (43%) vs.	
						87/363 (24%) (p<0.001 for A or B vs. C)	

Author, Year Country Study Name	Interferon		Protease Inhibitor			Number Screened/ Eligible/ Enrolled/	Age Sex	Genotype Severity of Liver Disease Proportion
-	Regimen	Ribavirin Regimen	Regimen	Eligibility	Exclusion	Analyzed	Race	Treatment-Naïve
Sherman, 2011 ²¹ Europe and US Response-Guided Telaprevir Combination Treatment for Hepatitis C Virus Infection Overall Quality: Fair	Regimen A. Pegylated interferon alfa-2a 180 mcg weekly for 24 weeks B. Pegylated interferon alfa-2a 180 mcg weekly for 48 weeks C. Pegylated interferon alfa-2a 180 mcg weekly for 48 weeks C. Pegylated interferon alfa-2a 180 mcg weekly for 48 weeks (not randomized) Randomization to A and B was done at week 20 in those with an extended rapid virologic response (undetectable HCV RNA in week 4 and week 12). Subjects not achieving ERVR were assigned to	Ribavirin Regimen A. Ribavirin 1000-1200 mg daily for 24 weeks B. Ribavirin 1000-1200 mg daily for 48 weeks C. Ribavirin 1000-1200 mg daily for 48 weeks (not randomized) Randomization to A and B was done at week 20 in those with an extended rapid virologic response (undetectable HCV RNA in week 4 and week 12). Subjects not achieving ERVR were assigned to group C	Regimen A. Telaprevir 750 mg tid for 12 weeks B. Telaprevir 750 mg tid for 12 weeks C. Telaprevir 750 mg for 12 weeks	Treatment-naïve Ages between 18 and 70 years Chronic HCV genotype 1 infection Detectable HCV RNA Diagnosis for at least 6 months before screening Neutrophils ≥ 1500/mm³ Hemoglobin≥12 g/dL for women and ≥13 g/dL for men Platelets ≥ 90K/mm³ Liver biopsy in past year	HIV HBV Hepatic decompensation Clinically significant liver disease of other etiology Active cancer in previous 5 years (except basal-cell	NR/544/322/3 22 Subjects treated for 20 weeks prior to randomization . Only subjects who completed 20 weeks and had an early rapid virologic response were randomized.	Race A vs. B Age median: 51 vs. 50 Female: 36% vs. 39% Non White: 17% vs. 18%	Treatment-Naïve A vs. B Genotype 1: all Treatment-naïve: 100% Cirrhosis: 11% vs. 8% Minimal or no fibrosis: 27% HCV RNA ≥ 800K IU/ml: 77% vs. 79%

Author, Year Country Study Name Quality	Duration of Followup	Outcome	Subgroup Analyses	Subgroup Analyses	Histologic Response	Adverse Events	Funding Source
Sherman, 2011 ²¹	72 weeks	A vs. B vs. C	A vs. B	A vs. B	NR	A vs. B vs. C	Vertex,
Europe and US		ETR: 159/162 (98%) vs. 154/160 (96%)	Black: 15/17 (88%) vs. 15/17 (88%)	HCV genotype 1a: 103/115 (90%) vs. 10/117 (88%)		Overall withdrawals (after randomization): 1/162 (1%) vs. 41/160 (26%) vs. 39/118 (33%)	Tibotec
Continued		vs. 97/118 (82%); As B p=NS	White: 126/135 (93%) vs. 114/131 (87%)	HCV genotype 1b: 45/46 (98%) vs. 37/43 (86%)		Withdrawals for adverse events: 1/162 (1%) vs. 20/160 (13%) vs. 12/118 (10%)	
		SVR: 149/162 (92%) vs. 140/160 (88%) vs. 76/118 (64%); A non inferior to B	Asian/other: 8/10 (80%) vs. 11/12 (92%) BMI>=30: 55/61 (90%) vs. 43/49 (88%)	Bridging fibrosis or cirrhosis: 31/38 (82%) vs. 29/33 (88%) no Bridging fibrosis or		Serious adverse events: 4/162 (2) vs. 16/160 (10%) vs. 7/118 (6%) Deaths: NR Fatigue: 110/162 (68%) vs. 111/160 (69%) vs. 81/118 (69%)	
			BMI>=25 to <30: 51/56 (91%) vs. 46/51 (90%) BMI <25: 42/44 (95%) vs51/60 (85%)	cirrhosis: 118/124 (95%) vs. 111/127 (87%)		Nausea: 71/162 (44%) vs. 76/160 (48%) vs. 61/118 (52%) Diarrhea: 48/162 (30%) vs. 54/160 (34%) vs. 38/118 (32%) Pruritus: 95/162 (59%) vs. 83/160 (52%) vs. 55/118 (47%) Rash: 60/162 (37%) vs. 62/160 (39%) vs. 47/118 (40%) Headache: 61/162 (38%) vs. 57/160 (36%) vs. 51/118 (43%) Insomnia: 50/162 (31%) vs. 62/160 (39%) vs. 44/118 (37%)	
						Anemia: 68/162 (42%) vs. 66/160 (41%) vs. 38/118 (32%)	

Evidence Table 4. Quality rating: Trials of protease inhibitors plus pegylated interferon and ribavirin

Author, Year	Randomization adequate?	Allocation concealment adequate?	Groups similar at baseline?	Eligibility criteria specified?	Outcome assessors masked?	Care provider masked?	Patient masked?	Attrition and withdrawals reported?	Loss to followup: differential/ high?	Intention- to-treat analysis	Quality	Funding
Hezode 2009 ¹⁴	Unclear	Yes	Yes	Yes	Unclear	Unclear	Unclear	Yes	Unclear	Yes	Fair	Vertex Pharma- ceuticals
Jacobson 2011 ¹⁵	Unclear	Yes	Yes	Yes	Yes	Yes	Yes	Yes	No	Yes	Good	Vertex, Tibotec
Kumada 2011 ¹⁶	Unclear	Unclear	Yes	Yes	No	No	No	Yes	No	Yes	Fair	NR
Kwo 2010 ¹⁷	Yes	Yes	Yes	Yes	No	No	No	Yes	No	Yes	Fair	Merck
Marcellin 2011 ¹⁸	Unclear	Unclear	Yes	Yes	No	No	No	Yes	No	No	Fair	Janssen, Vertex Pharma-ceuticals
McHutchison 2009 ¹⁹	Unclear	Unclear	Yes	Yes	Unclear	Unclear	Unclear	Yes	Unclear	Yes	Fair	Vertex Pharma- ceuticals
Poordad 2011 ²⁰	Yes	Yes	Yes	Yes	Yes	Yes	Yes	Yes	Yes	Yes	Fair	Schering-Plough now Merck)
Sherman 2011 ²¹ ILLUMINATE Study	Unclear	Yes	Yes	Yes	No	No	No	No	No	Yes	Fair	Vertex

Evidence Table 5. Trials of dual therapy with pegylated interferon plus ribavirin: duration effects

						Number		Genotype
Author, Year						Screened/		Severity of Liver
Country			Protease			Eligible/	Age	Disease
Study Name			Inhibitor			Enrolled/	Sex	Proportion
Quality	Interferon Regimen	Ribavirin Regimen	Regimen	Eligibility	Exclusion	Analyzed	Race	Treatment-Naïve
Andriulli, 2009 ²²	A: Pegylated interferon	A: 1000-1200	None	Treatment-naïve	Neutrophils <3000	NR/NR/149/	A vs. B:	A vs. B:
Italy	alpha-2a 180 mcg /	mg/day depending		Ages 18-70 years	Platelets < 80K	120		
	week for 12 weeks	of body weight for 6		Detectable HCV-RNA	Hemoglobin <12 g/dL for females and		Age	Genotype 1: none
Early	B: Pegylated interferon	weeks		levels	<13 g/dL for males		mean: 53	Treatment-naïve: all
discontinuation	alpha-2a 180 mcg /	B: 1000-1200		Infection with genotype	HIV co-infection		vs. 53	Fibrosis stage 3 or
of ribavirin in	week for 12 weeks	mg/day depending		2 or 3	Alcohol intake >30 g daily		Female:	platelets <140K: 14%
HCV-2 and		of body weight for		Abnormal ALT	Drug abuse		41% vs.	vs. 10%
HCV-3 patients		12 weeks			Chronic disease		51%	HCV-RNA >600K:
responding to					Psychiatric disorders		non white:	64% vs. 52%
Peg-interferon		Patients with rapid			Autoimmune diseases		NR	Cirrhosis: NR
alpha-2a and		virologic response			Pregnancy or lactation			
ribavirin		(undetectable HCV-						
		RNA) at week 4						
Overall Quality:		were randomized to						
Fair		A or B above						

Author, Year Country Study Name Quality	Duration of Followup	Outcome	Subgroup Analyses	Subgroup Analyses	Histologic Response	Adverse Events	Funding Source
Andriulli, 2008 ²²	Followup	A vs. B:	NR (only one arm	A vs. B:	NR	A vs. B:	Investigator
Italy	visits at 24		reported)			Overall withdrawals: NR	funded
		SVR: 32 /59 (54%) vs.		Baseline HCV		Withdrawals for adverse events: 5/120 (4%) vs.	
Continued	completion of	50 / 61 (82%); p<0.001		RNA<300K: 12/14		2/24(8%); p=0.33	
	treatment			(86%) vs. 17/21		Serious adverse events: NR	
				(81%); p=NS		Deaths: NR	
				Baseline HCV RNA		Interferon-related adverse events: 66% vs. 63%	
				300K-700K: 7/10		Neutrophils <1000 at 12 weeks: 17% vs. 16%	
				(70%) vs. 10/14			
				(71%); p=NS			
				Baseline HCV RNA			
				>700K: 13/35 (37%)			
				vs. 23/26 (88%);			
				p<0.001			

Author, Year Country Study Name			Protease Inhibitor			Number Screened/ Eligible/ Enrolled/	Age Sex	Genotype Severity of Liver Disease Proportion
Quality	Interferon Regimen	Ribavirin Regimen		Eligibility	Exclusion	Analyzed	Race	Treatment-Naïve
Berg, 2006 ²³ Germany Extended treatment duration for hepatitis C virus type 1: Comparing 48 vs. 72 weeks of	A: Pegylated interferon alfa-2a 180 mcg/week for 48 weeks B: Pegylated interferon alfa-2a 180 mcg/week for 72 weeks	A: 400 mg twice daily for 48 weeks B: 400 mg twice daily for 72 weeks	None	Treatment naïve Ages 18-70 years of age HCV genotype 1 infection HCV RNA >1000 IU/mL Increased ALT at screening Liver biopsy within the preceding 18 months	HCV genotype other than type 1 Decompensated liver disease Liver disease of other etiology HBV or HIV co-infection Autoimmune disorder Clinically significant cardiovascular disease Organ grafts Systemic infections	467/459/455 /455	A vs. B: Age mean: 43 vs. 43 Female: 44% vs. 46% non	A vs. B: Genotype 1: all Treatment-naïve: all Fibrosis stage 3-4: 7% vs. 9% HCV RNA (log IU/mL) mean: 5.8 vs. 5.8
pegylated interferon alfa- 2a plus ribavirin Overall Quality: Fair				showing chronic hepatitis Neutrophils > 1500 Platelets > 90K Hemoglobin > 12g/dL for women and > 13 g/dL for men Creatinine <1.5 mg/dL	Clinically significant bleeding disorders Malignant neoplasm Concomitant immunosuppressive medication use Alcohol or drug abuse in the past year		White: 3% vs. 5%	

Author, Year Country Study Name Quality	Duration of Followup	Outcome	Subgroup Analyses	Subgroup Analyses	Histologic Response	Adverse Events	Funding Source
Berg T, 2006 ²³ Germany Continued	Followup visits at 24 weeks after completion of treatment	A vs. B: SVR: 121/230 (53%) vs. 121/225 (54%); p=0.8	A vs. B: SVR: White: 115/222 (52%) vs. 115/213 (54%); p=NS non White: 6/8 (75%)	A vs. B: Genotype 1b: 75/155 (48%) vs. 66/132 (50%); p=NS Genotype 1a: 38/60 (63%) vs. 40/67	NR	A vs. B: Overall withdrawals: 55/230 (24%) vs. 92/225 (41%); p<0.001 Withdrawals due to adverse events: 21/230 (9%) vs. 26/225 (12%); p=NS Serious adverse events: 15.6% vs. 11.1%; p=NS Deaths: NR	Roche
			vs. 6/12 (50%); p=NS Male: 73/128 (57%) vs. 66/122 (54%); p=NS Female: 48/102 (47%) vs. 55/103 (53%); p=NS	(60%); p=NS Genotype 1a/1b: 4/6 (67%) vs. 13/18 (72%); p=NS Fibrosis Stage 0-2: 117/214 (55%) vs. 116/205 (57%); p=NS Fibrosis Stage 3-4: 4/16 (25%) vs. 5/20 (25%); p=NS			

						Number		Genotype
Author, Year						Screened/		Severity of Liver
Country			Protease			Eligible/	Age	Disease
Study Name			Inhibitor			Enrolled/	Sex	Proportion
Quality	Interferon Regimen	Ribavirin Regimen	Regimen	Eligibility	Exclusion	Analyzed	Race	Treatment-Naïve
Berg T, 2009 ²⁴	A: Pegylated interferon	A: 800-1400 mg	None	Treatment-naïve	HCV genotype other than type 1	438/433/433	A vs. B:	A vs. B:
Germany	alfa-2b 1.5 mcg/kg for a	daily for a duration		Ages 18-70 years	Decompensated liver disease	/433		
	duration determined by	determined by the		HCV genotype 1	HBV or HIV co-infection		Age	Genotype 1: all
Continued	the time required to	time required to		infection	Liver disease of other causes		mean: 43	
	achieve HCV-RNA	achieve HCV-RNA		Positive test for anti-	Autoimmune disorder		vs. 43	Treatment-naïve: all
	negativity at weeks	negativity at weeks		HCV antibodies	Concomitant immunosuppressive			
	3,4,5,6,7, or 8 (times a	3,4,5,6,7, or 8		HCV-RNA >1000	medication use		Female:	Fibrosis stage 3-4:
	factor of 6)	(times a factor of 6)		IU/mL	Clinically significant bleeding disorders		46% vs.	15% vs. 13%
	B: Pegylated interferon	B: 800-1400 mg		Increased ALT	Clinically significant cardiac		43%	
	alfa-2b 1.5 mcg/kg for	daily for 48 weeks		Liver biopsy within 24	abnormalities			HCV-RNA mean: 5.7
	48 weeks			months of enrollment	Organ grafts		Non	vs. 5.7
				confirming chronic	Systemic infection		White:	
				hepatitis	Preexisting severe psychiatric condition		NR	
				Neutrophils > 1500	Neoplastic disease			
				Platelets >80K	Excessive alcohol intake			
				Hemoglobin >12 g/dL	Drug abuse in the past year			
				for females and >13 g/dL	Unwillingness to use contraception			
				for males				
				Creatinine <1.5 mg/dL				

Author, Year Country Study Name	Duration of				Histologic		Funding
Quality	Followup	Outcome	Subgroup Analyses	Subgroup Analyses	Response	Adverse Events	Source
Berg, 2009 ²⁴ Germany	visits at 24	A vs. B:	NR	NR	NR	A vs. B: Overall withdrawals: 63/208 (30.3%) vs. 71/225	Schering- Plough
Individualized treatment strategy according to early viral kinetics in hepatitis C virus type 1-infected patients	weeks after completion of treatment	SVR: 72/208 (35%) vs. 108/225 (48%); p=0.005				(31.6%); p=NS Withdrawals for adverse events: 4 / 208 (1.9%) vs. 7/226 (3.1%); p=NS Serious adverse events: 5/208 (2.6%) vs. 14/225 (6.2%); p=NS Deaths: NR Other edverse events not reported	
Quality: Poor						Other adverse events not reported	

Author, Year Country Study Name Quality	Interferon Regimen	Ribavirin Regimen	Protease Inhibitor Regimen	Eligibility	Exclusion	Number Screened/ Eligible/ Enrolled/ Analyzed	Age Sex Race	Genotype Severity of Liver Disease Proportion Treatment-Naïve
Brandao, 2006 ²⁵	A: Pegylated interferon	A: 400 mg twice	None	Treatment naïve	Treatment with systemic antivirals,	NR/NR/63/6	A vs. B:	A vs. B:
Brazil	alfa-2a 180 mcg/week	daily for 24 weeks		Aged >18 years	antineoplastics, immunomodulators, or	3		
	for 24 weeks	B: 400 mg twice		HCV RNA >1000 IU/mL	any other investigational drugs with		Age	Genotype 1: all
Continued	B: Pegylated interferon	daily for 48 weeks		ALT above upper limit	perceived effect against HCV		mean: 41	HCV RNA >800,000
	alfa-2a 180 mcg/week			of normal on two			vs. 41	IU/mL: 72% v 61%
	for 48 weeks			occasions within the last			Female:	Bridging fibrosis:
				6 months			41% vs.	16% vs. 6%
				Liver biopsy in the last			39%	
				18 month consistent with			Non	
				chronic hepatitis C			white:	
							19% vs.	
							16%	

Author, Year Country Study Name Quality	Duration of Followup	Outcome	Subgroup Analyses	Subgroup Analyses	Histologic Response	Adverse Events	Funding Source
Brandao, 2006 ²⁵ Brazil Continued	Followup visits at 24 weeks after completion of treatment	A vs. B: SVR: 6/32 (19%) vs. 15/31 (48%)	NR	A vs. B: Baseline HCV RNA <800K IU/mL: 3/9 (33%) vs. 7/12 (58%); p=NS Baseline HCV RNA >800K IU/mL: 3/23 (13%) vs. 8/19 (43%); p=NS Bridging fibrosis: 0/5 (0%) vs. 1/2 (50%); p=0.04 non bridging fibrosis: 6/27 (22%) vs. 14/29 (48%); p=0.04	NR	A vs. B: Overall withdrawals: 2/32 (6%) vs. 0/31 (0%); p=NS Withdrawals for adverse events: 2/32 (6.3 %) vs. 0/31 (0%); p=NS Serious adverse events: 3/32 (9.4%) vs. 1/31 (3.2%); p=NS Deaths: NR Headache: 14/32 (44%) vs. 16/31 (52%); p=NS Pyrexia: 13/32 (41%) vs. 16/31 (52%); p=NS Influenza-like illness 8/32 (25%) vs. 10/31 (32%); p=NS Neutropenia: 8/32 (25%) vs. 14/31 (45%); p=NS Myalgia: 7/32 (22%) vs. 14/31 (45%); p=NS Asthenia: 7/32 (22%) vs. 13/31 (42%); p=NS Pruritus: 9/32 (28%) vs. 6/31 (19%); p=NS Irritability: 8/32 (25%) vs. 7/31 (23%); p=NS Thrombocytopenia: 3/32 (9%) vs. 7/31 (23%); p=NS Leukopenia: 4/32 (13%) vs. 6/31 (19%); p=NS Nausea: 6/32 (19%) vs. 9/31 (29%); p=NS Diarrhea: 9/32 (22%) vs. 8/31 (26%); p=NS Diarrhea: 9/32 (22%) vs. 5/31 (16%); p=NS Depression: 5/32 (16%) vs. 5/31 (16%); p=NS Rigors: 3/32 (9%) vs. 6/31 (19%); p=NS Cough: 4/32 (13%) vs. 7/31 (23%); p=NS	Roche

Author, Year Country Study Name			Protease Inhibitor			Number Screened/ Eligible/ Enrolled/	Age	Genotype Severity of Liver Disease Proportion
	Interferon Regimen	Ribavirin Regimen		Eligibility	Exclusion			
Study Name Quality Buti, 2010 ²⁶ International Randomized trial of pegylated interferon alfa- 2b and ribavirin for 48 or 72 weeks in patients with hepatitis C virus genotype 1 and slow virologic response Overall Quality: Fair	undetectable HCV RNA at week 12 were continued until week 48 (group C). Subjects with a 2 log drop in HCV RNA at week 12 and detectable HCV RNA at 12 weeks were continued for another 12 weeks. Subjects with undetectable HCV RNA at week 24 (slow responders) were randomized to groups A or B A: Pegylated interferon alfa-2b 1.5 mcg/kg/week for 48 weeks	undetectable HCV RNA at week 12 were continued until week 48 (group C). Subjects with a 2 log drop in HCV RNA at week 12 and detectable HCV RNA at 12 weeks were continued for another 12 weeks. Subjects with undetectable HCV RNA at week 24 (slow responders) were randomized to groups A or B A: 800-1400 mg/day based on body weight for 48 weeks B: 800-1400 mg/day based on body weight for 72 weeks Nonrandomized	Inhibitor Regimen None	Eligibility Treatment naïve Aged 18-70 years Compensated HCV with confirmed diagnosis of hepatitis by ALT and liver biopsy	Exclusion Weight >125 kg HIV HBV Liver disease of other etiologies	Enrolled/ Analyzed NR/1427/15 9/159	Sex Race A vs. B: Age mean: 45 vs. 47 Female: 40% vs. 37% Non white: 0% vs. 4.1%	Proportion Treatment-Naïve A vs. B: Genotype 1: all HCV RNA>800,000: 87 vs. 93%
		C: 800-1400 mg/day based on body weight for 48 weeks						

Author, Year							
Country							
Study Name	Duration of				Histologic		Funding
Quality	Followup	Outcome	Subgroup Analyses	Subgroup Analyses	Response	Adverse Events	Source

Author, Year Country Study Name Ouality	Duration of	Outcome	Subgroup Analyses	Subaroun Andreas	Histologic	Adverse Events	Funding Source
Buti, 2010 ²⁶	Followup	Outcome A vs. B:	Subgroup Analyses NR	Subgroup Analyses NR	Response NR		
International	Followup visits at 24	A VS. B:	NK	NK	NK	(A vs. B vs. C [only A and B randomized]):	Schering-
International	weeks after	CVD. 27/96 (420/) via				Overall withdrawals: 8/86 (9.3%) vs. 17/73	Plough (now Merck)
Continued	completion of	SVR: 37/86 (43%) vs. 35/73 (47.9%); p=NS				(23.3%) vs. 100/816 (12.3%); A vs. B p=NS Withdrawals for adverse events: 3/86 (3.5%) vs.	Merck)
Continued	treatment	33/73 (47.9%), p=N3				6/73 (8.2%) vs. 39/816 (5.0%); A vs. B p=NS	
	treatment					Serious adverse events: 6/86 (7.0%) vs. 6/73	
						(8.2%) vs. 57/816 (7.0%); A vs. B p=NS	
						(6.2%) vs. 57/610 (7.0%), A vs. B p-1vs	
						Influenza-like illness: 36/86 (41.9%) vs. 34/73	
						(46.6%) vs. 347/816 (42.5%); A vs. B p=NS	
						Fatigue: 24/86 (27.9%) vs. 18/73 (24.7%) vs.	
						202/816 (24.8%); A vs. B p=NS	
						Myalgia: 22/86 (25.6%) vs. 12/73 (16.4%) vs.	
						162/816 (19.9%); A vs. B p=NS	
						Pyrexia: 21/86 (24.4%) vs. 18/73 (24.7)% vs.	
						245/816 (30%); A vs. B p=NS	
						Pruritus: 20/86 (23.3%) vs. 12/73 (16.4%) vs.	
						176/816 (21.6%); A vs. B p=NS	
						Neutropenia: 18/86 (20.9%) vs. 16/73 (21.9%) vs.	
						175/816 (21.4%); A vs. B p=NS	
						Nausea: 18/86 (20.9%) vs. 15/73 (20.5%) vs.	
						159/816 (19.5%); A vs. B p=NS	

Author, Year Country Study Name			Protease Inhibitor			Number Screened/ Eligible/ Enrolled/	Age Sex	Genotype Severity of Liver Disease Proportion
Quality 2000 ²⁷	Interferon Regimen	Ribavirin Regimen	0	Eligibility	Exclusion	Analyzed	Race	Treatment-Naïve
Dalgard, 2008 ²⁷	All patients were	All patients were	None	Treatment naïve	Injection drug use or alcohol abuse in	NR/428/298/	(A vs. B	(A vs. B vs. C)
Denmark,	treated for 4 weeks.	treated for 4 weeks.		HCV RNA positive	the prior 6 months	298	vs. C)	C 4 2/2 11
Sweden, Norway	Subjects with rapid	Subjects with rapid		HCV genotype 2 or 3	Poorly controlled psychiatric illnesses		A	Genotype 2/3: all
Da anda d	virologic response after 4 weeks were	virologic response after 4 weeks were		Elevated ALT at least	Decompensated cirrhosis		Age median:	Proportion treatment- naïve: all
Pegylated interferon alpha	randomized to A. or B.	randomized to A or		once during the prior 6 months	HBV positive HIV positive		38 vs. 38	Fibrosis: NR
and ribavirin for	Subjects without rapid	B. Subjects without		months	Liver disease of other etiologies		vs. 43	HCV RNA >400,000:
12 vs. 24 weeks	virologic response were	rapid virologic			Liver disease of other enologies		V3. 43	64% vs. 58% vs. 75%
in patients with	allocated to group C.	response were					Female:	Cirrhosis: NR
hepatitis C virus		allocated to group					36% vs.	
genotype 2 or 3	A: Pegylated interferon	C.					35% vs.	
and rapid	alfa-2b 1.5						41%	
virological	mcg/kg/week for 14	A: 800-1400						
response	weeks	mg/day based on					Non	
	B: Pegylated interferon	body weight for 14					white: NR	
Overall Quality:	alfa-2b 1.5	weeks						
Fair	mcg/kg/week for 24	B: 800-1400 mg/day						
	weeks	based on body						
	C: Pegylated interferon	weight for 24 weeks						
	alfa-2b 1.5	C: 800-1400 mg/day						
	mcg/kg/week for 24	based on body						
	weeks	weight for 24 weeks						

Author, Year Country							
Study Name Quality	Duration of Followup	Outcome	Subgroup Analyses	Subgroup Analyses	Histologic Response	Adverse Events	Funding Source
Dalgard, 2008 ²⁷ Denmark, Sweden, Norway Continued	Up to 24 weeks after treatment completion (week 48)	(A vs. B vs. C): ETR: 136/148 (91.9%) vs. 144/150 (96.0%) vs. NR; A vs. B p=NS SVR: 120/148 (81.1%) vs. 136/150 (90.7%) vs. 69/126 (58.5%); A vs. B p=NS	` / · I	A vs. B: HCV RNA >400K IU/ml: 77/88 (87%) vs. 75/85 (88%); p=NS HCV RNA <400K IU/ml: 35/42 (83%) vs. 55/55 (100%); p=NS Genotype 3: 93/110 (84%) vs. 106/115 (92%); p=NS Genotype 2: 27/29 (93%) vs. 30/31 (97%); p=NS	NR	A vs. B: Treatment discontinuations (<80% of prescribed injections): 9/148 (6%) vs. 32/150 (21%); p=0.02 Hemoglobin <10g/dL: 9/148 (6.1%) vs. 13/150 (8.7%); p=0.39 Neutrophils <700/mm3: 9/148 (6.1%) vs. 15/149 (10.1%); p=0.31 Depression: 29/110 (26.4%) vs. 37/124 (29.8%); p=0.56	Schering- Plough (now Merck)

Author, Year Country Study Name Quality	Interferon Regimen	Ribavirin Regimen	Protease Inhibitor Regimen	Eligibility	Exclusion	Number Screened/ Eligible/ Enrolled/ Analyzed	Age Sex Race	Genotype Severity of Liver Disease Proportion Treatment-Naïve
2010 ²⁸ Austria Pegylated	treated for 4 weeks. Subjects with rapid virologic response (HCV-RNA <50	mg/day depending of body weight for 48 weeks B: 1000-1200		Ages 18-65 years Chronic HCV genotype 1 or 4 infection Positive HCV antibody	Evidence of decompensation Co-infection with HBV or HIV Systematic immunomodulatory or antineoplastic therapy within previous 6	289	Age mean: 45 vs. 44	Genotype 1: 91% vs. 89% Treatment-naïve: all
interferon alfa- 2a and ribavirin for 24 weeks in	IU/mL) were treated with 24 weeks. Subjects without rapid virologic	mg/day depending of body weight for 72 weeks		test Quantifiable HCV RNA Elevated ALT	months Diabetes mellitus treated with insulin Severe psychiatric disorders		Female 36% vs. 35%	HCV-RNA level >800K IU: 38% vs. 44%
hepatitis C type 1 and 4 patients with rapid virologic	response continued to week 12 and were re- evaluated. Subjects with early virologic response			Histologic findings consistent with chronic hepatitis C on liver biopsy within the	History of immunologically mediated disease Other severe chronic or uncontrolled disease		non White: NR	Fibrosis stage 3-4: 20% vs. 19%
response Overall Quality: Poor	(HCV RNA <600 IU/mL or a 2 log decrease in serum HCV RNA) were randomized			previous 6 months Neutrophils >3000 Platelets >100K Hemoglobin > 12 g/dL in				
	to complete either 48 weeks or 72 weeks of treatment.			women and > 13 g/dL in men Serum creatinine <1.5 times the upper limit of				
	A: Pegylated interferon alfa-2a 180 mcg/week for 48 weeks B: Pegylated interferon			normal Thyroid-stimulating hormone within normal limits				
	alfa-2a 180 mcg/week for 72 weeks			mines				

Author, Year Country Study Name Quality	Duration of Followup	Outcome	Subgroup Analyses	Subgroup Analyses	Histologic Response	Adverse Events	Funding Source
Ferenci, 2010 ²⁸	Followup	A vs. B:	NR	A vs. B:	NR	A vs. B:	Roche
Austria	visits at 24					Overall withdrawals: 26/139 (18.7%) vs. 48 / 150	
	weeks after	SVR: 71 / 139 (51.1%)		Genotype 1: 65/127		(32.0%); p<0.01	
Continued	completion of	vs. 88 / 150 (58.7%);		(51.2%) vs. 81/134		Withdrawals for adverse events: 7/139 (5.07%) vs.	
	treatment	p=NS		(60.4%); p=NS		8/150 (5.3%); p=NS	
				Genotype 4: 6/12		Serious adverse events: 38 / 139 (27.3%) vs. 51 /	
				(50.0%) vs. 7/16		150 (34.0%); p=NS	
				(43.8%); p=NS		Deaths: NR	
				Baseline HCV-RNA		Serious hematologic adverse event: 1/139	
				>400K IU/mL:		(0.007%) vs. 2 / 150 (1.3%); p=NS	
				51/105 (48.6%) vs.		Serious gastrointestinal adverse event: 5/139	
				64/113 (56.6%);		(3.6%) vs. 2/150 (1.3%); p=NS	
				p=NS		Serious infectious adverse event: 2/139 (1.4%) vs.	
				Baseline HCV-		8/150 (5.3%); p=NS	
				RNA<400K IU/mL:		Serious pulmonary adverse event; 3/139 (2.2%)	
				20/34 (58.8%) vs.		vs. 5/150 (3.3%); p=NS	
				24/37 (64.9%); p=NS		Serious neuropsychiatric adverse event:	
						5/139(3.6%) vs. 4/150 (2.7%); p=NS	
				Fibrosis F3-4: 18/32		Serious cardiovascular adverse event: 3/139	
				(56.3) vs. 19/34		(2.2%) vs. 3/150 (2.0%); p=NS	
				(55.9%); p=NS		Serious skin adverse event: 1/139 (0.007%) vs.	
				Fibrosis F0-2: 53/107		1/150 (1.3%); p=NS	
				(49.5%) vs. 69/116			
				(59.5%); p=NS			

						Number		Genotype
Author, Year						Screened/		Severity of Liver
Country			Protease			Eligible/	Age	Disease
Study Name			Inhibitor			Enrolled/	Sex	Proportion
Quality	Interferon Regimen	Ribavirin Regimen	Regimen	Eligibility	Exclusion	Analyzed	Race	Treatment-Naïve
Hadziyannis,	A: Pegylated interferon	A: ("Low dose -24"	None	Treatment naive adults	Neutropenia (neutrophil count <1.5	1736/1373/1	(A vs. B	(A vs. B vs. C vs. D)
2004 ²⁹	alpha-2a 180 μg/week	or "24-LD")		with serum hepatitis C	x109 cells/L)	311/1284	vs. C vs.	Genotype, n (%):
Europe, North &	for 24 weeks	Ribavirin 800		virus (Genotype) RNA	Thrombocytopenia (platelet count		D):	Genotype 1 -
South America,		mg/day for 24		concentration greater	<90x109 cells/L)		Age	101/207(49%) vs.
Australia, New	B: Pegylated interferon	weeks		than 2000 copies/mL	Anemia (hemoglobin level <120 g/L in		(mean):	118/280(42%) vs.
Zealand, and	alpha-2a 180 μg/week			Elevated serum alanine	women and <130 g/L in men) - or a		41 vs. 42	250/361(69%) vs.
Taiwan (99	for 24 weeks	B: ("Standard dose-		aminotransferase(ALT)	medical condition that would be		vs. 43 vs.	271/436(62%)
centers world-		24" or "24-SD")		level documented on 2 or	clinically significantly worsened by		43 years	Genotype 2 -
wide)	C: Pegylated interferon	Ribavirin 1000		more occasions 14 days	anemia			39/207(19%) vs.
	alpha-2a 180 μg/week	mg/day for 24		or more apart within the	Serum creatinine level more than 1.5		Female:	53/280(19%) vs.
Peginterferon-	for 48 weeks	weeks, (Body		previous 6 months	times the upper limit of normal		32% vs.	46/361(13%) vs.
α2a and		weight <75kg)		Compensated liver	Co-infection with hepatitis A or B virus		34% vs.	66/436(15%)
Ribavirin	D: Pegylated interferon	or		disease and a liver	or HIV		27% vs.	Genotype 3 -
Combination	alpha-2a 180 μg/week	Ribavirin 1200		biopsy specimen	History of bleeding from esophageal		34%	57/207(28%) vs.
Therapy in	for 48 weeks	mg/day for 24		consistent with chronic	varices or other conditions consistent			91/280(33%) vs.
Chronic		weeks, (Body		hepatitis C obtained in	with Decompensated liver disease		Race:	53/361(15%) vs.
Hepatitis C		weight >75kg)		the previous 15 months	Organ transplant		White -	87/436(20%)
				Patients with	Severe or poorly controlled psychiatric		88% vs.	Other - 106/207(51%)
Overall Quality:		C: ("Low dose-48"		compensated cirrhosis or	disease (especially depression)		91% vs.	vs. 162/280(58%) vs.
Fair		or "48-LD")		transition to cirrhosis	malignant neoplastic disease		87% vs.	111/361(31%) vs.
		Ribavirin 800		(Child–Pugh class A)	Severe cardiac or chronic pulmonary		90%	165/436(38%)
		mg/day for 48		Negative pregnancy test	disease		Non	Histologic diagnosis
		weeks		result 24 hours before the	Immunologically mediated disease		White -	using Ishak scores:
				first dose of study	(except controlled thyroid disease)		12% vs.	Non cirrhotic -
		D: ("Standard dose-		medications	Seizure disorder		9% vs.	163/207(79%) vs.
		48" or "48-SD")			Severe retinopathy		13% vs.	209/280(75%) vs.
		Ribavirin 1000			Alcohol or drug dependence within 1		10%	270/361(75%) vs.
		mg/day for 48			year of study entry			321/436(74%)
		weeks, (Body			Clinically significant co morbid medical			Cirrhosis -
		weight <75kg)			conditions			10/207(5%) vs.
		or			Pregnancy or unwillingness to practice			20/280(7%) vs.
		Ribavirin 1200			contraception			25/361(7%) vs.
		mg/day for 48						35/436(8%)
		weeks, (Body						
		weight >75kg)						

Author, Year Country Study Name Quality	Interferon Regimen	Ribavirin Regimen	Protease Inhibitor Regimen	Eligibility	Exclusion	Number Screened/ Eligible/ Enrolled/ Analyzed	Age	Genotype Severity of Liver Disease Proportion Treatment-Naïve
Hadziyannis, 2004 ²⁹ Europe, North & South America, Australia, New Zealand, and Taiwan (99 centers world- wide)	6	8	9		Severe psychiatric disease was defined as treatment with an antidepressant medication or major tranquilizer for major depression or psychosis - for 3+months /or period of disability due to psychiatric disease History of a suicide attempt/hospitalization			Bridging fibrosis - 34/207(16%) vs. 51/280(18%) vs. 66/361(18%) vs. 80/436(18%) 100% Treatment naive
Continued								

Author, Year	1						
Country							
Study Name	Duration of				Histologic		
Ouality	Followup	Outcome	Subgroup Analyses	Subgroup Analyses	Response	Adverse Events	Funding Source
Hadziyannis, 2004 ²⁹	Followup visits	(A vs. B vs. C vs. D):	NR	(A vs. B vs. C vs. D):	NR	(A vs. B vs. C vs. D):	Roche, Basel,
Europe, North & South	at 24 weeks	(A vs. B vs. C vs. B).	INK	(A vs. B vs. C vs. B).	TVIC	Pre-mature withdrawal:	Switzerland
America, Australia, New	post-treatment	SVR: 112/207 (54%) vs.		SVR:		(for any reason): 14/207(7%) vs. 22/280(8%) vs.	Switzeriand
Zealand, and Taiwan (99	post treatment	177/280 (63%) vs. 180/361		Genotype 1 -		117/361(32%) vs. 117/436(27%)	
centers world-wide)		(50%) vs. 259/436 (59%)		29/101(29%) vs.		(for AE/abnormal labs): 10/207(5%) vs. 13/280(5%) vs.	
		A vs. C p=NS		42/118(36%) vs.		59/361(16%) vs. 67/436(15%)	
Peginterferon-α2a and		A vs. B p=0.04		41/250(16%) vs.		(insufficient response): 0/207(0%) 0/280(0%) vs.	
Ribavirin Combination		B vs. D p<0.0001		52/271(19%)		31/361(9%) vs.24/436(6%)	
Therapy in Chronic		C vs. D p=0.007		Genotype 2/3 -		Deaths: vs. 0/207(0%) 0/280(0%) vs. 1/361(<1%) vs.	
Hepatitis C				79/96(82%) vs.		2/436(<1%)	
				117/144(81%) vs.		Severe Adverse Events: 46/207(22%) vs. 63/280(23%)	
Overall Quality: Fair				77/99(78%) vs.		vs. 116/361(32%) vs. 114/436(32%)	
				113/153(74%)			
						Adverse events:	
				Bridging fibrosis or		Headache - 102/207(49%) vs. 136/280(49%) vs.	
				cirrhosis: 21/43 (49%)		187/361(52%) vs. 239/436(55%)	
				vs. 36/66 (55%) vs.		Fatigue - 98/207(47%) vs. 135/280(48%) vs.	
				33/87 (38%) vs. 56/111		182/361(50%) vs. 211/436(48%)	
				(50%)		Myalgia - 91/207(44%) vs. 120/280(43%) vs.	
				No Bridging fibrosis or cirrhosis: 89/154 (58%)		154/361(43%) vs. 163/436(37%)	
				vs. 130/196 (66%) vs.		Pyrexia - 81/207(39%) vs. 114/280(41%) vs. 156/361(43%) vs. 173/436(40%)	
				146/262 (56%) vs.		Insomnia - 69/207(33%) vs. 99/280(35%) vs.	
				210/313 (67%)		146/361(40%) vs. 146/436(33%)	
				210/313 (07/0)		Nausea - 64/207(31%) vs. 91/280(33%) vs.	
				HCV RNA>200:		107/361(30%) vs. 151/436(35%)	
				63/117 (54%) vs.		Rigors - 64/207(31%) vs. 87/280(31%) vs. 87/361(24%)	
				93/148 (63%) vs.		vs. 119/436(27%)	
				116/260 (45%) vs.		Irritability - 59/207(29%) vs. 76/280(27%) vs.	
				163/294 (55%)		96/361(27%) vs. 112/436(26%)	
				HCV RNA<200: 49/90		Alopecia - 53/207(26%) vs. 74/280(265) vs.	
				(54%) vs. 84/132 (64%)		106/361(29%) vs. 92/436(21%)	
				vs. 64/101 (63%) vs.		Arthralgia - 50/207(24%) vs. 70/280(25%) vs.	
				96/142 (68%)		106/361(29%) vs. 105/436(24%)	
						Pruritus - 56/207(27%) vs. 60/280(21%) vs.	
						81/361(22%) vs. 111/436(25%)	
						Depression - 43/207(21%) vs. 42/280(15%) vs.	
						79/361(22%) vs. 104/436(24%)	
						Diarrhea - 44/207(21%) vs. 46/280(16%) vs.	
						65/361(18%) vs. 96/436(22%)	
						Dermatitis - 34/207(16%) vs. 49/280(185) vs.	
						69/361(19) vs. 86/436(20%) Decreased appetite - 30/207(14%) vs. 41/280(15%) vs.	
						66/361(18%) vs. 91/436(21%)	

Author, Year Country Study Name		Du D .	Protease Inhibitor	TH. 11-114		Number Screened/ Eligible/ Enrolled/	Age Sex	Genotype Severity of Liver Disease Proportion
				. 8				
Quality Ide, 2009 Japan ³⁰ A Randomized Study of Extended Treatment with Pegylated interferon alpha- 2b Plus Ribavirin Based on Time to HCV RNA Negative-	Interferon Regimen A: (Standard group - received a 48-week course of treatment) Pegylated interferon α-2b - 1.5 μg/kg/week for 48 weeks B: (Extended group – treatment course performed for 44 weeks after HCV RNA first became negative) Pegylated interferon α-2b - 1.5 μg/kg/week for 48-68 weeks	A: (Standard group - received a 48-week course of treatment) Ribavirin by body weight: < 60 kg - 600 mg/day for 48 weeks 60-80 kg - 800 mg/day for 48 weeks > 80 kg - 1000 mg/day for 48 weeks B: (Extended group - treatment course performed for 44 weeks after HCV RNA first became negative) Ribavirin by body weight: < 60 kg - 600 mg/day for 48-68 weeks 60-80 kg - 800		Eligibility Male and female patients aged 20–75 years Compensated chronic HCV genotype 1b infection Positive for HCV RNA by a quantitative reverse-transcription PCR with a concentration >100K IU / ml At least one elevated serum alanine aminotransferase level at the time of screening or entry into the trial	Patients with an HCV genotype other than 1b infection Hepatitis B surface antigen Autoimmune hepatitis Primary biliary cirrhosis Sclerosing cholangitis Decompensated cirrhosis (Child – Pugh class B or C) Evidence of hepatocellular carcinoma Patients with platelet counts of < 8 × 10 4/mm3, leukocyte counts of 2,500/ml or less, or hemoglobin levels of < 12 g/dl	Analyzed NR/NR/113/ 113	Race A vs. B: Age (Mean): 55.3 vs. 54.6 years Female: 53.6% vs. 47.4% Non white: NR	Treatment-Naïve A vs. B: Genotype 1b: 100% Fibrosis Stage (Desmet et al 1994): 1/2 - 67.8% vs. 52.6% 3/4 - 19.6% vs. 19.3% Treatment naïve: NR
		mg/day for 48-68 weeks > 80 kg - 1000 mg/day for 48-68 weeks						

Author, Year Country Study Name Quality	Duration of Followup	Outcome	Subgroup Analyses	Subgroup Analyses	Histologic Response	Adverse Events	Funding Source
Ide, 2009 ³⁰	Followup	A vs. B:	NR	NR	NR	A vs. B:	Internal
Japan	visits at 24					Overall withdrawals: 11/56 (20%) vs. 9/57 (16%);	Funding
	weeks after	SVR: 20/56(36%) vs.				p=NS	
Continued	completion of	30/57(53%), p=0.07				Withdrawal due to adverse event: 7/56 (13%) vs.	
	treatment					6/57 (11%); p=NS	

Author, Year Country Study Name Quality	Interferon Regimen	Ribavirin Regimen	Protease Inhibitor Regimen	Eligibility	Exclusion	Number Screened/ Eligible/ Enrolled/ Analyzed	Age Sex Race	Genotype Severity of Liver Disease Proportion Treatment-Naïve
Kamal, 2005 ³¹	A: Pegylated interferon	A: Ribavirin 10.6	None	Documented chronic	Previous IFN-a therapy	335/287/279	(A vs. B	(A vs. B vs. C):
Egypt	alfa-2b 1.5 µg/kg for 24	mg/kg/day for 24		hepatitis C according to	Other liver diseases such as hepatitis A,	/271	vs. C):	, ,
C5 1	weeks	weeks		the following criteria:	hepatitis B, schistosomiasis,		ĺ	Genotype 4: 100%
Pegylated				elevated serum alanine	autoimmune hepatitis, alcoholic liver		Age	
interferon alpha-	B: Pegylated interferon	B: Ribavirin 10.6		aminotransferase (ALT)	disease, drug induced hepatitis, or		(Mean):42	(Ishak et al 1995)
2b and ribavirin	alfa-2b 1.5 µg/kg for 36	mg/kg/day for 36		above the upper limit of	decompensated liver disease		vs. 44 vs.	Inflammation grade
therapy in	weeks	weeks		normal (40 U/l) on two	Co infection with schistosomiasis or		41	(mean): 8.2 vs. 7.6 vs.
chronic hepatitis				occasions during the	human immunodeficiency virus			9.1
C genotype 4:	C: Pegylated interferon	C: Ribavirin 10.6		preceding six months	Neutropenia (1,500/mm3)			Fibrosis stage (mean):
impact of	alfa-2b 1.5 µg/kg for 48	mg/kg/day for 48		Anti-HCV positive	Thrombocytopenia (90,000/mm3)			1.8 vs. 2.3 vs. 2.1
treatment	weeks	weeks		antibody status assessed	Creatinine concentration >1.5 x the		vs. 48%	HCVRNA mean: 2.8
duration and				by second generation	upper limit of normal			vs. 2.7 vs. 2.8
viral kinetics on				enzyme linked	Serum a fetoprotein concentration >25		Non	
sustained				immunosorbent assay	ng/ml		white: NR	Treatment naïve:
virological				Positive polymerase	Organ transplant			100%
response				chain reaction for HCV	Neoplastic disease			
				RNA	Severe cardiac or pulmonary disease			
Overall Quality:				Genotype 4	Unstable thyroid dysfunction			
Fair				Chronic hepatitis C in	Psychiatric disorder			
				liver biopsy performed	Current pregnancy or breast feeding			
				within the preceding year	Therapy with immunomodulatory			
				with no signs of cirrhosis	agents within the last six months			
				or				
				bridging fibrosis on				
				pretreatment liver biopsy				

Author, Year Country Study Name Quality	Duration of Followup	Outcome	Subgroup Analyses	Subgroup Analyses	Histologic Response	Adverse Events	Funding Source
Kamal, 2005 ³¹	Followup	(A vs. B vs. C):	NR	NR	(A vs. B vs.	(A vs. B vs. C):	Fulbright
Egypt	visits at 48				C):	Deaths: NR	Foundation
	weeks after	ETR: 45/95(48%) vs.				Life-threatening Adverse Events: NR	Grants(NIAID
Continued	completion of	65/96(68%) vs.			All patients	Severe Adverse Events: NR	(R2)
	treatment	67/96(70%),			underwent	Overall Treatment Withdrawals: 3/95 (3%) vs.	AI054887) &
		p=0.04 (A and B);			liver biopsy	5/96 (5%) vs. 5/96 (5%)	the Alexander
		p=0.02 (A and C),			before and	Withdrawals due to Adverse Events: 1(2%) vs.	von Humboldt
		p=0.4(B and C)			after	2(2%) vs. 4(4%)	Foundation
1		GLID 20/07 (200/)			treatment.	Neutropenia (<500/mm3) 1/95 (1%) vs. 1/96 (1%)	(Germany)
		SVR: 28/95 (29%) vs.			Pair wise	vs. 3/96 (3%)	
		63/96 (66%) vs. 66/96 (69%),			comparison of	Fatigue- 56/95(60%) vs. 59/96(64%) vs. 62/96(66%)	
		p=0.001 (A and B);			histological	Influenza-like illness- 53 (57%) vs. 58/96(63%)	
		p=0.001 (A and B); p=0.001(A and C);			grading and	vs. 59/96(63%)	
		p=0.5(B and C)			staging	Headache- 49/95(53%) vs. 52/96(57%) vs.	
		p=0.5(B and C)			scores for	58/96(62%)	
					the initial	Myalgia- 48/95(52%) vs. 52/96(57%) vs.	
					and	58/96(62%)	
					followup	Pyrexia- 41/95(44%) vs. 50/96(54%) vs.	
					biopsies	53/96(62%)	
					showed no	Insomnia- 31/95(33%) vs. 35/96(38%) vs.	
					deterioration	46/96(49%)	
					or	Injection site erythema - 28/95(30%) vs.	
						34/96(37%) vs. 39/96(42%)	
						Irritability- 26/95(28%) vs. 33/96(36%) vs.	
					any patient	30/96(32%)	
					and	Back pain- 23/95(25%) vs. 25/96(27%) vs.	
					improvemen	` '	
					t (>2 point	Rigors- 16/95(17%) vs. 17/96(18%) vs.	
					necro-	21/96(22%) Somethmost 12/05(140%) via 16/06(170%) via	
					y score	Sore throat- 13/95(14%) vs. 16/96(17%) vs. 20/96(21%)	
					improvemen	` '	
					t) was	20/96(21%)	
					detected in	Pruritus- 10/95(11%) vs. 15/96(16%) vs.	
					155 patients	18/96(19%)	
					(54%):	Anorexia- 9/95(10%) vs. 14/96(15%) vs.	
I					(= ./0/.	18/96(19%)	
						10/90(19%)	

Author, Year Country Study Name Quality	Duration of Followup	Outcome	Subgroup Analyses	Subgroup Analyses	Histologic Response	Adverse Events	Funding Source
Kamal, 2005 ³¹					Histological	Arthralgia- 8/95(9%) vs. 12/96(13%) vs.	
Egypt					response	17/96(18%)	
					was more	Dyspnea- 8/95(9%) vs. 11/96(12%) vs.	
Continued					likely in	15/96(16%)	
					those who received	Rash- 7/95(8%) vs. 10/96(11%) vs. 12/96(13%)	
					longer	Depression- 3/95(3%) vs. 3/96(3%) vs. 9/96(9%) Dry mouth- 5/95(5%) vs. 7/96(8%) vs. 8/96(9%)	
					treatment	Alopecia- 4/95(4%) vs. 6/96(7%) vs. 7/96(7%)	
						Nausea- 4/95(4%) vs. 4/96(4%) vs. 7/96(7%)	
						Dizziness- 3/95(3%) vs. 5/96(5%) vs. 6/96(6%)	
					_	Abdominal pain- 3/95(3%) vs. 5/96(5%) vs.	
					t was	7/96(7%)	
					detected in:	Dry skin- 2/95(2%) vs. 6/96(7%) vs. 7/96(7%)	
					(>2 point	Diarrhea- 2/95(2%) vs. 6/96(7%) vs. 8/96(9%)	
					necro-	Vomiting- 1/95(2%) vs. 3/96(3%) vs. 5/96(5%)	
					inflammator		
					y score		
					improvemen t):		
					12/95(12.6%		
) vs.		
					67/96(69.8%		
) vs. 71/96		
					(73.9%)		

Author, Year Country Study Name Quality	Interferon Regimen	Ribavirin Regimen	Protease Inhibitor Regimen	Eligibility	Exclusion	Number Screened/ Eligible/ Enrolled/ Analyzed	Age Sex Race	Genotype Severity of Liver Disease Proportion Treatment-Naïve
Lagging,	A: Pegylated interferon	A: Ribavirin 800	None	Adults age 18 years and	NR	392/382/382	A vs. B:	A vs. B:
2008^{32}	alfa-2a 180 µg/week for	mg/day (2 equal		older		/382		
Denmark &	12 weeks	doses) for 12 weeks		Compensated liver			Age	Genotype 2: 28% vs.
Finland				disease			(Mean):	26%
	B: Pegylated interferon	B: Ribavirin 800		Treatment-naive for			42 vs. 42	Genotype 3: 71% vs.
Randomized	alfa-2a 180 µg/week for	mg/day (2 equal		hepatitis C			years	74%
Comparison of	24 weeks	doses) for 24 weeks		Seronegative for				
12 or 24 Weeks				hepatitis B surface			Female:	Bridging fibrosis
of Pegylated				antigen and for			37% vs.	(Ishak stage 3-4):
interferon alpha-				antibodies to human			44%	39% vs. 40%
2a and Ribavirin				immunodeficiency virus				Cirrhosis (Ishak stage
in Chronic				Positive test for anti-			Non	5-6): 13% vs. 13%
Hepatitis C				HCV antibody			white: NR	Steatosis present
Virus Genotype				Infection with HCV				(grade 1-3): 64% vs.,
2/3 Infection				genotypes 2 and/or 3 but				69%
				not genotypes 1, 4, 5, or				Moderate or severe
Overall Quality:				6				steatosis (grade 2-3):
Fair				HCV-RNA 600 IU/mL				29% vs. 27%
				within 6months of				
				treatment initiation				Treatment naive:
				Liver biopsy consistent				100%
				with chronic hepatitis C				
				within 24 months of				
				entry				

	Duration of Followup	Outcome	Subgroup Analyses	Subgroup Analyses	Histologic Response	Adverse Events	Funding Source
Denmark & Finland Continued	Followup visits at 24 weeks after completion of treatment	A vs. B: SVR: 114/194 (59%) vs. 147/188 (78%); p<0.0001	A vs. B: SVR: Age <40: 61/76 (80%) vs. 63/76 (83%);p=NS Age >40: 53/118 (45%0 vs. 84/112 (84%); p<0.0001	A vs. B: No significant fibrosis - 59/85(69%) vs. 69/83(84%); p=0.022 Bridging fibrosis - 36/70(51%) vs. 53/70(76%); p=0.0051 Cirrhosis - 19/23(84%) vs. 13/23(57%); p=NS Genotype 2: 31/55 (56%) vs. 40/49 (82%); p=0.0057 Genotype 3: 79/137 (58%) vs. 108/139 (78%); p=0.0015	NR	A vs. B: Deaths: NR Life-threatening Adverse Events: NR Severe Adverse Events: NR Withdrawals: 12/194 (6%) vs. 46/188 (24%); p<0.001 Withdrawals due to adverse events: 2/194(1%) vs. 20/188 (11%); P=0.0001	Swedish Society of Medicine, Swedish Medical Council, Swedish Society of Microbiology, Avtal om lakarutbildning och forskning (ALF) Funds, and Roche affiliates (Nordic region)

Author, Year Country Study Name Quality	Duration of Followup	Outcome	Subgroup Analyses	Subgroup Analyses	Histologic Response	Adverse Events	Funding Source
Lagging, 2008 ³² Denmark & Finland Continued	Followup visits at 24 weeks after completion of	A vs. B: SVR: 114/194 (59%) vs. 147/188 (78%);	A vs. B: SVR: Age <40: 61/76 (80%)	A vs. B: No significant fibrosis - 59/85(69%)	NR	A vs. B: Deaths: NR Life-threatening Adverse Events: NR Severe Adverse Events: NR	Swedish Society of Medicine, Swedish
Continued	treatment	p<0.0001	vs. 63/76 (83%);p=NS Age >40: 53/118 (45%0 vs. 84/112 (84%); p<0.0001	vs. 69/83(84%);		Withdrawals: 12/194 (6%) vs. 46/188 (24%); p<0.001 Withdrawals due to adverse events: 2/194(1%) vs. 20/188 (11%); P=0.0001	Medical Council, Swedish Society of Microbiology, Avtal om lakarutbildning och forskning (ALF) Funds, and Roche affiliates (Nordic region)

Author, Year Country			Protease			Number Screened/ Eligible/	Age	Genotype Severity of Liver Disease
Study Name			Inhibitor			Enrolled/	Sex	Proportion
Quality	Interferon Regimen	Ribavirin Regimen	Regimen	Eligibility	Exclusion	Analyzed	Race	Treatment-Naïve
Liu, 2008 ³⁴ Taiwan Pegylated Interferon-alpha-	A: Pegylated interferon alfa-2a 180 μg/week for 24 weeks B: Pegylated interferon alfa-2a 180 μg/week for 48 weeks	Ribavirin Regimen A: (24-week group) Ribavirin by body weight: < 75 kg - 1000 mg/day for 24 weeks > 75 kg - 1200 mg/day for 24 weeks B: (48-week group) Ribavirin by body weight: < 75 kg - 1000 mg/day for 48 weeks > 75 kg - 1200 mg/day for 48 weeks	Regimen None	Eligibility Treatment-naive patients with chronic hepatitis C Aged >18 years Presence of anti-HCV antibody Detectable serum HCV RNA level determined by real-time RT-PCR analysis for 16 months HCV-1 infection confirmed by a reverse hybridization assay Serum alanine aminotransferase (ALT) level > upper limit of normal Liver histologic characteristics consistent with chronic viral hepatitis within the previous 3 months	Exclusion Anemia (hemoglobin level,<13 g/dL for men and <12 g/dL for women) Neutropenia (neutrophil count, <1500 cells/mm3) Thrombocytopenia (platelet count, <70,000 cells/mm3) Mixed infection with HCV-1 and another genotype of HCV Co infection with hepatitis B virus or HIV Chronic alcohol abuse (daily alcohol consumption, 120 g/day) Decompensated cirrhosis (Child-Pugh class B or C) Serum creatinine level 11.5x the upper limit of normal Autoimmune liver disease Neoplastic disease Organ transplantation or immunosuppressive therapy Evidence of drug abuse Pregnancy Poorly controlled autoimmune disease Cardiopulmonary disease Neuropsychiatric disorders		Race A vs. B: Age (Mean): 54 vs. 53 years Female: 42.9% vs. 43.5% Non white: NR	Treatment-Naïve A vs. B: Genotype 1a: 2.6% vs. 1.9% Genotype 1b: 92.9% vs. 94.2% Genotype 1a & 1b: 4.5% vs. 3.9% Fibrosis (Ishak 1995) > 3: 78.6% vs. 76.0% 6: 22.7% vs. 20.1% Steatosis- present: 44.2% vs. 41.6% absent: 55.8% vs. 58.4% Treatment naïve: 100%
					Diabetes mellitus with retinopathy Unwillingness to receive contraception during the study period			

Author, Year							
Country							
Study Name	Duration of				Histologic		Funding
Quality	Followup	Outcome	Subgroup Analyses	Subgroup Analyses	Response	Adverse Events	Source
Liu, 2008 ³⁴	Followup	A vs. B:	NR	NR	Histological	A vs. B:	National
Taiwan	visits at 24				Response:	Overall withdrawals: 7/154 (5%) vs. 4/154 (3%);	Taiwan
	weeks after	SVR: 49/87(56%) vs.			42/71(59%)	p=NS	University
Continued	completion of	89/117(76%), P<.001			vs.	Withdrawal due to adverse events: 6/154(4%) vs.	Hospital,
	treatment				76/97(78%),	4/154 (3%) p=NS	National
					p=0.001	Dose reduction due to Adverse Events:	Science
						69/154(45%) 82/154(53%) p=NS	Council, and
					ALT	Deaths: 0/154(0%) vs. 1/154(<1%); p=NS	Department of
					normalizatio	Serious Adverse Event: 4/154(2%) vs.	Health,
					n:	11/154(7%); p=NS	Executive
					38/75(51%)		Yuan, Taiwan
					vs.	Adverse Events:	
					77/107(72%	Fever - 35/154(23%) vs. 33/154(21%); p=NS	
), p<0.001	Rigor - 19/154 (12%) vs. 13/154(8%); p=NS	
						Fatigue - 88/154 (57%) vs. 100/154(65%); p=NS	
						Headache - 28/154 (18%) vs. 35/154(23%); p=NS	
						Myalgia - 40/154(26%) vs. 36/154(23%); p=NS	
						Arthralgia - 8/154(5%) vs. 13/154(8%); p=NS	
						Insomnia - 61/154(40%) vs. 69/154(45%); p=NS	
						Irritability - 19/154(12%) vs. 22/154(14%); p=NS	
						Depression - 36/154(23%) vs. 26/154(17%);	
						p=NS	
						Anorexia - 63/154(41%) vs. 80/154(52%); p=NS	
						Constipation - 10/154(6%) vs. 15/154(10%);	
						p=NS	
						Diarrhea - 14/154(9%) vs. 18/154(12%); p=NS	
						Body weight loss - 29/154(19%) vs. 46/154(30%);	
						p=0.02	
						Hair loss/alopecia - 24/154(16%) vs.	
						36/154(23%); p=NS	

Author, Year Country Study Name	Duration of				Histologic		Funding
Quality	Followup	Outcome	Subgroup Analyses	Subgroup Analyses	Response	Adverse Events	Source
Liu, 2008 ³⁴						Aphthous ulcer - 22/154(14%) vs. 34/154(22%);	
Taiwan						p=NS	
						Cough - 28/154(18%) vs. 32/154(21%); p=NS	
Continued						Nasal congestion - 13/154(8%) vs. 17/154(11%);	
						p=NS	
						Tinnitus - 13/154(8%) vs. 20/154(13%); p=NS	
						Dermatitis - 44/154(29%) vs. 48/154(31%); p=NS	
						Injection reaction - 22/154(14%) vs. 29/154(19%);	
						p=NS	
						Anemia - 60/154(39%) vs. 68/154(44%); p=NS	
						Neutropenia - 34/154(22%) vs. 42/154(27%);	
						p=NS	
						Thrombocytopenia - 25/154(16%) vs.	
						23/154(15%); p=NS	

						Number		Genotype
Author, Year						Screened/		Severity of Liver
Country			Protease			Eligible/	Age	Disease
Study Name			Inhibitor			Enrolled/	Sex	Proportion
Quality	Interferon Regimen	Ribavirin Regimen	Regimen	Eligibility	Exclusion	Analyzed	Race	Treatment-Naïve
Mangia,	A: Pegylated interferon	A: (control standard	None	18 to 70 years of age	Leukocyte count < 3000/cubic	NR/NR/283/	A vs. B:	A vs. B:
2005 ³⁵	alfa-2b 1.0 µg/kg/week	duration group)		Presence of antibodies to	millimeter	283		
Italy	for 24 weeks (control	Ribavirin by body		HCV	Platelet count < 80,000/cubic millimeter		Age	Genotype 2: 76% vs.
	standard duration	weight:		Infection with genotype	Hemoglobin level <12 g/deciliter for		(Mean):	75%
Pegylated	group)	< 75 kg - 1000		2 or 3	women and <13 g/deciliter for men		49.7 vs.	Genotype 3: 24% vs.
interferon alfa-		mg/day for 24		Abnormal alanine	Infection with the human		46.6 years	25%
2b and Ribavirin	B: Pegylated interferon	weeks		aminotransferase levels	Immunodeficiency virus (HIV)			
for 12 vs. 24	alfa-2b 1.0 µg/kg/week	> 75 kg - 1200		Treatment naïve	Alcohol intake > 20 g daily		Female:	HCV-RNA (>800,00
	for 12 or 24 weeks	mg/day for 24			Presence of drug abuse		44% vs.	IU/mL): 66% vs. 64%
	depending on if HCV	weeks			Presence of Chronic disease		44%	Liver fibrosis
	RNA at week 4				Presence of Psychiatric disease			(Scheuer 1991):
Overall Quality:	(variable duration	B: (variable			Presence of Autoimmune disease		Non	stage $> 3 - 23\%$ vs.
Fair	group)	duration group)			Presence of Pregnancy and lactation		white: NR	16%
		Ribavirin by body						Steatosis:
		weight:						(moderate/severe) -
		< 75 kg - 1000						36% vs. 31%
		mg/day for 48						_
		weeks						Treatment naïve:
		> 75 kg - 1200						100%
		mg/day for 48						
		weeks						

Author, Year Country							
Study Name	Duration of				Histologic		Funding
Quality	Followup	Outcome	Subgroup Analyses	Subgroup Analyses	Response	Adverse Events	Source
Mangia, 2005 ³⁵	Followup	A vs. B:	NR	A vs. B:	NR	A vs. B:	Italian branch
Italy	visits at 24					Withdrawals: 4/70 (6%) vs. 5/213 (2.3%); p=NS	of Schering-
	weeks after	SVR: 53/70(76%) vs.		SVR:		Withdrawals due to adverse events: NR	Plough
Continued	completion of	164/213(77%)		Genotype 2: 40/53		Deaths: NR	
	treatment			(75%) vs. 131/160		Serious adverse events: NR	
				(82%); p=NS			
				Genotype 3: 13/17			
				(76%) vs. 33/53			
				(62%); p=NS			

Author, Year Country Study Name			Protease Inhibitor			Number Screened/ Eligible/ Enrolled/	Age Sex	Genotype Severity of Liver Disease Proportion
Quality	Interferon Regimen	Ribavirin Regimen	Regimen	Eligibility	Exclusion	Analyzed	Race	Treatment-Naïve
Manns 2011 ³⁶ International Reduced dose and duration of peginterferon alfa-2b and weight-based ribavirin in patients with genotype 2 and 3 chronic hepatitis C Overall Quality: Fair	A: Pegylated interferon alfa-2b - 1.5 lg/kg/ Wk for 24 weeks B: (reduced-dose treatment) Pegylated interferon alfa-2b - 1.0 lg/kg/wk for 24 weeks	A: Ribavirin (weight-based) - 800–1200 mg/day for 24 weeks: <65 kg – 800 mg/day 65–85 kg – 1000 mg/day >85 kg – 1200 mg/day B: Ribavirin (weight-based) - 800–1200 mg/day for 24 weeks: <65 kg – 800 mg/day 65–85 kg – 1000 mg/day 75–85 kg – 1000 mg/day 85 kg – 1200 mg/day C: Ribavirin (weight-based) - 800–1200 mg/day >85 kg – 1200 mg/day C: Ribavirin (weight-based) - 800–1200 mg/day 85 kg – 1000 mg/day 65–85 kg – 1000 mg/day 65–85 kg – 1000 mg/day >85 kg – 1200 mg/day >85 kg – 1200 mg/day	None	Patients who had CHC G2 or G3 infection and were treatment naive. All patients had detectable hepatitis C virus (HCV) RNA, abnormal alanine aminotransferase, and compensated liver disease, and were eligible for treatment according to current consensus guidelines [10,11]. Patients were required to have hemoglobin levels P11 g/dl (women) or P12 g/dl (men), platelet count P100,000 cells/mm3, neutrophil count P1500 cells/mm3, and thyroid stimulating hormone levels within normal limits	Patients with human immunodeficiency virus (HIV) or hepatitis B coinfection, creatinine clearance <50 ml/min, cause of liver disease other than CHC, evidence of advanced liver disease, preexisting psychiatric conditions or history of severe psychiatric disorder. Patients with a history of substance abuse were required to have remained abstinent for 6 months prior to study entry and patients receiving buprenorphine were required to have been stable for 6 months	NR/696/696/ 602	A vs. B vs. C: Age (Mean): 38.8 vs. 39.9 vs. 39.7 years Female: 39.6% vs. 34.8% vs. 35.1% Race: NR	A vs. B: Genotype 2: 16.5% vs. 21.9% vs. 21.1% Genotype 3: 83.5% vs. 78.1% vs. 78.9% HCV-RNA (>600,00 IU/mL): 51.7% vs. 53.6% vs. 53.9% (<600,00 IU/mL): 47.4% vs. 46% vs. 45.2% Treatment naïve: 100%

Author, Year Country Study Name Quality	Duration of Followup	Outcome	Subgroup Analyses	Subgroup Analyses	Histologic Response	Adverse Events	Funding Source
Manns 2011 ³⁶ International Continued	24 weeks after end of treatment	A vs. B vs. C: SVR (Hep-Net + International cohort): 153/230(66.5%) vs. 144/224(64.3%) vs. 129/228(56.6%), p=0.495	A vs. B vs. C: SVR: HCV RNA: ≥600,000 IU/ml – 77/109(70.6%) vs. 70/103(68%) vs. 59/103(57.3%) <600,000 IU/ml – 75/119(63%) vs. 74/120(61.7%) vs. 69/123(56.1%)	A vs. B vs. C: SVR: Genotype 2 – (Hep-Net cohort, n=84): 21/27(77.8% vs. 19/314(61.3%) vs. 14/26(53.8%) (International cohort n=51): 8/11(72.7%) vs. 12/18(66.7%) vs. 16/22(72.7%) Genotype 3 – (Hep-Net cohort, n=263): 47/89(52.8%) vs. 50/84(59.5%) vs. 41/90(45.6%) (International cohort n=284): 77/103(74.8%) vs. 63/91(69.2%) vs. 58/90(64.4%)	NR	A vs. B vs. C: Deaths - <1% vs. <1% vs. 0% AE leading to interruption, reduction, or increase 15.7% vs. 4.9% vs. 12.3% AE leading to discontinuation 1.3% vs. 1.3% vs. 2.2% Pyrexia-37.8% vs. 37.1% vs. 44.3% Fatigue-22.6% vs. 22.3% vs. 15.8% Headache-22.6% vs. 25.4% vs. 25.4% Alopecia-20.9% vs. 16.1% vs. 13.6% Asthenia-19.1% vs. 27.7% vs. 19.7% Myalgia-15.2% vs. 12.1% vs. 14.9% Influenza-like illness- 12.6% vs. 9.4% vs. 10.1% Pruritus-12.6% vs. 19.6% vs. 10.1% Weight-decrease-12.6% vs. 9.6% Nausea-11.7% vs. 11.6% vs. 14.0% Injection-site erythema-11.3% vs. 13.8% vs. 7.5% Depressed mood-11.3% vs. 7.1% vs. 8.3% Arthralgia-10.9% vs. 7.6% vs. 10.5% Anemia-10.0% vs. 4.9% vs. 10.5% Anemia-10.0% vs. 4.9% vs. 10.5% Diarrhea-9.6% vs. 12.1% vs. 7.0% Dry skin-5.7% vs. 11.2% vs. 6.6% Treatment-emergent SAE-6.1% vs. 4.9% vs. 3.1% Treatment-emergent-7.0% vs. 4.5% vs. 5.3%	Schering- Plough (now Merck)

						Number		Genotype
Author, Year						Screened/		Severity of Liver
Country			Protease			Eligible/	Age	Disease
Study Name			Inhibitor			Enrolled/	Sex	Proportion
Quality	Interferon Regimen	Ribavirin Regimen	Regimen	Eligibility	Exclusion	Analyzed	Race	Treatment-Naïve
Mecenate,	Patients with negative	Patients with	None	HCV-RNA positive	History of injected drugs or alcohol	NR/210/143/	(All	(All groups - not
2010^{37}	HCV RNA at week 4	negative HCV RNA		HCV genotype 2 or 3	abuse (>40 g ethanol/day) within the 6	143	groups -	broken down by arm)
Italy	randomized to either 12	at week 4		Elevated alanine	months prior to study entry		not broken	
	or 24 weeks of	randomized to either		aminotransferase (>40	Poorly controlled psychiatric illness		down by	Genotype 2: 55%
Short vs.	treatment	12 or 24 weeks		UI/L) at least 8 months	Decompensated cirrhosis		arm)	Genotype 3: 45%
standard				prior to study entry	Positive for human immunodeficiency			
treatment with	A1: Pegylated	A1: Ribavirin 800-		Histologically proven	antibody virus (HIV) or positive for		Age	Cirrhosis (Ishak stage
pegylated	interferon alpha-2a 180	1200 mg daily for		chronic HCV hepatitis	hepatitis		(Mean):	5-6):
interferon alfa-	μg/week for 12 weeks	12 weeks			B surface antigen (HBV)		43 years	10%
2A plus ribavirin					Pregnancy			Bridging fibrosis
in patients with	A2: Pegylated	A2: Ribavirin 800-			Lactation		Female:	(Ishak stage 3-4):
hepatitis C virus	interferon alpha-2a 180	1200 mg daily for			Impaired renal function		19%	19%
genotype 2 or 3:	μg/week for 24 weeks	24 weeks			Other concurrent medical conditions of			
the CLEO trial					the liver different from HCV infection		Non	Treatment naïve: NR
	B: Pegylated interferon	B: Ribavirin 800-					white: NR	
Overall Quality:	alpha-2a 180 μg/week	1200 mg daily for						
Fair	for 24 weeks	24 weeks						
	(nonrandomized arm of	(nonrandomized						
	patient without rapid	arm of patient						
	virologic response)	without rapid						
		virologic response)						

Author, Year Country Study Name Quality	Duration of Followup	Outcome	Subgroup Analyses	Subgroup Analyses	Histologic Response	Adverse Events	Funding Source
Mecenate, 2010 ³⁷ Italy Continued	Followup visits at 24 weeks after completion of treatment	(A1 vs. A2): SVR: 60/72(83%) vs. 53/71(75%) p=NS	NR	(A1 vs. A2): SVR: Genotype 2: 32/60(53%) vs. 31/53(50%); p=NS Genotype 3: 28/60(47%) vs. 22/53(42%); p=NS	NR	(A1 vs. A2): Withdrawals: 0/72 (0%) vs. 5/71 (7%) Discontinuation due to Adverse Events - 0/72(0%) vs. 5/71(7%) Deaths: NR Life-threatening Adverse Events: NR Serious Adverse Events: NR Adverse events: Anemia: 5/72(7%) vs. 6/71(8%); p=NS Neutropenia: 2/72(3%) vs. 1/71(1%); p=NS Depression: 2/72(3%) vs. 2/71(3%); p=NS Cutaneous rash: 0/72(0%) vs. 0/71(0%); p=NS Alopecia: 0/72(0%) vs. 1/71(1%); p=NS Fatigue: 2/72(3%) vs. 4/71(5%); p=NS	NR

Author, Year Country			Protease			Number Screened/ Eligible/	Age	Genotype Severity of Liver Disease
Study Name			Inhibitor			Enrolled/	Sex	Proportion
Quality	Interferon Regimen	Ribavirin Regimen	Regimen	Eligibility	Exclusion	Analyzed	Race	Treatment-Naïve
Pearlman,	A: (Standard) Pegylated		None	Chronic HCV genotype	HCV/human immunodeficiency virus	NR/112/101/	A vs. B:	A vs. B:
2007 ³⁸	interferon α-2b - 1.5	Ribavirin by body		1-infected patients	co infection	101		
Atlanta, GA -	μg/kg/week for 48	weight:		Baseline elevated serum	HCV genotype other than 1		Age	Genotype 1: 100%
USA	weeks	< 64 kg - 800		alanine aminotransferase	Decompensated cirrhosis		(Mean):	
		mg/day for 48		levels	Other causes of liver disease, including		56 vs. 54	Fibrosis (METAVIR)
Treatment	B: (Extended)	weeks		Detectable serum HCV-	co infection with hepatitis B		years	F3/F4 - 27% vs. 25%,
Extension of 72	Pegylated interferon α-	65 - 84 kg - 1000		RNA via nucleic acid	Creatinine clearance <50 mL/minute			p=0.86
Weeks of	2b - 1.5 μg/kg/week for	mg/day for 48		testing	(modification of diet in renal disease		Female:	
Pegylated	72 weeks	weeks		Treatment-naive	equation)		33% vs.	Treatment-naïve:
interferon and		85 - 104 kg - 1200		Age >18 years	Platelet count <80x109/L		35%	100%
Ribavirin in		mg/day for 48		Liver biopsy in the past 2	Neutrophil count <1.5x109/L			
Hepatitis C		weeks		years consistent with	Hemoglobin concentration 13 g/dL and		Non	
Genotype 1-		>105 kg - 1400		chronic hepatitis	12 g/dL in men and women		white:	
Infected Slow		mg/day for 48			Co-existing uncontrolled psychiatric or		47% vs.	
Responders		weeks			cardiopulmonary disorders		48%	
					Hemoglobinopathy			
Overall Quality:		B: (Extended)			Sarcoidosis			
Fair		Ribavirin by body			Malignant neoplasm			
		weight:			Receipt of immunosuppressive or			
		< 64 kg - 800			immunomodulatory therapy in the			
		mg/day for 72			previous 6 months			
		weeks			Pregnancy			
		65 - 84 kg - 1000			Men whose partners were pregnant or			
		mg/day for 72			unwilling to use contraception during			
		weeks			the study period			
		85 - 104 kg - 1200			Patients were also excluded if they			
		mg/day for 72			imbibed significant amounts of alcohol			
		weeks			(30 g/day)			
		>105 kg - 1400			Active substance abusers in the past 6			
		mg/day for 72			months			
		weeks						

Author, Year Country							
Study Name	Duration of				Histologic		Funding
Quality	Followup	Outcome	Subgroup Analyses	Subgroup Analyses	Response	Adverse Events	Source
Pearlman, 2007 ³⁸	Followup	A vs. B:	A vs. B:	NR	NR	A vs. B:	NR
Atlanta, GA - USA	visits at 24					Overall withdrawals: 7/49(14%) vs. 8/52(15%);	
	weeks after	SVR: 9/49 (18%) vs.	SVR:			p=NS	
Continued	completion of	20/52 (38%), p=0.03	African Americans:			Withdrawals due to adverse events: 6/49(12%) vs.	
	treatment		12% vs. 21%, p=0.02			5/52(10%); p=NS	
						Deaths: NR	
						Life-threatening Adverse Events: NR	
						Serious Adverse Events: NR	
						Dose Reduction due to Adverse Event: (Week 1 -19) - 14/49(29%) vs. 15/52(29%); p=NS (Week 24-48) - 4/49(8%) vs. 2/52(4%); p=NS	
						Discontinuation due to Adverse Event: (Week 24-48) - 7/49(14%) vs. 8/52(15%); p=NS	

Author, Year Country Study Name Quality Sanchez-Tapias,	Interferon Regimen Patients with positive	Ribavirin Regimen Patients with	Protease Inhibitor Regimen None	Eligibility Treatment-naive patients	Exclusion Decompensated liver disease	Number Screened/ Eligible/ Enrolled/ Analyzed NR/NR/522/	Age Sex Race (A vs. B	Genotype Severity of Liver Disease Proportion Treatment-Naïve (A vs. B vs. C vs. D):
2006 ³⁹ Spain Peginterferon-Alfa2a Plus Ribavirin for 48 Vs. 72 weeks in Patients with Detectable Hepatitis C Virus RNA at Week 4 of Treatment	HCV RNA at week 4 randomized to 48 or 72 weeks A: (Total treatment duration, 48 weeks) Pegylated interferonalfa-2a 180 µg/week for 48 weeks B: (Total treatment duration, 72 weeks) Pegylated interferon-	positive HCV RNA at week 4 randomized to 48 or 72 weeks A: (Total treatment duration, 48 weeks) Ribavirin 800 mg/day for 48 weeks B: (Total treatment duration, 72 weeks) Ribavirin 800 mg/day for 72		with CHC consecutively referred to 28 specialist hepatology centers in Spain Older than 18 years Persistent increase of serum alanine transaminase levels during the past 6 months Positive anti-HCV antibody test Serum HCV-RNA concentration greater	Co-existing serious medical or psychiatric illness Liver disease other than that caused by HCV infection Neutrophil count less than 1.5 x109/L Platelet count less than 90x109/L Hemoglobin concentration less than 12 g/dL in women or less than 13 g/dL in men Serum creatinine level greater than 1.5 times the upper limit of the normal range Presence of co-infection with hepatitis	Randomized population: 326/326	vs. C vs. D): Age (Mean): 42.8 vs. 43.2 vs. 39.3 vs. 42.4 years Female: 21% vs. 27% vs.	Genotype 1: 90.3% vs. 88.2% vs. 30.4% vs. 97% Genotype 2: .6% vs6% vs. 12.2% vs. 0% Genotype 3: 4% vs. 5% vs. 50.7% vs. 0% Genotype 4: 5% vs. 5% vs. 6.8% vs. 3% Other (not-typeable): 0% vs. 1.2% vs. 0% vs. 0%
Overall Quality: Fair	Pegylated interferonalfa-2a 180 μg/week for 72 weeks Arms C and D not randomized (24 or 48 by genotype) C: (Total treatment duration, 24 weeks: RVR at week 4 and HCV-RNA levels <800,000 IU/mL) Pegylated interferonalfa-2a 180 μg/week for 24 weeks D: (Total treatment duration, 48 weeks: Genotype 1/4, RVR at week 4 and HCV-RNA levels >800,000 IU/mL) Pegylated interferonalfa-2a 180 μg/week for	mg/day for /2 weeks Arms C and D not randomized (24 or 48 by genotype) C: (Total treatment duration, 24 weeks: RVR at week 4 and HCV-RNA levels <800,000 IU/mL) Ribavirin 800 mg/day for 24 weeks D: (Total treatment duration, 48 weeks: Genotype 1/4, RVR at week 4 and HCV- RNA levels >800,000 IU/mL) Ribavirin 800 mg/day for 48 weeks		concentration greater than 600 IU/mL Histologic evidence of chronic hepatitis in a liver biopsy specimen obtained within the preceding 24 months Written informed consent to participate in the study All participants had to use 2 forms of effective contraception during treatment and throughout the 24-week followup phase of the study	Presence of co-infection with hepatitis A virus Hepatitis B virus or human immunodeficiency virus (HIV) Patients who received any systemic antiviral, antineoplastic, or immunomodulatory therapy within 6 months before the study Pregnant and breast-feeding women and male partners of pregnant women		27% vs. 30% vs. 44% Non white: NR	vs. 0% HCV-RNA>800,00 IU/mL (Mean): 963 vs. 1110 vs. 648 vs. 1612 Treatment naive: 100%

Author, Year Country Study Name Quality	Duration of Followup	Outcome	Subgroup Analyses	Subgroup Analyses	Histologic Response	Adverse Events	Funding Source
Sanchez-Tapias, 2006 ³⁹ Spain Continued	Followup visits at 24 weeks after completion of treatment	A vs. B: SVR: 53/165(32%) vs. 72/161(45%)	NR	NR	NR	A vs. B: Deaths: NR Serious Adverse Events: 4.8% vs. 8%; p=NS Treatment discontinuation - 29/165(18%) vs. 58/161(36%); p<0.001 Discontinuation due to Adverse event - 14/165(8%) vs. 19/161 (12%); p=NS Dose reduction - 74/165(45%) vs. 96/161 (59%); p=NS Adverse Events: Asthenia - 98/165(59%) vs. 95/161 (59%); p=NS Headache - 50/165(30%) vs. 53/161 (33%); p=NS Fever - 45/165(27%) vs. 45/161 (28%); p=NS Neutropenia - 40/165(24%) vs. 41/161 (25%); p=NS Influenza-like symptoms - 39/165(24%) vs. 28/148 (17%); p=NS Pruritus - 34/165(21%) vs. 41/161 (25%); p=NS Insomnia - 29/165(18%) vs. 41/161 (25%); p=NS Anorexia - 34/165(21%) vs. 23/161 (14%); p=NS Irritability - 28/165(17%) vs. 35/161 (22%); p=NS Anemia - 30/165(18%) vs. 34/161 (21%); p=NS Depression - 19/165(12%) vs. 31/161 (19%); p=NS Myalgia - 23/165(14%) vs. 22/161 (14%); p=NS Alopecia - 22/165(13%) vs. 27/161 (17%); p=NS	NR

Author, Year Country Study Name Quality	Duration of Followup	Outcome	Subgroup Analyses	Subgroup Analyses	Histologic Response	Adverse Events	Funding Source
Sanchez-Tapias, 2006 ³⁹ Spain Continued						Leukopenia - 18/165(11%) vs. 18/161 (11%); p=NS Injection site reaction - 12/165(7%) vs. 19/161 (12%); p=NS	

Author, Year Country Study Name	Latanfanan Bariman	Dikaninin Daniman	Protease Inhibitor	Fit all life.	Englasian	Number Screened/ Eligible/ Enrolled/	Age Sex	Genotype Severity of Liver Disease Proportion
Quality			Regimen	Eligibility	Exclusion	Analyzed	Race	Treatment-Naïve
Shiffman,	A: Pegylated interferon	A: Ribavirin 800	None	Eligible patients were	Other liver diseases	1810/1469/1	A vs. B:	A vs. B:
2007 ⁴⁰	alfa-2a 180 µg/week for			those who were 18 years	Human immunodeficiency virus (HIV)	469/1465		G
132 centers	16 weeks	weeks		of age or older	Hepatocellular carcinoma		Age	Genotype 2: 50.8%
worldwide				Infected with HCV	Severe depression or another severe		(Mean):	vs. 48.7%
	B: Pegylated interferon			genotype 2 or 3	psychiatric disease		46 vs.	Genotype 3: 49.2%
Pegylated	alfa-2a 180 µg/week for			Had a quantifiable serum	Clinically significant cardiovascular or		45.6 years	vs. 51.3%
interferon alfa-	24 weeks	weeks		HCV RNA level (>600	renal disease			
2a and Ribavirin				IU per milliliter)	Uncontrolled seizure disorder		Female:	Steatosis (% of
for 16 or 24				Elevated serum alanine	Severe retinopathy		39% vs.	hepatocytes):
Weeks in HCV				transaminase level	Previously received interferon or		37%	none - 20% vs. 21%
Genotype 2 or 3				Findings on liver biopsy	ribavirin (not treatment naive)			>0-5% - 26% vs. 25%
				consistent with chronic	Patients with cirrhosis had to have a		Non	6-33% - 12% vs. 12%
Overall Quality:				HCV infection	Child–Pugh score of less than 7 to be		white:	34-66% - 7% vs. 7%
Good					eligible		13% vs.	>66% - 2% vs. <1%
							13%	unknown - 33% vs.
								34%
								Treatment naïve:
								100%

Author, Year Country Study Name	Duration of	0.4	G. L		Histologic	Adams Francis	Funding
Quality Shiffman et al, 2007	Followup	Outcome A vs. B:	Subgroup Analyses NR	Subgroup Analyses A vs. B:	Response NR	Adverse Events A vs. B:	Source Roche
132 centers worldwide	Followup	A VS. B:	NK		NK	Deaths: NR	Roche
132 centers worldwide	weeks after	ETD: 651/722(800/) via		(p-value for		Life-threatening Adverse Events: NR	
Continued	completion of	ETR: 651/732(89%) vs. 599/731(82%)		interaction)		Serious Adverse Events: 5% vs. 6%	
Continued	treatment	399/731(82%)		Ganatura 2:		Withdrawals: 41/736(5%) vs. 91/731(12%);	
	treatment	SVD. 455/722(620/) via		Genotype 2: 232/358(62%) vs.		p<0.0001	
		SVR: 455/732(62%) vs. 515/731(70%); p<0.001		268/356(75%);		Withdrawal due to Adverse Events: 30/736(4%)	
		313/731(70%), p<0.001		p=0.06		vs. 25/731(5%); p=ns	
				Genotype 3:		Neutropenia (Grade 4): 13/733 (2%) vs. 20/732	
				221/358(62%) vs.		(3%); p=ns	
				244/369(66%);		Anemia (<8.5 g/dL): 4/733 (<1%) vs. 4/732	
				244/309(00%),		(<1%); p=ns	
				HCVRNA >800:			
				280/506 (55%) vs.			
				344/501 (67%):			
				p=0.26			
				HCVRNA 400-800:			
				43/65 (66%) vs.			
				59/80 (74%)			
				HCVRNA<400:			
				132/161 (82%) vs.			
				122/150 (81%)			
				Cirrhosis or bridging			
				fibrosis: 88/185			
				(48%) vs. 95/165			
				(58%); p=0.82			
				No Cirrhosis or			
				bridging fibrosis:			
				367/547 (67%) vs.			
				420/566 (74%)			

Author, Year Country Study Name Quality	Interferon Regimen	Ribavirin Regimen	Protease Inhibitor Regimen	Eligibility	Exclusion	Number Screened/ Eligible/ Enrolled/ Analyzed	Age Sex Race	Genotype Severity of Liver Disease Proportion Treatment-Naïve
Von Wagner, 2005 ⁴¹ Germany Peginterferonalpha-2a (40KD) and Ribavirin for 16 or 24 Weeks in Patients with Genotype 2 or 3 Chronic Hepatitis C Overall Quality: Fair	Patients with negative HCV RNA at week 4 randomized to either 16 or 24 weeks of treatment A: Pegylated interferon alfa-2a 180 µg/week for 16 weeks B: Pegylated interferon alfa-2a 180 µg/week for 24 weeks C: Pegylated interferon alfa-2a 180 µg/week for 24 weeks (non randomized patients who did not achieve RVR)	A: Ribavirin by body weight: < 65 kg - 800 mg/day for 16 weeks 65 - 85 kg - 1000 mg/day for 16 weeks > 85 kg - 1200 mg/day for 16 weeks B: Ribavirin by body weight: < 65 kg - 800 mg/day for 24 weeks 65 - 85 kg - 1000 mg/day for 24 weeks C: (Nonrandomized): Ribavirin by body weight: < 65 kg - 800 mg/day for 24 weeks 5 kg - 1200 mg/day for 24 weeks 5 kg - 1200 mg/day for 24 weeks 5 kg - 800 mg/day for 24 weeks 5 kg - 800 mg/day for 24 weeks 5 kg - 1000 mg/day for 24 weeks 5 85 kg - 1200 mg/day for 24 weeks 5 85 kg - 1200 mg/day for 24 weeks 5 85 kg - 1200 mg/day for 24	None	Male and female patients above 18 years of age with compensated chronic HCV infection not previously treated with interferon and/or ribavirin Tested positive for anti-HCV antibody and for HCV RNA (600 IU/mL by quantitative reverse transcription-polymerase chain reaction) Had a liver biopsy specimen taken within 18 months prior to the screening visit showing chronic hepatitis Had at least 1 serum alanine aminotransferase (ALT) level elevated at screening or entry into the trial Entry neutrophil and platelet counts at least 1500/ L and 90,000/ L, respectively Hemoglobin values at entry visit at least 12 g/dL for females and at least 13 g/dL for males	Any other cause of liver disease or other relevant disorders including human immunodeficiency or hepatitis B virus co infection Clinically significant hematologic, hepatic, metabolic, renal, rheumatologic, neurologic, or psychiatric disease Clinically significant cardiac or cardiovascular abnormalities; Organ grafts Systemic infection Clinically significant bleeding disorders Evidence of malignant neoplastic disease Concomitant immunosuppressive medication Excessive daily intake of alcohol or drug abuse within the past year Pregnancy and lactation, and male partners of pregnant women	NR/153/153/ 153	(A vs. B vs. C): Age (Mean): 38 vs. 39 vs. 42 years Female: 26% vs. 42% vs. 64% Non white: MR	(A vs. B vs. C): Genotype 2: 27% vs. 27% vs. 9% Genotype 3: 72% vs. 73% vs. 91% Fibrosis (Mean Ishak score): A (interface hepatitis) - 1 vs. 1.1 vs. 1.4 B (confluent necrosis) - 0.3 vs. 0.4 vs. 0.4 C (focal inflammation) - 1.4 vs. 1.4 vs. 1.4 D (portal inflammation) - 1.6 vs. 1.7 vs. 1.8 A-D(total inflammation) - 4.3 vs. 4.6 vs. 5.0 F (fibrosis) - 1.6 vs. 1.6 vs. 1.6 vs. 2.4 Cirrhosis: NR Treatment naive: 100%

Author, Year Country Study Name Quality	Duration of Followup	Outcome	Subgroup Analyses	Subgroup Analyses	Histologic Response	Adverse Events	Funding Source
Von Wagner, 2005 ⁴¹	Followup	A vs. B:	NR	A vs. B:	NR	A vs. B:	Hoffman-La
Germany	visits at 4, 12					Withdrawals: 1/71 (1.4%) vs. 6/71 (8.5%)	Roche
	and 24 weeks	SVR: 58/71(82%) vs.		SVR:		Withdrawal due to Adverse Events: NR	(Grenzach,
Continued	after	57/71(80%); p=NS		Genotype 2:18/19		Deaths: NR	Germany) &
	completion of			(95%) vs. 18/19		Life-threatening Adverse Events: NR	the German
	treatment			(95%); p=NS		Severe Adverse Events: NR	Hepatitis
				Genotype 3: 39/51		Withdrawals: NR	Network of
				(76%) vs. 39/52		Adverse events:	Competence
				(75%); p=NS		Flu-like symptoms: 37/71(52.1%) vs. 33/71 (46.5%); p=NS	(Hep-Net)
				HCVRNA <800:		Fatigue: 26/71(36.6%) vs. 30/71 (42.3%); p=NS	
				33/35 (94%) vs.		Pruritus: 19/71(26.8%) vs. 24/71 (33.8%); p=NS	
				27/31 (87%); p=NS		Headache: 18/71(25.4%) vs. 22/71 (31.0%); p=NS	
				HCVRNA>800:		Anorexia: 16/71(22.5%) vs. 19/71 (26.8%); p=NS	
				24/35 (69%) vs.		Alopecia: 15/71(21.1%) vs. 18/71 (25.4%); p=NS	
				30/40 (75%); p=NS		Asthenia: 12/71(16.9%) vs. 18/71(25.4%); p=NS	
						Pain: 9/71(12.7%) vs. 16/71(22.5%); p=NS	
						Dyspnea: 10/71(14.1%) vs. 16/71(22.5%); p=NS	
						Sleeping disturbance: 9/71(12.7%) vs. 16 (22.5%); p=NS	
						Pyrexia: 10/71(14.1%) vs. 13/71(18.3%); p=NS	
						Dry skin: 13/71(18.3%) vs. 9/71(12.7%); p=NS	
						Aggressivity: 8/71(11.3%) vs. 12/71(16.9%);	
						p=NS	
						Depression: 8/71(11.3%) vs. 10/71 (14.1%); p=NS	
						Chills: 10/71(14.1%) vs. 8/71(11.3%); p=NS	
						Nausea: 5/71(7.0%) vs. 11/71(15.5%); p=NS	
						Dry Mouth: 4/71(5.6%) vs. 8/71(11.3%); p=NS	

Author, Year Country Study Name	Landau	Dia dia pada dia	Protease Inhibitor	Fig. 1. 14	For location	Number Screened/ Eligible/ Enrolled/	Age Sex	Genotype Severity of Liver Disease Proportion
Quality	Interferon Regimen	Ribavirin Regimen		Eligibility	Exclusion	Analyzed	Race	Treatment-Naïve
Yu, 2006 ⁴²	A: Pegylated interferon	A: Ribavirin by	None	Eligible subjects were	Patients with HCV genotype other than	NR/NR/60/6	A vs. B:	A vs. B:
Taiwan	alpha-2b by body	body weight:		previously untreated Taiwanese chronic	1b infection Hepatitis B surface antigen	0	A 00	Canatuma 1h. 1000/
A randomized	weight: < 60 kg - 80 μg/week	< 75 kg - 1000					Age (Mean):	Genotype 1b: 100%
trial of 24- vs.	for 24 weeks	mg/day for 24 weeks		hepatitis C patients	Human immunodeficiency virus infection		45.4 vs.	Fibrosis Score
	> 60 kg - 100 μg/week	> 75 kg - 1200		18 to 65 years old, who:	Autoimmune hepatitis		45.4 vs. 45.1 years	(Knodell, 1981):
of PEG	for 24 weeks	mg/day for 24		(1) Were seropositive for HCV antibodies and	Primary biliary cirrhosis		43.1 years	Score 0–2 - 71.1% vs.
interferon alpha-	101 24 Weeks	weeks		HCV RNA by	Sclerosing cholangitis		Female:	73.3%
•	B: Pegylated interferon	WEEKS		polymerase chain	Wilson's disease		38% vs.	Score 3–4 - 28.9% vs.
for genotype-1b-	alpha-2b by body	B: Ribavirin by		reaction (PCR);	a1-antitrypsin deficiency		27%	26.7%
infected chronic	weight:	body weight:		(2) Had undergone a	Decompensated cirrhosis (Child–Pugh		2770	20.770
hepatitis C	< 60 kg - 80 μg/week	< 75 kg - 1000		liver biopsy within 1	class B or C)		Non	Treatment naïve:
patients: a pilot	for 48 weeks	mg/day for 48		vear before entry that	Overt hepatic failure		white: NR	100%
study in Taiwan	$> 60 \text{ kg} - 100 \mu\text{g/week}$	weeks		was consistent with	History of alcohol abuse		Willie. Tite	10070
stady in raiwan	for 48 weeks	> 75 kg - 1200		chronic hepatitis;	Psychiatric condition			
Overall Quality:	Tot to weeks	mg/day for 48		(3) Had displayed	Previous liver transplantation or with			
Fair		weeks		elevated serum alanine	evidence of hepatocellular carcinoma			
				transaminase (ALT),				
				defined as >1.5 times the				
				upper limit of the normal				
				range for at least two				
				measurements within 6				
				months preceding the				
				trial entry;				
				(4) Possessed an HCV				
				genotype 1b infection				
				Neutrophil count greater				
				than 1500/mm ³				

Author, Year Country Study Name Quality	Interferon Regimen	Ribavirin Regimen	Protease Inhibitor	Eligibility			Age Sex Race	Genotype Severity of Liver Disease Proportion Treatment-Naïve
Yu, 2006 ⁴²	Interferon Regimen	Kibavii iii Kegiiileii	Regimen	Platelet count greater	Exclusion	Anaryzeu	Nacc	11 catilient-1 (aive
Taiwan				than 1x105/mm3				
1 11 11 11 11				Hemoglobin level greater				
Continued				than 13 g/dl for males				
				and 12 g/dl for females				
				Serum creatinine level				
				less than 1.5 mg/dl				
				No pregnancy or				
				lactation and the use of a				
				reliable method of				
				contraception				

Author, Year Country Study Name Quality	Duration of Followup	Outcome	Subgroup Analyses	Subgroup Analyses	Histologic Response	Adverse Events	Funding Source
Yu, 2006 ⁴² Taiwan Continued	Followup visits at 24 weeks after completion of treatment	A vs. B: SVR: 22/45(48.9%) vs. 12/15(80%)	A vs. B: SVR: Male: 14/28 (50%) vs. 8/11 (72%); p=NS Female: 8/17 (47%) vs. 4/4 (100%); p=NS	A vs. B: SVR: Fibrosis score 0-2: 18/32(56.3%) vs. 8/11(72.7%), p=NS Fibrosis score 3-4: 4/13(30.8%) vs. 4/4(100%), p=0.029 Baseline HCV-RNA <400,000 IU/mL: 14/22(63.6%) vs. 4/5(80%), p=NS Baseline HCV-RNA >400,000 IU/mL: 8/23(34.8%) vs. 8/10(80%), p=0.026	NR NR	A vs. B: Deaths: NR Life-threatening Adverse Events: NR Severe Adverse Events: NR Withdrawals: 1/45 (2%) vs. 3/15 (20%); p=0.02 Withdrawal due to Adverse Events: 1/45 (2%) vs. 2/15 (13%); p=NS Dose reduction due to Adverse Events: 19/45 (42.2%) vs. 7/15 (46.7%); p=NS Adverse Events: Fever - 31/45 (68.9%) vs. 10/15 (66.7%); p=NS Chills - 10/45 (22.2%) vs. 4/15 (26.7%); p=NS Myalgia - 26/45 (57.7%) vs. 6/15 (40.0%); p=NS Headache - 32/45 (71.1%) vs. 9/15 (60.0%); p=NS Anorexia - 14/45 (31.1%) vs. 3/15 (20.0%); p=NS Nausea - 16/45 (35.6%) vs. 6/15 (40.0%); p=NS Diarrhea - 3/45 (6.7%) vs. 3/15 (20.0%); p=NS Anxiety/depression - 19/45 (42.2%) vs. 8/15 (53.3%); p=NS Insomnia - 26/45 (57.7%) vs. 10/15 (66.6%); p=NS Hair loss - 24/45 (53.3%) vs. 10/15 (66.6%); p=NS Skin rash - 30/45 (66.7%) vs. 9/15 (60.0%); p=NS	Taiwan Liver Research Foundation
						i	

Author, Year Country							
Study Name	Duration of				Histologic		Funding
Quality	Followup	Outcome	Subgroup Analyses	Subgroup Analyses	Response	Adverse Events	Source
Yu, 2006 ⁴²						Body weight loss - 8/45 (17.7%) vs. 2/15 (13.3%);	
Taiwan						p=NS	
						Anemia (hemoglobin o10 g/dl) - 20/45 (44.4%)	
Continued						vs. 8/15 (53.3%); p=NS	
						Leukopenia	
						White cell count <3000/mm3 - 34/45 (75.5%) vs.	
						11/15 (73.3%); p=NS	
						White cell count <1500/mm3 - 1/45 (2.2%) vs.	
						2/15 (13.3%); p=NS	
						Thrombocytopenia (<100 K/mm3) - 20/45	
						(44.4%) vs. 4/15 (26.6%); p=NS	
						Abnormal thyroid function tests - 4/45 (8.8%) vs.	
						1/15 (6.6%); p=NS	

Author, Year Country			Protease			Number Screened/ Eligible/	Age	Genotype Severity of Liver Disease
Study Name			Inhibitor			Enrolled/	Sex	Proportion
Quality	Interferon Regimen	Ribavirin Regimen	Regimen	Eligibility	Exclusion	Analyzed	Race	Treatment-Naïve
Yu, 2007 ⁴³	A: Pegylated interferon	A: Ribavirin by	None	Eligible patients were	Patients with an HCV genotype	326/152/150	A vs. B:	A vs. B:
Taiwan	alfa-2a 180 µg/week for	body weight:		previously untreated	infection other than type 2 infection	/150		
	24 weeks	< 75 kg - 1000		Taiwanese patients with	Hepatitis B surface antigen		Age	Genotype 2 - 100%
A randomized		mg/day for 24		CHC, aged 18–65 years,	HIV infection		(Mean):	
study of	B: Pegylated interferon	weeks		who:	Autoimmune hepatitis		49.4 vs.	Fibrosis (Knodell)
pegylated	alfa-2a 180 µg/week for	> 75 kg - 1200		(1) Were seropositive for	Primary biliary cirrhosis		50.2 years	F 0-2 - 80% vs. 78%
interferon and	16 weeks	mg/day for 24		HCV antibodies	Sclerosing cholangitis			F 3-4 - 20% vs. 22%
ribavirin for 16		weeks		(2) Had undergone a	Wilson's disease		Female:	Steatosis
vs. 24 weeks in				liver biopsy within 1	a1-antitrypsin deficiency		40% vs.	None (0) - 67% vs.
patients with		B: Ribavirin by		year before entry, the	Decompensated cirrhosis (Child–Pugh		34%	68%
genotype 2		body weight:		result of which was	class B or C)			Mild (1) - 28% vs.
chronic hepatitis		< 75 kg - 1000		consistent with chronic	Overt hepatic failure		Non	26%
C		mg/day for 16		hepatitis	Current alcohol misuse or history of		white: NR	Moderate to severe
		weeks		(3) Displayed an	alcohol misuse (>20 g/day)			(2–3) - 5% vs. 6%
Overall Quality:		> 75 kg - 1200		increased serum alanine	Psychiatric condition			
Fair		mg/day for 16		transaminase level,	Previous liver transplantation			Treatment naïve: NR
		weeks		defined as >1.5 times the	Evidence of hepatocellular carcinoma			
				upper limit of the normal	were excluded from the study			
				range for at least two				
				measurements within 6				
				months preceding the				
				trial entry				
				(4) Had HCV2 infection				

Author, Year Country Study Name			Protease Inhibitor			Number Screened/ Eligible/ Enrolled/	Age Sex	Genotype Severity of Liver Disease Proportion
Quality	Interferon Regimen	Ribavirin Regimen	Regimen	Eligibility	Exclusion	Analyzed	Race	Treatment-Naïve
Yu, 2007 ⁴³				Neutrophil count				
Taiwan				>1500/mm3				
				Platelet count				
Continued				>9x104/mm3				
				Hemoglobin				
				concentration >12 g/dl				
				for men, and 11 g/dl for				
				women				
				Serum creatinine				
				concentration < 1.5				
				mg/dl				
				No pregnancy or				
				lactation				
				Use of a reliable method				
				of contraception for				
				women				

Author, Year							
Country							
Study Name	Duration of				Histologic		Funding
Quality	Followup	Outcome	Subgroup Analyses	Subgroup Analyses	Response	Adverse Events	Source
Yu, 2007 ⁴³	Followup	A vs. B:	A vs. B:	A vs. B:	NR	A vs. B:	Taiwan Liver
Taiwan	visits at 24					Deaths: NR	Research
	weeks after	95/100 (95%) vs. 47/50	Age:	Fibrosis F0-2: 76/80		Life-threatening Adverse Events: NR	Foundation
Continued	completion of	(94%); p=NS	<50 years -	(95%) vs. 34/39		Severe Adverse Events: NR	
	treatment		46/46(100%) vs.	(95%); p=NS		Withdrawals: 1/100(1%) vs. 0/50(0%); p=1	
			19/19(100%);p=NS	Fibrosis F3-4: 19/20		Withdrawal due to Adverse Events: 1/100(1%) vs.	
			>50 years - 49/54(91%)	(95%) vs. 10/11		0/50(0%); p=1	
			vs. 28/31(90%); p=NS	(91%); p=NS		Dose reduction due to Adverse Events -	
						54/100(54%) vs. 26/50 (52%), p=0.817	
			Female: 38/42(91%) vs.	HCVRNA <800K:			
			16/18(89%); p=NS	81/85 (95%) vs.		Adverse Events:	
			Male: 57/58 (98%) vs.	39/41 (95%); p=NS		Fever: 55/100 (55%) vs. 29/50 (58%), p=0.727	
			31/32 (97%): p=NS	HCVRNA>800K:		Chills: 28/100 (28%) vs. 12/50 (24%), p=0.602	
			DM 05 40/52 (020/)	14/15 (93%) vs. 8/9		Headache: 39/100 (39%) vs. 21 /50 (42%),	
			BMI <25: 49/53 (93%)	(89%); p=NS		p=0.724	
			vs. 25/27 (93%); p=NS			Anorexia: 46/100 (46%) vs. 20/50 (40%), p=0.601	
			BMI>25: 46/47 (98%)			Nausea: 15/100 (15%) vs. 3/50 (6%), p=0.181	
			vs. 22/23 (96%); p=NS			Diarrhea: 9/100 (9%) vs. 5/50 (10%), p=1	
						Anxiety: 7/100 (7%) vs. 4/50 (8%), p=1 Depression: 10/100 (10%) vs. 3/50 (6%), p=0.545	
						Insomnia: 57/100 (57%) vs. 23/50 (46%), p=0.227 Hair loss: 49/100 (49%) vs. 10/50 (20%),	
						p=0.001*	
						Skin rash: 54/100 (54%) vs. 22/50 (44%), p=	
						0.248	
						Leukopenia(white cell count,1500/mm3): 2/100	
						(2%) vs. 1/50 (2%), p=1	
						Anemia (hemoglobin level<10g/dl): 53/100(53%)	
						vs. 27/50 (54%), p=0.908	
						Thrombocytopenia(<50,000/mm3: 1/100 (1%) vs.	
						0/50 (0%), p=1	
						Abnormal thyroid function tests: 13/100 (13%) vs.	
						4/50 (8%), p=0.362	

						Number		Genotype
Author, Year						Screened/		Severity of Liver
Country			Protease			Eligible/	Age	Disease
Study Name			Inhibitor			Engible/ Enrolled/	Sex	Proportion
Quality	Interferon Regimen	Ribavirin Regimen		Eligibility	Exclusion	Analyzed	Race	Treatment-Naïve
Yu, 2008 ⁴⁴	A: Pegylated interferon	A: Ribavirin by	None	Eligible patients were	Patients with an HCV genotype	NR/NR/200/	A vs. B:	A vs. B:
Taiwan	alfa-2a 180 µg/week for	body weight:	None	previously untreated	infection other than type 1 infection	200	A vs. D.	A vs. D.
Taiwaii	24 weeks	< 75 kg - 1000		Taiwanese patients with	Hepatitis B surface antigen	200	Age	Genotype 1 - 100%
Rapid	24 weeks	mg/day for 24		CHC, aged 18–65 years,	HIV infection		(Mean):	Genotype 1 - 100%
	D. Dl. d. intenferen	weeks		who:	Autoimmune hepatitis		49.7 vs.	E:1:- (V1-11)
Virological	B: Pegylated interferon							Fibrosis (Knodell)
Response and	alfa-2a 180 µg/week for			(1) Were seropositive for	Primary biliary cirrhosis		49.1 years	F 0–2 - 75% vs. 81%
Treatment	48 weeks	mg/day for 24		HCV antibodies	Sclerosing cholangitis		Б 1	F 3–4 - 25% vs. 19%
Duration for		weeks		(2) Had undergone a	Wilson's disease		Female:	TT
Chronic		D D'1 ' ' 1		liver biopsy within 1	al-antitrypsin deficiency		43% vs.	Treatment naïve: NR
Hepatitis C		B: Ribavirin by		year before entry, the	Decompensated cirrhosis (Child–Pugh		42%	
Genotype 1		body weight:		result of which was	class B or C)			
Patients: A		< 75 kg - 1000		consistent with chronic	Overt hepatic failure		Non	
Randomized		mg/day for 48		hepatitis	Current alcohol misuse or history of		white: NR	
Trial		weeks		(3) Displayed an	alcohol misuse (>20 g/day)			
		> 75 kg - 1200		increased serum alanine	Psychiatric condition			
Overall Quality:		mg/day for 48		transaminase level,	Previous liver transplantation			
Fair		weeks		defined as >1.5 times the	Evidence of hepatocellular carcinoma			
				upper limit of the normal	were excluded from the study			
				range for at least two				
				measurements within 6				
				months preceding the				
				trial entry				
				(4) Had HCV2 infection				
				Neutrophil count				
				>1500/mm3				
				Platelet count				
				>9x104/mm3				
				Hemoglobin				
				concentration >12 g/dl			1	
				for men, and 11 g/dl for				
				women			1	
				Serum creatinine				
				concentration < 1.5			1	
				mg/dl				

Author, Year Country							
Study Name	Duration of				Histologic		Funding
Quality	Followup	Outcome	Subgroup Analyses	Subgroup Analyses	Response	Adverse Events	Source
Yu, 2008 ⁴⁴	Followup	A vs. B:	A vs. B:	A vs. B:	NR	A vs. B:	Taiwan Liver
Taiwan	visits at 24					Withdrawals: 3/100(3%) vs. 10/100(10%),	Research
	weeks after	SVR: 59/100(59%) vs.	SVR:	SVR:		p=0.045	Foundation
Continued	completion of	79/100(79%)	Male: 34/57 (60%) vs.	Fibrosis score F0-2:		Withdrawal due to Adverse Events: 3/100 (3%)	
	treatment		46/58 (79%); p=NS	48/75 (64%) vs.		vs. 9/100 (9%); p=NS	
			Female: NR	62/84 (77%); p=NS		Deaths: NR	
				Fibrosis score F3-4:		Life-threatening Adverse Events: NR	
				11/25 (44%) vs.		Serious Adverse Events: 1/100 (1%) vs. 1/100	
				17/19 (89%);		(1%); p=NS	
				p=0.002		Dose reduction due to adverse events:	
				HCV RNA <400K:		54/100(54.0%) vs. 65/100(65.0%), p=0.113	
				34/45 (76%) vs.		Influenza-like symptoms (fever, chills, headache):	
				36/44 (82%); p=NS		76/100(76%) vs. 74/100(74%), p=0.744	
				HCV RNA>400K:		Anorexia and/or nausea - 50 (50%) vs. 53 (53%),	
				25/55 (45%) vs.		p=0.671	
				43/56 (77%);		Diarrhea - 18 (18%) vs. 26 (26%), p=0.172	
				p<0.001		Anxiety - 31 (32%) vs. 36/100(36%), p=0.454	
						Depression - 24 (24%) vs. 34/100(34%), p=0.119	
						Insomnia - 59 (59%) vs. 65/100(65%), p=0.382	
						Hair loss – 66/100(66%) vs. 72/100(72%),	
						p=0.359	
						Skin rash – 54/100(54%) vs. 66/100(66%),	
						p=0.083	
						Leukopenia (white cell count < 1500 mm-3) –	
						5/100(5%) vs. 8/100(8%), p=0.39	
						Anemia (hemoglobin < 10 g/dl) – 39/100(39%)	
						vs. 48/100(48%), p=0.199	
						Thrombocytopenia (< 50,000 mm-3) – 2/100(2%)	
						vs. 6/100(6%), p=0.279	
						Abnormal thyroid function tests – 13/100(13%)	
						vs. 15/100(15%), p=0.684	

						Number		Genotype
Author, Year						Screened/		Severity of Liver
Country			Protease			Eligible/	Age	Disease
Study Name			Inhibitor			Enrolled/	Sex	Proportion
Quality	Interferon Regimen	Ribavirin Regimen	Regimen	Eligibility	Exclusion	Analyzed	Race	Treatment-Naïve
Zeuzem, 2004 ⁴⁵	A: Pegylated interferon	A: Ribavirin 800	None	Treatment-naive patients	No histologic evidence of liver disease	NR/NR/514/	A vs. B	(A vs. B vs. C):
Australia,	alfa-2a 180 µg/week for	mg/day (2 equal		aged 18 years or older	One or more elevated ALT values (i.e.,	491	vs. C:	
Europe, New	24 weeks	doses) for 24 weeks		with a positive antibody	greater than the ULN) within the		Age	Genotype 1: 68% vs.
Zealand, North				to hepatitis C virus	previous 18 months		(Mean):	67% vs. 68%
& South	B: Pegylated interferon	B: Ribavirin 800		(HCV) antibody test	Patients with transition to cirrhosis or		44 vs. 44	Genotype 1a: 36% vs.
America	alfa-2a 180 µg/week for	mg/day (2 equal		Detectable HCV RNA in	cirrhosis on liver biopsy		vs. 41	42% vs. 38%
	48 weeks	doses) for 48 weeks		serum	History of bleeding from esophageal		years	Genotype 1b: 31% vs.
Pegylated				Biopsy findings	varices		Female:	25% vs. 30%
interferon alfa-	C: No treatment	C: No treatment		consistent with a	Other conditions consistent with		58% vs.	Genotype (other type
2a (40				diagnosis of chronic	decompensated liver disease were		61% vs.	1): 1% vs. 0% vs. 0%
Kilodaltons) and				hepatitis C	excluded to avoid the possibility of		62%	Genotype 2: 18% vs.
Ribavirin in				Persistently normal ALT	including individuals whose ALT levels		Non	20% vs. 19%
Patients with				levels (equal to or below	had returned to the normal range as a		white	Genotype 3: 9% vs.
Chronic				the upper limit) of	consequence of advanced liver disease		race:	9% vs. 9%
Hepatitis C and				normal (ULN)	Neutropenia (absolute neutrophil count		14%	Genotype 4: 4% vs.
Normal				documented on at least 3	1500 cells/mm3)		vs.14%	4% vs. 3%
Aminotransferas				occasions, a minimum of	Thrombocytopenia (90,000		vs. 17%	Genotype 5: 1% vs.
e Levels				4 weeks apart, with at	platelets/mm3)			0% vs. 0%
				least one value obtained	Anemia (hemoglobin concentration 12			Genotype 6: 1% vs.
Overall Quality:				during the 42-day	g/dL in women and 13 g/dL in men) or			1% vs. 1%
Fair				screening period and at	a medical condition that would be			
				least one value obtained	significantly worsened by anemia			Cirrhosis: 0% vs.
				6-18 months before	Serologic evidence of infection with			1% vs. 0%
				screening.	human immunodeficiency virus or			
					hepatitis A or B virus, and serum			Fibrosis (Ishak):
					creatinine level 1.5 times the ULN			0-1: 66% vs. 69% vs.
					Organ transplant recipients			77%
					Individuals with severe cardiac disease			2: 21% vs. 20% vs.
					History of severe psychiatric disease			14%
					(especially depression)		1	3-4: 12% vs. 9% vs.
					Evidence of drug abuse (including		1	7%
					excessive alcohol consumption) within the preceding year			>4: 0% vs. 1% vs. 0%
								Treatment naive:
								100%

Author, Year Country Study Name			Protease Inhibitor			Number Screened/ Eligible/ Enrolled/	Age Sex	Genotype Severity of Liver Disease Proportion
Quality	Interferon Regimen	Ribavirin Regimen		Eligibility	Exclusion		Race	Treatment-Naïve
Zeuzem, 2004 ⁴⁵					Other serious systemic disease	•		
Australia,					Pregnant or lactating women and male			
Europe, New					partners of pregnant women. All fertile			
Zealand, North					men and women who participated in the			
& South					trial were required to use two forms of			
America					effective contraception during treatment			
					and for 6 months after the end of			
Continued					treatment			

Author, Year Country			Protease Inhibitor			Number Screened/ Eligible/ Enrolled/	Age Sex	Genotype Severity of Liver Disease
Study Name Quality	Interferon Regimen	Ribavirin Regimen		Eligibility	Exclusion	Analyzed	Race	Proportion Treatment-Naïve
Zeuzem, 2004 ⁴⁵				8 1	Other serious systemic disease			
Australia,					Pregnant or lactating women and male			
Europe, New					partners of pregnant women. All fertile			
Zealand, North					men and women who participated in the			
& South					trial were required to use two forms of			
America					effective contraception during treatment			
					and for 6 months after the end of			
Continued					treatment			

Author, Year							
Country							
Study Name	Duration of				Histologic		Funding
Quality	Followup	Outcome	Subgroup Analyses	Subgroup Analyses	Response	Adverse Events	Source
Zeuzem, 2004 ⁴⁵	Followup	A vs. B:	NR	A vs. B:	NR	A vs. B:	Roche (Basel,
Australia, Europe,	visits at 24					Withdrawals: 20/212 (9%) vs. 58/210 (28%);	Switzerland)
New Zealand, North	weeks after	ETR: NR		ETR: NR		p<0.001	
& South America	completion of					Withdrawals due to adverse events: 15/212 (7%)	
	treatment	SVR: 63/212(30%) vs.		SVR:		vs. 38/210 (18%); p<0.001	
Continued		109/210(52%); p<0.001		Genotype 1 -		Severe adverse events 56/212 (26%) vs. 70/210	
				19/144(13%) vs.		(33%); p=NS	
				57/141(40%);		Life-threatening adverse events - 3/212 (1%) vs.	
				p<0.001		8/210 (4%)	
				Genotypes 2/3 - 42/58(72%) vs.		Serious adverse events - 18/212 (8%) vs. 34/210 (16%); p=0.02	
				46/59(78%); p=NS		Deaths - 0/212(0%) vs. 0/210(0%); p=NS	
				Genotypes 4 -		Dose reduction due to adverse events -	
				1/8(13%) vs.		65/212(32%) vs. 102/210(49%); p<0.001	
				5/9(56%); p=NS		05/212(32/0) vs. 102/210(15/0), p vs. 001	
						Adverse Events:	
				HCV RNA <800		Headache - 93/212 (44%) vs. 117/210 (56%);	
				IU/mL: 39/123		p=0.02	
				(32%) vs. 72/127		Fatigue - 109/212 (51%) vs. 107/210 (51%);	
				(57%); p<0.001		p=NS	
				HCV RNA >800		Myalgia - 81/212 (38%) vs. 93/210 (44%); p=NS	
				IU/mL: 24/87(28%)		Pyrexia – 64/212 (30%) vs. 90/210 (43%); p<0.01	
				vs. 36/82(44%);		Insomnia - 74/212 (35%) vs. 76/210 (36%); p=NS	
				p=0.03		Nausea - 68/212 (32%) vs. 84/210 (40%); p=NS	
						Arthralgia - 68/212 (32%) vs. 62/210 (30%);	
						p=NS	
						Depression - 55/212 (26%) vs. 57/210 (27%); p=NS	
						p=NS Irritability - 58/212 (27%) vs. 55/210 (26%);	
						p=NS	
						Rigors - 50/212 (24%) vs. 53/210 (25%); p=NS	
						Alopecia - 43/212 (20%) vs. 59/210 (28%); p=NS	
						Asthenia - 47/212 (22%) vs. 48/210 (23%); p=NS	

Author, Year Country							
Study Name	Duration of				Histologic		Funding
Quality	Followup	Outcome	Subgroup Analyses	Subgroup Analyses	Response	Adverse Events	Source
Zeuzem, 2004 ⁴⁵						Diarrhea - 40/212 (19%) vs. 55/210 (26%); p=NS	
Australia, Europe,						Pruritus - 34/212 (16%) vs. 42/210 (20%); p=0.03	
New Zealand, North						Hemoglobin <10.0 to >8.5 g/dL - 10/212 (5%) vs.	
& South America						24/210 (11%); p=0.01	
						Hemoglobin <8.5 g/dL - 3/212 (1%) vs. 1/210 (
Continued						1%); p=NS	
						Neutrophils <0.5 x109/L - 10/212 (5%) vs. 10/210	
						(5%); p=NS	
						Platelets <50 x109/L - 3/212 (1%) vs. 4/210 (2%);	
						p=NS	
						Hypothyroidism - 0/212 (0%) vs. 5/210 (2%);	
						p=NS	
						Hyperthyroidism - 1/212 (1%) vs. 3/210 (1%);	
						p=NS	

Evidence Table 6. Quality rating: Trials of dual therapy with pegylated interferon plus ribavirin: duration effects

Author, Year	Randomization adequate?	Allocation concealment adequate?	Groups similar at baseline?	Eligibility criteria specified?	Outcome assessors masked?	Care provider masked?	Patient masked?	Attrition and withdrawals reported?	Loss to followup: differential/high?	Intention- to-treat analysis	Quality	Funding
Andriulli, 2009 ²²	Unclear	Yes	Unclear	Yes	No, open label	No, open label	No, open label	Yes	No	Yes	Fair	Investigator funded
Berg, 2006 ²³	Unclear	Unclear	Yes	Yes	No, open label	No, open label	No, open label	yes	No	Yes	Fair	Roche
Berg, 2009 ²⁴	Unclear	Unclear	Unclear	Yes	No, open label	No, open label	No, open label	Yes	Yes	Yes	Poor	Schering- Plough
Brandao, 2006 ²⁵	Yes	Unclear	Yes	Yes	No, open label	No, open label	No, open label	Yes	No	Yes	Fair	Roche
Bronowicki, 2006 ⁴⁶	Yes	Unclear	Unclear	Yes	No, open label	No, open label	No, open label	Yes	No	Yes	Fair	Roche
Buti, 2010 ²⁶	Yes	Unclear	Yes	Yes	No, open label	No, open label	No, open label	Yes	No	Yes	Fair	Schering- Plough (now Merck)
Dalgard, 2008 ²⁷	Unclear	Unclear	Yes	Yes	No, open label	No, open label	No, open label	Yes	No	Yes	Fair	Schering- Plough (now Merck)
Ferenci, 2010 ²⁸	Unclear	unclear	Unclear	Yes	Unclear	Unclear	Unclear	Yes	Yes	Yes	Poor	Roche
Ide, 2009 ³⁰	Unclear	Yes	Unclear	Yes	Unclear	Unclear	Unclear	Yes	Yes	Yes	Fair	Internal Funding
Kamal, 2005 ³¹	Unclear	Unclear	Yes	Yes	Unclear	Unclear	Unclear	Yes	No	Yes	Fair	Fulbright Foundation Grants(NIAID (R2) AI054887) & the Alexander von Humboldt Foundation (Germany)

Author, Year	Randomization adequate?	Allocation concealment adequate?	Groups similar at baseline?	Eligibility criteria specified?	Outcome assessors masked?	Care provider masked?	Patient masked?	Attrition and withdrawals reported?	Loss to followup: differential/high?	Intention- to-treat analysis	Quality	Funding
Lagging, 2008 ³²	Unclear	Unclear	Yes	Yes	No, open label	No, open label	No, open label	Yes	No	Yes	Fair	Swedish Society of Medicine, Swedish Medical Council, Swedish Society of Microbiology, Avtal om lakarutbildning och forskning (ALF) Funds, and Roche affiliates (Nordic region)
Lam, 2010 ³³	Unclear	Yes	Yes	Yes	No, open label	No, open label	No, open label	No	No	Yes	Fair	investigator initiated research grant from Roche Laboratories, LLC to Pacific Health Foundation
Liu, 2008 ³⁴	Unclear	Unclear	Yes	Yes	No, open label	No, open label	No, open label	Yes	No	Yes	Fair	National Taiwan University Hospital, National Science Council, and Department of Health, Executive Yuan, Taiwan
Mangia, 2005 ³⁵	Unclear	Unclear	Yes	Yes	No, open label	No, open label	No, open label	No	No	Yes	Fair	Italian branch of Schering- Plough
Manns 2011 ³⁶	No	Yes	Unclear	Yes	No, open label	No, open label	No, open label	Yes	Yes	Yes	Poor	Schering- Plough (now Merck)

Author, Year	Randomization adequate?	Allocation concealment adequate?	Groups similar at baseline?	Eligibility criteria specified?	Outcome assessors masked?	Care provider masked?	Patient masked?	Attrition and withdrawals reported?	Loss to followup: differential/high?	Intention- to-treat analysis	Quality	Funding
Mecenate, 2010 ³⁷	Unclear	Unclear	Unclear	Yes	No, open label	No, open label	No, open label	Yes	No	Yes	Fair	NR
Pearlman, 2007 ³⁸	Unclear	Unclear	Yes	Yes	No, not described	No, not described	No, not described	Yes	No	Yes	Fair	NR
Sanchez-Tapias, 2006 ³⁹	Yes	Yes	Unclear	Yes	No, open label	No, open label	No, open label	Yes	No	Yes	Fair	NR
Shiffman, 2007 ⁴⁰	Unclear	Yes	Yes	Yes	No	Yes	Yes	Yes	No	Yes	Good	Roche
Von Wagner, 2005 ⁴¹	Unclear	Unclear	Unclear	Yes	Unclear	Unclear	Unclear	Yes	No	Yes	Fair	Hoffman-La Roche (Grenzach, Germany) & the German Hepatitis Network of Competence (Hep-Net)
Yu, 2006 ⁴²	Yes	Unclear	Unclear	Yes	No, open label	No, open label	No, open label	Yes	No	Yes	Fair	Taiwan Liver Research Foundation
Yu, 2007 ⁴³	Yes	Unclear	Unclear	Yes	No, open label	No, open label	No, open label	Yes	No	Yes	Fair	Taiwan Liver Research Foundation
Yu, 2008 ⁴⁴	Yes	Yes	Unclear	Yes	No - open label	No - open label	No - open label	Yes	No	Yes	Fair	Taiwan Liver Research Foundation
Zeuzem, 2004 ⁴⁵	Unclear	Unclear	Yes	Yes	No, open label	No, open label	No, open label	Yes	Yes	Yes	Fair	Roche (Basel, Switzerland)

Evidence Table 7. Trials of dual therapy with pegylated interferon plus ribavirin: dose effects

						Number		
Author, Year						Screened/		
Country			Protease			Eligible/		Genotype
Study Name	Interferon	Ribavirin	Inhibitor			Enrolled/	Sex	Severity of Liver Disease
Quality	Regimen	Regimen	Regimen	Eligibility	Exclusion	Analyzed	Race	Proportion Treatment-Naïve
Abergel, 2006 ⁴⁷	A: (standard-	A: Ribavirin	None	Age between 18 and 75	Recent history of alcohol abuse	NR/210/	A vs. B	A vs. B
France	dose)	800		years	or IV drug addiction	210/203	Age(Mean):	Genotype 1 - 50/101(49.5%) vs.
	Pegylated	mg/day/48weeks		No previous treatment	Hemoglobin <12 g/dL in women		49.3 vs. 51.1	54/102(529%)
Pegylated	interferon	B: Ribavirin		with IFN and/or	and <13 g/dL in men		years	Genotype 2 - 11/101(10.9%) vs.
interferon alpha-	alpha-2b 1.5	800 mg/day/48		ribavirin	Platelets <75 000/lL			9/102(8.8%)
2b plus ribavirin	μg/kg	weeks		Alanine	Neutrophils <1500/IL		Female: 36%	Genotype 3 - 30/101(29.7%) vs.
for treatment of	1x/week/48			aminotransferase (ALT)	Decompensated cirrhosis		vs. 32%	28/102(27.5%)
chronic hepatitis	weeks			> upper limit of normal	(ascites, variceal hemorrhage			Genotype 4 - 5/101(5%) vs.
C with severe	B: (low-dose)			(ULN) at least once	encephalopathy)		Race: NR	4/102(3.9%)
fibrosis: a	Pegylated			during the last 12	Albumin <30 g/L			Genotype 5 - 5/101(5%) vs.
multicenter	interferon			months	Prothrombin <60%			7/102(6.9%)
randomized	alpha-2b 0.75			Positive serum HCV-	Bilirubin >34 lmol/L			
controlled trial	μg/kg			RNA using qualitative	HCC			Fibrosis stage:
comparing two	1x/week/48			polymerase chain	Chronic hepatitis B infection			F3 - 55/101(54.4%) vs.
doses of	weeks			reaction (PCR) and	HIV infection			44/102(43.1%)
Pegylated				severe fibrosis on liver				F4 - 46/101(45.6%) vs.
interferon alpha-				biopsy defined by a				58/102(56.9%)
2b				METAVIR fibrosis				
				stage of F3 or				Cirrhosis: 46% vs. 57%
Overall Quality:				F4 at histological				
Fair				examination of the liver				100% Treatment naïve

Author, Year Country	Duration						
Study Name	of				Histologic		Funding
Quality	Followup	Outcome	Subgroup Analyses	Subgroup Analyses	Response	Adverse Events	Source
Abergel, 2006 ⁴⁷	Followup	A vs. B	A vs. B	A vs. B	None	A vs. B	Schering-
France	visits at 24	ETR: 59/101(62.8%)	ETR: NR	ETR: NR		Discontinuation - 30/101(31 %) vs. 28/102(27 %)	Plough,
	weeks after	vs. 57/102(59.4%)				Discontinuation or treatment reduction –	France and
Continued	completion		SVR:	SVR:		53/101(54%) vs. 37/102(36 %), p < 0.03	Delegation
	of	SVR: 50/101(49.5%)	BMI <27 kg/m2 -	Genotypes 1, 4, 5, -		Treatment reduction - 36/101(37%) vs.	Regionale a la
	treatment	vs. 38/102(37.2%)	35/70 (50.0%) vs.	15/60(25.0%) vs. 11/65		13/102(12%), p < 0.0002	Recherche
			26/70 (37.1%); p=NS	(16.9%); p=NS		Overall withdrawals - NR	Clinique,
			$BMI \ge 27 \text{ kg/m2} -$	Genotype 1 - 12/50		Deaths - NR	Clermont-
			10/31 (32.3%) vs.	(24.0%) vs. 09/54			Ferrand,
			12/32 (37.5%); p=NS	(16.7%); p=NS		Severe Adverse Events:	France
			, ,	Genotypes 2, 3 - 30/41		Adverse event - 8/101(9%) vs. 4/102(3%)	
			gamma glutamyl	(73.2%) vs. 27/37		Cytopenia -7/101(7%) vs. 1/102(1%)	
			transpeptidase (GGT)	(73.0%); p=NS		Others - 7/101(8%) vs. 3/102(2 %)	
			used as a marker for	M; ; ,000,000		A 1	
			steatosis:	Viremia <800.000		Adverse events	
			GGT <1.6 ULN - 29/48 (60.4%) vs.	IU/mL - 25/55 (45.5%) vs. 20/47 (42.5%);		Adverse event - 15/101(16%) vs. 4/102(3%), p	
			23/48 (47.9%); p=NS	vs. 20/47 (42.3%), p=NS		Cytopenia - 20/101(21 %) vs. 9/102(8%), <0.03	
			GGT >1.6 ULN -	Viremia <u>></u> 800 000		Anemia - 9/101(10%) vs. 5/102(4%)	
			13/50 (26.0%) vs.	IU/mL - 20/44 (45.5%)		Neutropenia - 10/101(11 %) vs. 4/102(3%)	
			13/50 (25.5%); p=NS	vs. 17/53 (32.1%);		Thrombopenia - 3/101(3 %) vs. 0/102(0%)	
			13/31 (23.370), p=113	p=NS		Others - 2/101(1%) vs. 0/102(1%)	
				P=145		Hemoglobin < 10g/dL - 27/101(27 %) vs.	
				Cirrhosis (F4) - 18/46		16/102(15%), p=0.054	
				(39.1%) vs. 20/58		Neutrophils $< 750/ \mu L - 21/101(21\%)$ vs.	
				(34.5%); p=NS		8/102(7%), p <0.01	
				Severe fibrosis(F3) -		Platelets < 50 000/ μL - 7/101(7%) vs. 7/102(6 %)	
				27/55 (49.1%) vs. 18/44		Depression - 13/101(12%) vs. 15/102(14%)	
				(40.1%); p=NS		Suicide - 2/101(1%) vs. 0/102(0%)	
				. //1		Hypothyroidism (treated) - 9/101(10%) vs.	
						1/102(.5%)	

Author, Year Country Study Name Quality	Interferon Regimen	Ribavirin Regimen	Protease Inhibitor Regimen	Eligibility	Exclusion	Number Screened/ Eligible/ Enrolled/ Analyzed	Age Sex Race	Genotype Severity of Liver Disease Proportion Treatment-Naïve
Brady, 2010 ⁴⁸ United States Induction pegylated interferon alfa-2b combination with ribavirin in patients with genotype 1 and 4 chronic hepatitis C: a prospective, randomized, multicenter, open- label study Overall Quality: Fair	A. Pegylated interferon alfa-2b 3.0 mcg/kg/week for 12 weeks followed by 1.5 mcg/kg/week for 36 weeks B. Pegylated interferon alfa-2b 1.5 mcg/kg/week for 48 weeks	A. 800-1400 mg/day for 48 weeks B. 800-1400 mg/day for 48 weeks	NA	Treatment-naïve patients Genotype 1 or 4 Positive HCV antibodies and detectable HCV RNA Liver biopsy consistent with viral hepatitis within the past 48 months Cirrhosis no worse than Child-Pugh Class A Hemoglobin ≥12 g/dL in females and 13 g/dL in males White blood cells ≥3000 Neutrophil ≥1500 Platelet ≥ 65K Direct bilirubin within 20% of upper limits of normal Creatinine within 20% of upper limits of normal Albumin within normal limits	Non genotype 1 or 4 HCV infection Decompensated liver disease Evidence of coexisting liver disease Coinfection with HIV or HBV Hemochromatosis Alpha-1 antitrypsin deficiency Wilson disease Autoimmune hepatitis Alcoholic liver disease Hepatocellular carcinoma Pregnancy Psychiatric conditions Significant cardiovascular dysfunction within the past 1 year Poorly controlled diabetes mellitus Chronic pulmonary disease Clinically significant retinal abnormalities Immunologically mediated diseases Any medical condition requiring systemic steroids Active clinical gout Substance abuse in the past 6 months	NR/NR/ 623/610	A vs. B Age mean: 45 vs. 45 Female: 50% vs. 50% non White: 32% vs. 28%	A vs. B genotype 1: 99% vs. 99% Treatment-naïve: all Fibrosis stage 3 or 4: 26% vs. 23% HCV- RNA ≥800K: 71% vs. 62%

Author, Year Country Study Name Quality	Duration of Followup	Outcome	Subgroup Analyses	Subgroup Analyses	Histologic Response	Adverse Events	Funding Source
Brady, 2010 ⁴⁸ United States Continued	24 weeks following treatment completion	A vs. B ETR: 126/299 (42.1%) vs. 121/311 (38.9%); p= SVR: 96/299 (32.1%) vs. 92/311 (29.6%); p=0.434	A vs. B Black: 13/36 (36.1%) vs. 12/37 (32.4%); p=0.9 Hispanic: 29.9% vs. 22.5%; p=0.292 (absolute numbers NR) Weight <85 kg: 26% vs. 31% (p=NS); (absolute numbers NR) Weight ≥85 kg: 38% vs. 28% (p=0.08); (absolute numbers NR)	NR	NR	A vs. B Overall withdrawals: 146/299 (48.8%) vs. 133/311 (42.7%); p=0.2 Withdrawals for adverse events: NR Serious adverse events: NR Deaths: NR Neutropenia <500: 10/299 (3.4%) vs. 5/311 (1.6%); p=0.261 Anemia hemoglobin <10: 50/299 (16.7%) vs. 50/311 (16.1%); p=0.916 Thrombocytopenia platelets <50: 3/299 (1.0%) vs. 4/311 (1.3%); p=1.0 Pyrexia: 68/299 (22.7%) vs. 80/311 (25.7); p=0.445 Myalgia: 114/299 (38.1%) vs. 108/311 (34.7%); p=0.430 Rash: 34/299 (11.4%) vs. 58/311 (18.6%); p=0.016 Fatigue: 131/299 (43.8%) vs. 156/311 (50.2%); p=0.136 Headache: 30/299 (10.0%) vs. 47/311 (15.1%); p=0.077 Insomnia: 47/299 (15.7%) vs. 51/311 (16.4%); p=0.906 Depression: 55/299 (18.4%) vs. 70/311 (22.5%); p=0.247 Nausea: 37/299 (12.4%) vs. 40/311 (12.9%); p=0.953	Schering Plough

Author, Year Country Study Name Quality	Interferon Regimen	Ribavirin Regimen	Protease Inhibitor Regimen	Eligibility	Exclusion	Number Screened/ Eligible/ Enrolled/ Analyzed	Sex	Genotype Severity of Liver Disease Proportion Treatment-Naïve
Bronowicki,	A. Pegylated	All patients treated	NA	Treatment naïve	chronic liver disease of other	NR/516/	A vs. B	A vs. B
2006^{46}	interferon alfa-	for 24 weeks of		Aged ≥18 years	etiology	349/349		Genotype 1: all
France	2a 180	ribavirin 400 mg		HCV genotype 1	Evidence of decompensation		44.2 vs. 45.4	HCV RNA>800,000: 62% vs.
	mcg/week for	twice daily. At		infection	Coinfection with HBV or HIV		Female: 43%	71%
Effect of ribavirin	48 weeks	week 24 patients		HCV RNA >600 IU/mL	Neutrophils <1500/mm3		vs. 43%	Fibrosis score F3 or F4: 27% vs.
in genotype 1	B. Pegylated	with indictable		Increased ALT levels	platelets <90,000/mm3		Non White:	28%
patients with	interferon alfa-	HCV RNA were		documented 2 times in	Hemoglobin level less than 12		NR	
hepatitis C	2a 180	randomized at		last 6 months	g/dL (women) or less than 13			
responding to	mcg/week for	week 26 to 22		Liver biopsy consistent	g/dL (men)			
pegylated	48 weeks	more weeks (48		with chronic hepatitis C	Risk factor for anemia			
interferon alfa-2a		weeks total) of:		obtained within 18	Serum creatinine >1.5 times			
plus ribavirin		A. 400 mg twice		months before therapy	upper limit of number			
		daily			Severe psychiatric disease			
Overall Quality:		B. Placebo			Significant comorbid medical			
Fair					conditions			

Author, Year Country Study Name	Duration of				Histologic		Funding
Quality	Followup	Outcome	Subgroup Analyses	Subgroup Analyses	Response	Adverse Events	Source
Bronowicki,	24 weeks	A vs. B	NR	NR	NR	A vs. B	Roche
2006^{46}	following	SVR: 93/176				Overall withdrawals: NR	
France	treatment	(52.8%) vs. 118/173				Withdrawals for adverse events: 3/173 (1.7%) vs.	
	completion	(68.2%); p=0.004				4/176 (2.3%); p=NS	
Continued						Serious adverse events: 13/173 (7.5%) vs. 12/176	
		Hepatitis Quality of				(6.8%); p=NS	
		Life Questionnaire:				Deaths: 1/173 (0.5%) vs. 0/176 (0%); p=NS	
		Scores for all				Asthenia: 19/173 (10.6%) vs. 13/176 (7.3%); p=NS	
		domains not				Headache: 7/173 (3.9%) vs. 6/176 (3.4%); p=NS	
		significantly different				Depression: 13/173 (7.5%) vs. 16/176 (9.1%); p=NS	
		between two				Myalgia: 6/173 (3.4%) vs. 6/176 (3.4%); p=NS	
		treatment regimens at				Leukopenia: 5/173 (2.8%) vs. 5/176 (2.8%); p=NS	
		any point in time					

			1			Number		
Author, Year						Screened/		
			Protease				A	Compatents
Country	Interferon	Ribavirin	Inhibitor			Eligible/ Enrolled/	Age Sex	Genotype
Study Name				E11 11 1114	T. 1 .			Severity of Liver Disease
Quality	Regimen	Regimen	Regimen	Eligibility	Exclusion	Analyzed	Race	Proportion Treatment-Naïve
Ferenci, 2008 ⁴⁹	A: Pegylated	A: Ribavirin	None	Treatment-naive adult	Pregnant or breast-feeding	291/282/	A vs. B	A vs. B
Austria	interferon	800 mg/day/24		Aged 18 to 65 years	women and male partners of	250/250	Age (Mean):	Genotype 2 – 18/141(13%) vs.
	alpha-2a 180	weeks		Chronic hepatitis C	pregnant women		37 vs. 36	19/141(14%)
A Randomized,	μg/week/24	B: Ribavirin		HCV genotype 2 or 3	Received prior treatment with		years	Genotype 3 - 123/141(87%) vs.
Prospective Trial	weeks	400 mg/day/24		infection	interferon or ribavirin at any time			122/141(86%)
of Ribavirin 400	B: Pegylated	weeks		-			Female: 40%	
mg/Day Vs. 800	interferon			in serum and elevated	or human immunodeficiency		vs. 38%	Severity of liver disease-
mg/Day in	alpha-2a 180			serum ALT activity (1.5	virus			HCV RNA < 800,000 IU/mL - 5.9
Combination with	μg/week/24			times the upper limit of	Decompensated liver disease or		Race: NR	vs. 5.7
Pegylated	weeks			normal [ULN] in the	chronic liver disease attributable			Cirrhosis: NR
interferon Alfa-2a				previous 6 months and	to another cause			Minimal or no fibrosis: NR
in Hepatitis C				during screening)	Coronary heart disease			100% Treatment naïve
Virus Genotypes				Hemoglobin value 12	Diabetes mellitus requiring			
2 and 3				g/dL (women) or 13	insulin therapy			
				g/dL (men)	Autoimmune disorders			
Overall Quality:				Leukocyte count 3000/	Any other unstable chronic			
Fair				L	medical condition			
				Platelet count 100,000/	Severe psychiatric disease,			
				L	especially depression			
				Serum creatinine level	History of active alcohol or drug			
				1.5 times the ULN.	addiction within the previous 6			
				Women of childbearing	months			
				potential were required				
				to have a negative	*Patients on opiate substitution			
				pregnancy test within	therapy were eligible if they were			
				24 hours of the first	treated by the drug treatment			
				dose All fertile male	centre in the Department of			
				and female participants	Psychiatry, Medical University			
				were required to use	of Vienna			
				two forms of effective				
				contraception during				
				treatment and for 6				
				months after the end of				
				treatment				

Author, Year Country Study Name Quality	Duration of Followup	Outcome	Subgroup Analyses	Subgroup Analyses	Histologic Response	Adverse Events	Funding Source
Ferenci, 2008 ⁴⁹	Followup visits at 24	A vs. B ETR: NR	NR	A vs. B SVR:	NR	A vs. B	Roche,
Austria Continued	weeks after	SVR: 97/141(68.8%)		Genotype 2 - 14/18(77.8%) vs.		Overall withdrawals: 13/141 (9%) vs. 22/141 (16%) p=NS Withdrawals due to adverse events: NR	Austria
Continued	of	vs. 90/141(63.8%)		12/16(63.2%); p=NS		Deaths: NR	
	treatment			Genotype 3 - 83/12(67.5%) vs.		Severe Adverse Events: NR	
				78/122(63.9%); p=NS		Adverse events:	
						Pruritus: 48/141 (34%) vs. 50/141 (35%); p=NS Psychiatric events (mostly depression): 49/141	
						(35%) vs. 56/141 (40%); p=NS	
						Hemoglobin <8.5 g/dL: 2/141 (1.4%) vs. 1/141	
						(0.7%); p=NS Neutrophils <1000/mm3: 73/141 (52%) vs. 71/141	
						(50%); p=NS	
						Platelets <50K/mm3: 6/141(4%) vs. 6/141 (4%); p=NS	

Fried., 2008 W List of the providing of the parties	Author, Year Country Study Name Quality	Interferon Regimen	Ribavirin Regimen	Protease Inhibitor Regimen	Eligibility	Exclusion	Number Screened/ Eligible/ Enrolled/ Analyzed	Age Sex Race	Genotype Severity of Liver Disease Proportion Treatment-Naïve
Improved 2a 180 B. Pispaterin 1600 mg/day/48 weeks Meanth 15 Mean	Fried., 2008 ⁵⁰								-
	Fried., 2008 ⁵⁰ USA Improved Outcomes in Patients with Hepatitis C with Difficult-to-Treat Characteristics: Randomized Study of Higher Doses of Pegylated interferon ά-2a and Ribavirin Overall Quality:	A: Pegylated interferon alfa-2a 180 μg/week/48 weeks B: Pegylated interferon alfa-2a 180 μg/week/48 weeks C: Pegylated interferon alfa-2a 270 μg/week/48 weeks D: Pegylated interferon alfa-2a 270 μg/week/48	A: Ribavirin 1200 mg/day/48 weeks B: Ribavirin 1600 mg/day/48 weeks C: Ribavirin 1200 mg/day/48 weeks D: Ribavirin 1600		Treatment-naïve Age 18 years or older Weighing 85 kg Chronic hepatitis C infection with genotype 1 Baseline HCV RNA level 800,000 IU/mL determined by quantitative polymerase chain reaction (PCR) assay Positive anti- HCV antibody test Elevated serum alanine aminotransferase level within the previous 6 months Compensated liver disease Liver biopsy specimen consistent with chronic hepatitis C obtained within the previous 24	Infection with an HCV genotype other than 1 Previous treatment with interferon-based therapy, ribavirin, or any investigational drug for chronic hepatitis C History or other evidence of liver disease not associated with chronic hepatitis C Neutrophil count 1.5 x 10^9 cells/L Platelet count 90 109 cells/L Hemoglobin level 12 g/dL in women and 13 g/dL in men Increased risk of anemia or for whom anemia would be medically problematic Serum creatinine level more than 1.5 times the upper limit of normal Co infection with hepatitis B virus or human immunodeficiency virus Other serious chronic disease History of severe psychiatric disease (a history of a suicide attempt, hospitalization or period of disability due to psychiatric disease, and/or a Beck Depression Inventory score 20) Evidence of alcohol or drug	301/193/	A vs. B vs. C vs. D Age (Mean): 47.1 vs. 49.6 vs. 47.1 vs. 48.5 years Female: 20% vs. 13% vs. 26% vs. 21% Race: White - 70% vs. 62% vs. 74% vs. 68% Non White- 30% vs. 38% vs. 26% vs.	A vs. B vs. C vs. D Genotype 1 – 100% Histologic diagnosis: Non cirrhotic -83% vs. 81% vs. 83% vs. 81% Cirrhosis - 17% vs. 19% vs. 17% vs. 19% HCV RNA (IU/mLx106): 4.9 vs. 6.2 vs. 5.5 vs. 5.2

Author, Year Country Study Name Quality	Duration of Followup	Outcome	Subgroup Analyses	Subgroup Analyses	Histologic Response	Adverse Events	Funding Source
Fried, 2008 ⁵⁰	Followup	A vs. B vs. C vs. D	NR	NR	NR	A vs. B vs. C vs. D	Hoffman La
USA	visits at 24	ETR: 21/46(45.7%)				Overall withdrawals: 13/46(28%) vs. 9/47(19%) vs.	Roche
	weeks after	vs. 27/47(57.4%) vs.				15/47(32%) vs. 17/47(36%)	
Continued	completion	26/47(55.3%) vs.				Withdrawals for adverse events: 5/46(11%) vs.	
	of treatment	26/47(55.3%)				1/47(2%) vs. 7/47(15%) vs. 9/47(19%)	
						Deaths: NR	
		SVR: 13/46(28.3%)				Serious Adverse Events: 4/46(9%) vs. 6/47(13%) vs.	
		vs. 15/47(31.9%) vs.				6/47(13%) vs. 5/47(11%)	
		17/47(36.2%) vs.				Adverse events: (significant p-values noted for A vs. B,	
		22/47(46.8%)				A vs. C, or C vs. D)	
						Fatigue - 36/46(78%) vs. 32/47(68%) vs. 35/47(74%)	
						vs. 34/47(72%)	
						Headache - 24/46(52%) vs. 18/47(38%) vs. 22/47(47%) vs. 21/47(45%)	
						Insomnia - 18/46(39%) vs. 20/47(43%) vs. 22/47(47%) vs. 24/47(51%)	
						Nausea - 18/46(39%) vs. 20/47(43%) vs. 18/47(38%)	
						vs. 18/47(38%)	
						Chills - 15/46(33%) vs. 14/47(30%) vs. 19/47(40%) vs.	
						17/47(36%)	
						Myalgia - 14/46(30%) vs. 16/47(34%) vs. 19/47(40%)	
						vs. 16/47(34%)	
						Depression - 14/46 (30%) vs. 20/47(43%) vs.	
						12/47(26%) vs. 16/47(34%)	
						Arthralgia - 13/46(28%) vs. 16/47(34%) vs.	
						16/47(34%) vs. 15/47(32%)	
						Irritability - 14/46(30%) vs. 14/47(30%) vs.	
						12/47(26%) vs. 16/47(34%)	
						Pyrexia - 12/46(26%) vs. 14/47(30%) vs. 16/47(34%)	
						vs. 14/47(30%)	
						Rash - 12/46(26%) vs. 11/47(23%) vs. 15/47(32%) vs.	
						12/47(26%)	
						Diarrhea - 12/46(26%) vs. 9/47(19%) vs. 11/47(23%)	
						vs. 10/47(21%)	
						Cough - 9/46(20%) vs. 12/47(26%) vs. 12/47(26%) vs.	
						8/47(17%)	
						Dyspnea - 9/46(20%) vs. 12/47(26%) vs. 8/47(17%) vs. 12/47(26%)	
						Dizziness - 12/46(26%) vs. 9/47(19%) vs. 7/47(15%) vs. 9/47(19%)	
						Back pain - 1/46(2%) vs. 11/47(23%) vs. 4/47(9%) vs.	
						3/47(6%); (B vs. D p=0.02)	
						Injection site erythema - 10/46(22%) vs. 9/47(19%) vs.	
						6/47(13%) vs. 5/47(11%)	

						Number		
Author, Year						Screened/		
/			Protease				Age	Genotype
•	Interferon							
Ouality		Ribavirin Regimen		Eligibility	Exclusion		Race	
Country Study Name Quality Helbling, 2006 ⁵¹ Switzerland HCV-related advanced fibrosis/cirrhosis: randomized controlled trial of pegylated interferon α-2a and ribavirin Overall Quality: Fair	Interferon Regimen A: Pegylated interferon alpha-2a 180 μg/week/48 weeks B: Pegylated interferon alpha-2a 180 μg/week/48 weeks	Ribavirin Regimen A: (standard dose)Ribavirin <75 kg - 1000 mg/day/48 weeks >75 kg - 1200 mg/day in 2 divided doses/48 weeks B: (low dose) Ribavirin <75 kg - 600 mg/day/48 weeks >75 kg - 800 mg/day/48 weeks >75 kg - 800 mg/day in 2 divided doses/48 weeks	Protease Inhibitor Regimen None	Eligibility Age 18–70 years Biopsy proved (within ≤12 months) chronic hepatitis C with advanced fibrosis/cirrhosis (Ishak stage F4–F6 <7 Child–Pugh points No previous antiviral treatment Elevated alanine aminotransferase (ALT; on ≥2 occasions within >6 months) Serum HCV RNA positive Hemoglobin ≥11 g/dL Neutrophil count >1500/IL Platelet count ≥75 000/IL Serum creatinine ≤1.5 times upper limit of normal Normal fasting glucose	Exclusion Concomitant liver disease Ongoing substance abuse including alcohol (≥80 g/day) Hepatocellular carcinoma Clinically relevant disorders of other organs/systems Pregnancy or lactation Refusal to practice effective contraception during treatment/followup Immunomodulatory treatment within 6 months or treatment with any investigational drug within 30 days of study entry	Eligible/ Enrolled/ Analyzed NR/126/ 126/124	Age Sex Race A vs. B Age - Median: 47 vs. 47 years Female: 30% vs. 40% Race: NR	Genotype Severity of Liver Disease Proportion Treatment-Naïve A vs. B Genotype 1 – 30/64(47%) vs. 25/60(42%) Genotype 2 – 11/64(17%) vs. 7/60(12%) Genotype 3 - 18/64(28%) vs. 24/60(40%) Genotype 4 - 4/64(6%) vs. 3/60(4%) Histologic stage (Ishak): 3 - 3/64(5%) vs. 4/60(7%) 4 - 26/64(41%) vs. 18/60(30%) 5 - 19/64(30%) vs. 21/60(35%) 6 - 14/64(22%) vs. 13/60(22%) Cirrhosis: 57% vs. 52% Minimal or no fibrosis: 6% vs. 2% 100% Treatment naïve
				(or ≤8 µmol/L provided HbA1c ≤8.5%) Hbs-antigen negative antinuclear antibodies ≤1:160 Normal thyroid stimulating hormone Normal alpha-fetoprotein Focal lesions ruled out by ultrasound (within 1 month of study entry)				

Author, Year Country Study Name Quality	Duration of Followup	Outcome	Subgroup Analyses	Subgroup Analyses	Histologic Response	Adverse Events	Funding Source
Helbling, 2006 ⁵¹ Switzerland	Followup visits at 24 weeks post-	A vs. B ETR: NR	NR	A vs. B ETR: NR	NR	A vs. B Discontinuation: 15/64 (23%) vs. 16/60 (27%); p=NS	NR
Continued	treatment	SVR: 33/64(52%) vs. 23/60(38%), p=0.153		SVR: Fibrosis (Ishak): F4 - 15/26(58%) vs. 6/18(33%) F5-6 - 14/33(42%) vs. 14/34(41%) Genotype 1/4 - 11/34(32%) vs. 9/28(32%) Genotype 2/3 - 21/29(72%) vs. 14/31(45%)		Discontinuation (due to AE): 6/64(9%) vs. 9/60(15%); p=NS Overall withdrawals: 18/64(28%) vs. 23/60(38%); p=NS Deaths: 0/64(0%) vs. 2/60(3%); p=NS Severe Adverse Events: 9/64(14%) vs. 11/60(18%); p=NS Adverse events: Psychiatric - 1/64(2%) vs. 4/60(7%); p=NS Neurologic - 3/64 (5%) vs. 1/60(2%); p=NS Infectious - 1/64(2%) vs. 2/60(3%); p=NS Neoplastic - 2/64 (3%) vs. 1/60(2%); p=NS Skin - 0/64(0%) vs. 1/60(2%); p=NS Endocrine and Metabolism - 0/64(0%) vs. 1/60(2%); p=NS Eye - 1/64(2%) vs. 0/60(0%); p=NS Gastrointestinal - 0/64(0%) vs. 1/60(2%); p=NS Cardiovascular - 1/64(2%) vs. 0/60(0%); p=NS	

	1	1			1		1	1
						Number		
Author, Year						Screened/		
Country			Protease			Eligible/	Age	Genotype
Study Name	Interferon	Ribavirin	Inhibitor			Enrolled/	Sex	Severity of Liver Disease
Quality	Regimen	Regimen	Regimen	Eligibility	Exclusion	Analyzed	Race	Proportion Treatment-Naïve
Jacobson, 2007 ⁵²	A: Pegylated	A: Ribavirin 800	None	Treatment-naive	Positive test result for hepatitis B	Paper 1: NR/	A vs. B	A vs. B
USA (236	interferon alfa-	mg/day 24- 48		chronic hepatitis C	surface antigen or human	NR/ 5519/	Age - Mean:	Genotype 1 - 1512/2469 (61.2%)
practice sites	2b 1.5 μg/kg	weeks depending		patients	immunodeficiency virus (HIV)	4913	- 45.8 vs.	vs. 1506/2444 (61.6%)
nation-wide)	1x/week/24 -	on genotype		18 to 70 years old			45.8 years	Genotype 2 - 499/2469 (20.2%)
	48 weeks	B: Ribavirin 800-		Body weight less than		Paper 2:		vs. 525/2444 (21.5%)
Pegylated	depending on	1400 mg/day for		125 kg		4913/387/	Female -	Genotype 3 - 421/2469 (17.1%)
interferon alfa-2b	genotype	24-48 weeks		Treatment-naive adult		387/387	37.7% vs.	vs. 386/2444 (15.8%)
and Weight-	B: Pegylated	depending on		patients with HCV		(sub	36.2%	Genotype 4/5/6 - 33/2469 (1.3%)
Based or Flat-	interferon alfa-	genotype		RNA levels detectable		population		vs. 23/2444 (0.9%)
Dose Ribavirin in	2b 1.5 μg/kg			by (PCR)/branched		from	Race:	Genotype viral load >600,000
Chronic Hepatitis	1x/week/24 -	<65kg - Ribavirin		DNA assay		Jacobson,	White -	IU/mL - 1232/2469 (49.9%) vs.
C Patients: A	48 weeks	800 mg/week/48		Compensated liver		2007a)	80.7% vs.	1125/2444 (46.0%)
Randomized Trial	depending on	weeks		disease			78.8%	METAVIR stage:
	genotype	65-85 kg -		Liver biopsy showing			Non White -	F0-F2 - 1729/2469 (70.0%) vs.
Jacobson, 2007 ⁵³		Ribavirin 1000		HCV infection within			19.3% vs.	1709/2444 (69.9%)
(African-		mg/week/48 weeks		36 months prior to			21.2%	F3 - 486/2469 (19.7%) vs.
American sub-		>85-105 kg -		screening				489/2444 (20.0%)
group)		Ribavirin 1200		Elevated ALT at least			Paper 2:	F4 - 254/2469 (10.3%) vs.
USA (236		mg/week/48 weeks		once during the 6			Race:	246/2444 (10.1%)
practice sites		>105 kg but <125		months prior to			100% Non	ALT abnormal: 2119/2469
nation-wide)		kg - Ribavirin		screening			White	(85.8%) vs. 2105/2444 (86.1%)
		1400 mg/week/48		Alpha-fetoprotein level			(African-	HCV viral load (> 600,000
Impact of Weight-		weeks		of $\leq 100 \text{ ng/mL}$ in the			American)	IU/mL): 1232/2469(49.9%) vs.
based Ribavirin				year preceding entry				1125/2444(46%)
with Pegylated								100% Treatment naive
interferon alfa-2b								Paper 2: (African-Americans)
in African-								Genotype 1: 100%
Americans with								HCV viral load > 600,000 IU/mL
Hepatitis C Virus								- 119/202(59%) vs. 116/185(63%)
Genotype 1								METAVIR stage F3-F4 (%) -
								60/202(30%) vs. 58/185(31%)
Overall Quality:								Cirrhosis: 10% vs. 10%
Fair								Minimal or no fibrosis: NR
								100% Treatment naïve

Author, Year							
Country	Duration						
Study Name	of				Histologic		Funding
Quality	Followup	Outcome	Subgroup Analyses	Subgroup Analyses	Response	Adverse Events	Source

Author, Year Country Study Name Quality	Duration of Followup	Outcome	Subgroup Analyses	Subgroup Analyses	Histologic Response	Adverse Events	Funding Source
Jacobson, 2007 ⁵² USA (236 practice sites nation-wide) Jacobson, 2007 ⁵³ (African- American subgroup) USA (236 practice sites nation-wide) Continued	Followup visits at 24 weeks after completion of treatment	A vs. B ETR: 1193/2102(56.8%) vs. 1255/2121(59.2%), p= 0.082 SVR: 852/2102(40.5%) vs. 938/2121(44.2%), p=0.010	A vs. B 65-85 kg: 43.8% vs. 45.2% 85-105 kg: 38.8% vs. 42% >105 kg: 33.5% vs. 47.3% African-Americans Genotype 1: 19/188(10.1%) vs. 36/174(20.7%), p=0.006	A vs. B Genotype1: 337/1305 (29%) vs. 447/1313 (34%); p=0.005 Genotype 2/3: 462/777 (60%) vs. 479/775 (62%); p=0.252 Genotype 1 High Viral Load - 199/744(26.7%) vs. 246/789(31.2%), p=0.056 Genotype 1 Low Viral Load - 149/427(34.9%) vs. 151/381(39.6%); p=0.164	NR	A vs. B Discontinuation: 354/2444(14.5%) vs. 369/2469(14.9%); p=NS Overall withdrawals: 913/2444(37.3%) vs. 895/2469(36.2%); p=NS Death: 5/2444(<1%) vs. 9/2469(<1%); p=NS Serious Adverse Event: 279/2444(11.4%) vs. 287/2469(11.6%); p=NS Adverse events: Cardiovascular – 136/2444(5.6%) vs.162/2469(6.6%); p=NS Psychiatric - 1685/2444(68.9%) vs. 1667/2469(67.5%); p=NS Anemia - 473/2444(19.4%) vs. 721/2469(29.2%); p<0.001 Paper 2 (African Americans): Discontinuation: 85/202(42%) vs. 68/165(41%); p=NS Overall withdrawals: 35/202(17%) vs. 30/165(18%); p=NS Deaths: NR Severe Adverse Events: NR Adverse events: Nadir hemoglobin- <10 g/dL - 30/202(15%) vs. 37/185(20%); p=NS <8.5 g/dL - 2/202(1%) vs. 8/185(4%); p=0.04 RBV dose-reduction - 53/202(26%) vs. 69/185(37%);p=0.02 Nadir Absolute Neutrophil Count- <750 cells/mm3 - 56/202(28%) vs. 44/185(24%); p=NS <500 cells/mm3 - 10/202(5%) vs. 15/185(8%); p=NS Nadir platelets: <100 x 103 cells/mm3 - 30/202(15%) vs. 21/185(11%); p=NS <50 x 103 cells/mm3 - 2/202(1%) vs. 2/185(1%); p=NS	Schering- Plough Corp., Kenilworth, NJ

Quality Kawaoka, 2009 ⁵⁴	Interferon Regimen A: Pegylated interferon alpha-2a 1.0	Ribavirin Regimen A: Ribavirin 60 kg - 600 mg/week/24 weeks	Protease Inhibitor Regimen None	Eligibility Patients with chronic hepatitis C Age >20 years	Exclusion Patients treated with Shosaiko-to, a Japanese herbal medicine considered to improve	Number Screened/ Eligible/ Enrolled/ Analyzed NR/ 55/ 53/ 53	Sex	Genotype Severity of Liver Disease Proportion Treatment-Naïve A vs. B Genotype 2a: 13/26(50%) vs. 13/27(48%)
Dose comparison study of pegylated interferon-α-2b plus ribavirin in naïve Japanese	μg/kg/week/24 weeks B: Pegylated interferon alpha-2a 1.5 μg/kg/week/24 weeks	>60 kg-≤80 kg - 800 mg/week/24 weeks >80 kg - 1000 mg/week/24 weeks B: Ribavirin 60 kg - 600 mg/week/24 weeks >60 kg-<80 kg - 800 mg/week/24 weeks >80 kg - 1000 mg/week/24 weeks		Treatment naïve Genotype 2	liver function Patients with autoimmune hepatitis Patients with a history of hypersensitivity to Pegylated Interferon-alpha-2a or other interferons History of hypersensitivity to biological products, such as vaccines Decompensated liver cirrhosis (LC) Hepatocellular carcinoma (HCC) or malignant tumors in other tissues History of severe psychosis, such as being severely depressed and/or suicidal Women who were pregnant or lactating or who were suspected of being pregnant Patients judged by the investigator not to be appropriate for inclusion		Female: 65% vs. 44% Race: NR (study	Genotype 2b: 13/26(50%) vs. 14/27(52%) Histological stage (Desmet): F0 - 1/26(4%) vs. 0/27(0%) F1 - 14/26(51%) vs. 13/27(48%) F2 - 8/26(31%) vs. 9/27(33%) F3 - 3/26(12 %%) vs. 5/27(19%) Cirrhosis: None Minimal or no fibrosis: 55% vs. 48% 100% Treatment naive

Author, Year Country Study Name Quality	Duration of Followup	Outcome	Subgroup Analyses	Subgroup Analyses	Histologic Response	Adverse Events	Funding Source
Kawaoka, 2009 ⁵⁴ Japan Continued	24 weeks following treatment completion	A vs. B ETR: 23/26(88.5%) vs. 25/27(92.6%), p=0.13 SVR: 10/26(38.5%) vs. 20/27(74.1%), p=0.013	NR	NR	NR	A vs. B Overall withdrawals/drop-out: 2/26(7.2%) vs. 2/27(7.6%); p=NS Discontinuation (pre-mature withdrawal of treatment due to AE): 3/26(11.5%) vs. 2/27(7.4%); p=NS Depression - 1/26(3.8%) vs. 0/27(0%); p=NS Fatigue - 1/26(3.8%) vs. 1/27(4%); p=NS Excitability - 0/26(0%) vs. 1/27(4%); p=NS Deaths: NR Severe Adverse Events: NR Adverse events (leading to dose-reduction): Thrombocytopenia - 1/26(4%) vs. 0/27(0%); p=NS Fatigue - 1/26(4%) vs. 3/27(11%); p=NS Neutropenia - 0/26(0%) vs. 1/27(4%); p=NS Anemia - 15/26 (57.7%) vs. 10/27 (37%); p=NS Reduced Ribavirin - 21/26 (80.7%) vs. 22/27(81.5%); p=NS	NR

Author, Year Country Study Name Ouality	Interferon Regimen	Ribavirin Regimen	Protease Inhibitor Regimen	Eligibility	Exclusion	Number Screened/ Eligible/ Enrolled/ Analyzed	Age Sex Race	Genotype Severity of Liver Disease Proportion Treatment-Naïve
Krawitt, 2006 ⁵⁵	A: (low dose)	A: Ribavirin 1000	None	Age \geq 18 years older	Positive serum hepatitis B	NR/NR/	A vs. B	A vs. B
USA (New	Pegylated	mg/day/24 weeks		Detectable serum		314/301	Age:	Genotype 1 - 109/152(71.7%) vs.
York/New	interferon	(treatment		hepatitis C virus (HCV)	Any chronic liver disease other		. 50	119/162(73.5%)
England)	alpha-2b 50 µg/week/24	continued for additional 24		RNA Treatment naive	than chronic hepatitis C Hemoglobinopathies		> 50 years - 18% vs. 19%	Genotype 2/3 - 43/152(28.3%) vs. 43/162(26.5%)
A Study of Low	μg/week/24 weeks	weeks if HCV		Liver biopsy consistent	Evidence of hepatic		18% VS. 19%	43/102(20.3%)
Dose Pegylated	(treatment	RNA undetectable		with the diagnosis of	decompensation(ascites,		Female -	Histology
interferon Alpha-	continued for	by PCR at week		chronic hepatitis C,	encephalopathy, gastrointestinal			Fibrosis - 80/152(52.6%) vs.
2b with Ribavirin	additional 24	24)		performed not longer	bleeding secondary to portal			92/162(56.8%)
for the Initial	weeks if HCV	B: Ribavirin 1000		than 5 yr prior to entry,	hypertension)		Race:	Cirrhosis - 26/152(17.1%) vs.
Treatment of	RNA	mg/day/24 weeks		with histological	Other conditions that could			17/162(10.5%)
Chronic Hepatitis	,	(treatment		interpretation	interfere with participation in the		Non White -	
C	PCR at week	continued for		performed by	protocol - (i.e. coronary artery		4.6% vs.	Baseline HCV RNA:
011 01:4	24)	additional 24 weeks if HCV		pathologists at the study site locations	disease, uncontrolled		3.1%	$\leq 2 \times 10^6 \text{ copies/ml} - 67/152(44.1\%) \text{ vs. } 86/162(40.7\%)$
Overall Quality: Fair	B: (standard	RNA undetectable		Chronic hepatitis alone	hypertension, clinically significant retinal abnormalities,			$> 2 \times 10^6 \text{ copies/ml}$ -
Tan	interferon	by PCR at week		(F0)	pregnancy, nursing, severe			85/152(55.9%) vs. 96/162(59.3%)
	alpha-2b	24)		Chronic hepatitis with	preexisting psychiatric disorders			03/132(33.570) 13: 30/102(33.570)
	<75 kg - 100	,		fibrosis, including	Active substance dependency			100% Treatment naive
	μg/week/24			bridging fibrosis (F1-	within 6 months of screening for			
	weeks			F3)	entry into the study			
	≥75kg - 150			Chronic hepatitis with	Methadone maintenance (unless			
	μg/week/24			cirrhosis (F4)	a program of continual testing			
	weeks				was in use)			
	(treatment continued for				History of organ transplantation Participation in any other clinical			
	additional 24				trial or use of another			
	weeks if HCV				investigational drug within 30			
	RNA				days of entry			
	undetectable by							
	PCR at week							
	24)							

Author, Year Country Study Name Quality	Duration of Followup	Outcome	Subgroup Analyses	Subgroup Analyses	Histologic Response	Adverse Events	Funding Source
Krawitt, 2006 ⁵⁵	Followup	A vs. B	A vs. B	A vs. B	NR	A vs. B	Integrated
USA (New	visits at 24	ETR: NR	ETR: NR	ETR: NR		Total Discontinuation: 9/147(6%) vs. 28/154(18%);	Therapeutics
York/New	weeks Post-					p=0.0015	Group
England)	treatment	SVR: 50/152(33%) vs. 73/162(45%), p=0.02	SVR: Age:	SVR: HCV Genotype:		Discontinuation due to AE: 5/147(3%) vs. 14/154(9%); p=0.04	(Schering- Plough)
Continued		73/162(45%), p=0.02	Age: ≤ 40 years - 13/33(39%) vs. 18/38(47%), p= 0.63 > 40 - ≥ 50 years - 28/91(31%) vs. 40/93(43%), p= 0.09 > 50 years - 9/28 (32%) vs. 15/31 (48%), p= 0.29 Male: 29/94 (31%) vs. 44/110 (40%); p=0.14 Female - 21/58(36%) vs. 29/52(56%), p=0.06 Race: Caucasian - 50/145 (34%) vs. 70/157 (45%), p= 0.08 African-American - 0/6 (0%) vs. 3/4 (75%), p= 0.03 Hispanic/Other - 0/1 (0%) vs. 0/1 (0%), p= 1.00 Weight: < 75 kg - 20/50 (40%) vs. 24/42 (57%), p= 0.14 ≥ 75 kg - 30/102 (29%) vs. 49/120	HCV Genotype: Genotype 1 - 26/109 (24%) vs. 45/119 (38%), p= 0.03 Genotype 2/3 - 24/43 (56%) vs. 28/43 (65%), p= 0.51 Baseline HCV RNA: ≤ 2×106 copies/ml - 19/67 (28%) vs. 37/66 (56%), p= 0.002 > 2×106 copies/ml - 31/85 (36%) vs. 36/96 (38%), p= 1.00 Histology: No fibrosis or cirrhosis: 17/46 (37%) vs. 29/53 (55%); p=0.11 Fibrosis - 27/80 (34%) vs. 39/92 (42%), p= 0.27 Cirrhosis - 6/26 (23%) vs. 5/17 (29%), p= 0.73		p=0.04 Overall withdrawals: NR Deaths: NR Severe Adverse Events: NR	Plough)

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1						Number		
Author, Year			_			Screened/		~
Country			Protease			Eligible/	Age	Genotype
Study Name	Interferon	Ribavirin	Inhibitor			Enrolled/	Sex	Severity of Liver Disease
Quality	Regimen	Regimen	Regimen	Eligibility	Exclusion	Analyzed	Race	Proportion Treatment-Naïve
Manns, 2001 ⁵⁶	A: Pegylated	A: (weight-based)	NA	Eligible patients were	Patients were excluded if they	NR/2316/153	A vs. B:	A vs. B
US & UK	interferon alfa-	Ribavirin 1000–		previously untreated		0/1530		Genotype 1: 68% vs. 68%
	2b 1·5 g/kg/4	1200 mg/day/48		adults who had HCV	serum-fetoprotein concentration		Age (Mean):	Genotype 2/3: 30% vs. 29%
Peginterferon	weeks	weeks		RNA detectable in	of more than 50 g/L, HIV		44 vs. 43	Genotype 4, 5, or 6: 2% vs.3%
alfa-2b plus	followed by	75 kg > 1000 mg		serum by PCR, who had	infection, previous organ		years	
ribavirin	Pegylated	75 kg < 1200 mg		undergone a liver	transplantation, other causes of		Female:	Histology
compared with	interferon 0.5	B: (weight-based)		biopsy within 1 year	liver disease, pre-existing		168/514(33	Mean (SD) baseline Knodell
interferon alfa-2b	g/kg/week/44	Ribavirin 1000–		before entry that was	psychiatric disease, seizure		%) vs.	inflammatory score: $7.9 (2.3)$ vs.
plus ribavirin for	weeks	1200 mg/day/48		consistent with chronic	disorders, cardiovascular disease,		169/505(33	7.8 (2.5)
initial treatment	B: interferon	weeks		hepatitis, and who had	hemoglobinopathies, hemophilia,		%)	Bridging fibrosis/cirrhosis:
of chronic	alfa-2b 3	75 kg > 1000 mg		high serum values of	poorly controlled diabetes, or		Race: NR	146/491(30%) vs. 132/468(28%)
hepatitis C: a	million	75 kg < 1200 mg		alanine	autoimmune type disease, or if			
randomized trial	units/3x			aminotransferase	they were unable to use			Treatment naive: 100%
	week/48 weeks			(above the upper limit	contraception.			
Overall Quality:				of normal >43 IU/L for				
Fair				men, >34 IU/L for				
				women) with minimum				
				hematological and				
				biochemical values of:				
				hemoglobin 120 g/L for				
				women and 130 g/L for				
				men; white-blood-cell				
				count 3 109/L;				
				neutrophil count 1.5				
				109/L; platelet count				
				100 109/L; and				
				bilirubin, albumin, and				
				creatinine within				
				normal limits.				

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Author, Year Country	Duration						
Study Name	of				Histologic		Funding
Quality	Followup	Outcome	Subgroup Analyses	Subgroup Analyses	Response	Adverse Events	Source
Manns, 2001 ⁵⁶	24 weeks	SVR:	NR	A vs. B vs. C:	NR	A vs B vs. C:	Schering
US & UK	following	333/511(65%) vs.	INIX	SVR:	INIX	Overall withdrawals: NR	Plough
US & UK	treatment	289/514(56%) vs.		Genotype 1: 42%		Withdrawals for adverse events: 42/511 vs.	Research
Continued	completion	271/505(54%),		(145/348) vs. 34%		36/514 vs. 34/505	Institute,
Continued	Completion	p<0.001 (A vs. B),		(143/349) vs. 33%		Serious adverse events: NR	Kenilworth.
		p=0.41 (A vs. C)		(114/343), p=0.02 (A		Deaths: NR	NJ, and
		p=0°+1 (/1 v3. 0)		vs. B), p=0.94(A vs		Bodins. WY	clinical
				C)		Adverse Events:	research
				Genotype 2/3: 82%		Anemia: 9/511 vs. 12/514 vs. 13/505	centre grants
				(121/147) vs. 80%		Neutropenia: 18/511 vs. 10/514 vs. 8/505	from
				(122/153) vs. 79%		Asthenia 18/511 vs. 16/514 vs. 18/505	Massachusetts
				(115/146), p=0·46(A		Fatigue 64/511 vs. 62/514 vs. 60/505	General
				vs. B), p=0.89 (A vs.		Fever 46/511 vs. 44/514 vs. 33/505	Hospital
				C)		Headache 62/511 vs. 58/514 vs. 58/505	(MO1-
				Genotype 4/5/6: 50%		Rigors 48/511 vs. 45/514 vs. 41/505	RR01066),
				(8/16) vs. 33% (4/12)		Weight decrease 29/511 vs. 17/514 vs. 20/505	Scripps Clinic
				vs. 38% (6/16),		Dizziness 21/511 vs. 21/514 vs. 17/505	(MO1-
				p=0.72 (À vs B),		Arthralgia 34/511 vs. 34/514 vs. 28/505	RR00833),
				p>0.99 (A vs. C)		Musculoskeletal pain 21/511 vs. 17/514 vs.	and University
				SVR by baseline		19/505	of Florida
				HCV:		Myalgia 56/511 vs. 48/514 vs. 50/505	(5MO1-
				>2 10 ⁶ /mL: 42%		Anorexia 32/511 vs. 29/514 vs. 27/505	RR00082).
				(149/351) vs. 42%		Diarrhea 22/511 vs. 16/514 vs. 17/505	
				(144/345) vs. 42%		Nausea 43/511 vs. 36/514 vs. 33/505	
				(145/344)		Vomiting 14/511 vs. 14/514 vs. 12/505	
				2 10 ⁶ /mL: 78%		Concentration impairment 17/511 vs. 16/514	
				(125/160) vs. 59%		vs. 21/505	
				(100/169) vs. 56%		Depression 31/511 vs. 29/514 vs. 34/505	
				(90/161)		Insomnia 40/511 vs. 40/514 vs. 41/505	
				SVR by degree of		Irritability 35/511 vs. 34/514 vs. 34/505	
				fibrosis:		Coughing 17/511 vs. 15/514 vs. 13/505	
				No/minimal fibrosis -		Dyspnea 26/511 vs. 23/514 vs. 24/505	
				57% (189/333) vs.		Alopecia 36/511 vs. 29/514 vs. 32/505	
				51% (175/345) vs.		Pruritus 29/511 vs. 26/514 vs. 28/505	
				49% (164/336)		Rash 24/511 vs. 22/514 vs. 23/505	
				Bridging		Dry skin 24/511 vs. 18/514 vs. 23/505	
				fibrosis/cirrhosis -		Injection-site inflammation 25/511 vs. 27/514	
				44% (60/136) vs.		vs. 18/505	
				43% (63/146) vs.		Injection-site reaction 58/511 vs. 59/514 vs.	
]		41% (54/132)		36/505	

						Number		
Author, Year						Screened/		
Country			Protease			Eligible/	1 00	Genotype
Study Name	Interferon	Ribavirin	Inhibitor			Enrolled/	Age Sex	Severity of Liver Disease
Ouality	Regimen	Regimen	Regimen	Eligibility	Exclusion	Analyzed	Race	Proportion Treatment-Naïve
C		A: Ribavirin				NR/NR/	1	A vs. B
Meyer-Wyss, 2006 ⁵⁷	A: Pegylated-		None	Treatment-naive	Subjects participating in any		A vs. B	1.75
Switzerland	interferon	800mg/day/24-48		patients	study within 30 days prior to	227/219	Age - Median: 39	Genotype 1 - 49/113(43%) vs.
Switzerland	alpha-2b 1.0	depending on		Aged 18–65 years	entry into the trial			64/106(60%)
	μg/kg/week/24	genotype		Biopsy-proven chronic	Pregnant or nursing women		vs. 42 years	Genotype 2 - 14/113(12%) vs.
Comparison of	-48 depending	B: Ribavirin		hepatitis C within ≤12	Positive human		E 1 420/	10/106(%)
two PEG-	on genotype	800mg/day/24-48		months	immunodeficiency virus		Female: 43%	Genotype 3 - 41/113(36%) vs.
interferon alpha-	B: Pegylated-	depending on		Up to moderate fibrosis	(HIV)status		vs. 28%	26/106(9%)
2b doses (1.0 or	interferon	genotype		(METAVIR score ≤F2)	Liver disease other than chronic		D ND	Genotype 4 - 9/113(8%) vs.
1.5μg/kg)	alpha-2b 1.5			with elevated alanine	hepatitis C		Race: NR	6/106(6%)
combined with	μg/kg/week/24			aminotransferase levels	Elevated levels of fasting blood			Histological state (METAVID
ribavirin in	-48 depending			(ALT; on at least two	glucose			Histological stage (METAVIR
interferon-naïve	on genotype			occasions, at least 6	Abnormal values of thyroid			score): 0 - 21/113(19%) vs. 13/106(12%)
patients with				months apart)	stimulating hormone			
chronic hepatitis				HCV-RNA positive	Hemophilia or			1 - 44/113(39%) vs. 39/106(37%)
C and up to moderate fibrosis				serum	Hemoglobinopathy			2 - 48/113(42%) vs. 54/106(51%)
moderate fibrosis					A 1			Cirrhosis: None
011 01:4					Any known pre-existing medical condition that could interfere			Cirriosis: None
Overall Quality:								Minimal of no fibrosis: NR
Poor					with the patient's participation			Millimat of no fibrosis: NK
					and completion of the study including:			100% Treatment naive
					History of severe psychiatric			100% Heatment harve
					disorders			
					Central nervous system			
					trauma/active seizure disorders			
					Significant cardiovascular			
					Pulmonary, or retinal disorders			
					Clinically manifested gout			
					Substance abuse			
					Chronic systemic administration			
					of steroids/other			
					immunosuppressants			
					Immunologically mediated			
					disease.			
	j				uisease.			

Author, Year Country Study Name Quality	Duration of Followup	Outcome	Subgroup Analyses	Subgroup Analyses	Histologic Response	Adverse Events	Funding Source
Meyer-Wyss , 2006 ⁵⁷ Switzerland Continued	Followup visits at 4 and 24 weeks post-treatment	A vs. B ETR: NR SVR: 61/113(53%) vs. 56/106(53%), p= ns	NR	A vs. B ETR: 17/39(49%) vs. 23/49(47%) SVR: Genotype 1/4: 22/58 (38%) vs. 27/70 (39%), p= ns Genotypes 2/3: 39/55 (71%) vs. 29 /36 (81%), p= ns >800K IU/mL: 28/48 (58%) vs. 40/69 (43%); p=NS <800 IU/mL: 34/65 (52%) vs. 40/69 (58%): p=NS	NR	A vs. B Discontinuation: 14/115(12%) vs. 28/112(25%); p=0.01 Deaths: 0/115(0%) vs. 1/112(0%); p=NS Life-threatening Adverse Events: 4/115(3%) vs. 9/112(9%); p=NS Severe Adverse Events: 62/115(54%) vs. 59/112(53%); p=NS Withdrawals due to AE: 22/115 (19%) vs. 34/112 (30%); p=0.05 Adverse events (only body systems listed with at least 10% of patients reporting): Thrombocytopenia: 1/115(1%) vs. 1/112(1%); p=NS Leukopenia: 9/115(8%) vs. 5/112(4%); p=NS Neutropenia: 20/115(17%) vs. 18/112(16%); p=NS Hemolytic anemia: 3/115(3%) vs. 3/112(3%); p=NS Blood and lymphatic system disorders - 44/115(38.3%)vs. 41/112 (36.6%); p=NS General disorders and administration site conditions - 112/115(97.4%) vs. 108/112(96.4%); p=NS Gastrointestinal disorders - 81/115(70.4%)vs. 84/112(75.0%); p=NS Metabolism and nutrition disorders -16/115(13.9%) vs. 29/112(25.9%); p=0.02 Musculoskeletal and connective tissue disorders - 27/115(23.5%) vs. 33/112(29.5%); p=NS Nervous system disorders - 70/115(60.9%) vs. 80/112(71.4%); p=NS Psychiatric disorders - 71/115(61.7%) vs. 76/112(67.9%); p=NS Respiratory, thoracic and mediastinal disorders 18/115(15.7%) vs. 24/112(21.4%); p=NS Skin and subcutaneous disorders - 83/115(72.2%) vs. 76/112(67.9%); p=NS	Essex Chemie AG, Lucerne

						Number		
Author, Year						Screened/		
Country			Protease			Eligible/	Age	Genotype
Study Name	Interferon	Ribavirin	Inhibitor			Enrolled/	Sex	Severity of Liver Disease
Quality	Regimen	Regimen	Regimen	Eligibility	Exclusion	Analyzed	Race	Proportion Treatment-Naïve

Mimidis, 2006 ⁵⁸	A. Pegylated	A. 800-1200 mg	NA	Treatment-naïve	HBV	NR/NR/	A vs. B	A vs. B
Greece	interferon alfa-	daily (11 mg/kg)		HCV RNA detected in	HIV coinfection	188/120	Age mean:	genotype 1/4: 46% vs. 52%
	2b 3.0 mcg/kg	B. 800-1200 mg		serum	Hemochromatosis		NR	Treatment-naïve: all
Hepatitis C virus	weekly for 12	daily (11 mg/kg)		Liver biopsy consistent	Alpha-1 anti-trypsin deficiency		Sex: 36% vs.	Fibrosis: NR
survival curve	weeks followed			with chronic hepatitis	Wilson's disease		38%	Cirrhosis: NR
analysis in naïve	by 1.5 mcg/kg			within 6 months before	Autoimmune hepatitis		non White:	HCV RNA≥ 800k IU/mL: NR
patients treated	weekly for 36			enrollment	Alcohol drug or obesity induced		NR	
with Pegylated	weeks			Elevated ALT at entry	liver disease			
interferon alpha-	B. Pegylated			and at least once in 6	Substance abuse			
2b plus ribavirin.	interferon alfa-			months before	Any known pre-existing			
A randomized	2b 1.5 mcg/kg			screening	condition that could interfere			
controlled trial for	weekly for 48				with patient's participation			
induction with	weeks				Creatinine >1.5 mg/dL			
high doses of					Neutrophils <1000/mL ³			
Pegylated					Platelets < 50K/mL ³			
interferon and					Hemoglobin <11 g/dL			
predictability of								
sustained viral								
response from								
early virologic								
data								
Overall Quality:								
Poor								

Author, Year Country Study Name Quality	Duration of Followup	Outcome	Subgroup Analyses	Subgroup Analyses	Histologic Response	Adverse Events	Funding Source
Mimidis, 2006 ⁵⁸	Week 72	A vs. B	NR	A vs. B	NA	NR	NR
Greece		ETR: NR		Genotype 1: 9/35			
				(25.7%) vs. 18/40			
Continued		SVR: 38/89 (42.7%)		(45%); p=NS			
		vs. 47/87 (54%)		Genotype 2/3: 23/48			
				(47.9%) vs. 25/42			
				(59.5%); p=NS			
				Genotype 4: 6/6 (100%)			
				vs. 4/5 (80%); p=NS			

Author, Year Country Study Name Quality	Interferon Regimen	Ribavirin Regimen	Protease Inhibitor Regimen	Eligibility	Exclusion	Number Screened/ Eligible/ Enrolled/ Analyzed	Age Sex Race	Genotype Severity of Liver Disease Proportion Treatment-Naïve
Reddy, 2010 ⁵⁹ International, 14 countries Induction pegylated interferon alfa-2a and high dose ribavirin do not increase SVR in heavy patients with HCV genotype 1 and high viral loads Overall Quality: Fair		A. 1400 - 1600 mg/day for 48 weeks depending on weight B. 1200 mg/day for 48 weeks C. 1400 - 1600 mg/day for 48 weeks depending on weight D. 1200 mg/day for 48 weeks	NA	Treatment-naïve Aged 18 years or older Weight ≥ 85 kg HCV genotype 1 infection HCV RNA ≥ 400k IU/mL Liver biopsy in past 24 months consistent with chronic hepatitis C	coinfection with HBV, HAV, or HIV Chronic liver disease of other origin Current or past history of chronic systemic disease including severe psychiatric disease Increased baseline risk of anemia Neutrophils <1500/mL³ Platelets <90K/mL³ Hemoglobin<12 g/dL in men or <13 g/dL in women Creatinine >1.5 times upper limit of normal Pregnant or breastfeeding women and male partners	NR/NR/ 1175/1145	A vs. B vs. C vs. D Age mean: 46 vs. 46 vs. 45 vs. 46 Female: 19% vs. 24% vs. 22% vs. 19% non White: 14% vs. 13% vs. 19% vs. 13%	A vs. B vs. C vs. D genotype 1: all Treatment-naïve: all Bridging fibrosis/cirrhosis: 12% vs. 8% vs. 10% vs. 12% HCV RNA ≥800k IU/mL: 86% vs. 83% vs. 84% vs. 82%

Author, Year Country Study Name Quality	Duration of Followup	Outcome	Subgroup Analyses	Subgroup Analyses	Histologic Response	Adverse Events	Funding Source
Reddy, 2010 ⁵⁹ International, 14 countries Continued	Week 72	A vs. B vs. C vs. D ETR: NR SVR: 156/383 (40.7%) vs. 166/382 (43.5%) vs. 81/189 (42.9%) vs. 72/191 (37.7%); (p=NS for all comparisons)	A vs. B vs. C vs. D (counts not reported) Weight <95 kg: 44% vs. 46% vs. 44% vs. 49% Weight ≥95 kg: 38% vs. 41% vs. 41% vs. 29%	A vs. B vs. C vs. D (counts not reported) Steatosis score <5%: 42% vs. 48% vs. 48% vs. 47% Steatosis score ≥5%: 36% vs. 30% vs. 32% vs. 13%	NA	A vs. B vs. C vs. D Overall withdrawals: 117/383 (31%) vs. 109/382 (29%) vs. 53/189 (28%) vs. 54/191 (28%); A vs. C p=NS; B vs. D p=NS Withdrawals for adverse events: 47/383 (12%) vs. 40/382 (10%) vs. 17/189 (9%) vs. 22/191 (12%); A vs. C p=NS; B vs. D p=NS Serious adverse events: 39/383 (10%) vs. 36/382 (9%) vs. 20/189 (11%) vs. 22/191 (12%); A vs. C p=NS; B vs. D p=NS Deaths: 2/383 (<1%) vs. 2/382 (<1%) vs. 3/189 (1%) vs. 1/191 (<1%); A vs. C p=NS; B vs. D p=NS Pyrexia: 205/383 (54%) vs. 176/382 (46%) vs. 78/189 (41%) vs. 83/191 (43%); A vs. C p=NS; B vs. D p=NS Fatigue: 182/383 (48%) vs. 185/382 (48%) vs. 102/189 (54%) vs. 66/191 (35%); A vs. C p=NS; B vs. D p=NS Headache: 168/383 (44%) vs. 152/382 (40%) vs. 76/189 (76%) vs. 75/191 (39%); A vs. C p=0.006; B vs. D p=0.002 Chills: 132/383 (34%) vs. 122/382 (32%) vs. 55/189 (29%) vs. 42/191 (22%); A vs. C p=NS; B vs. D p=0.001 Myalgia: 113/383 (30%) vs. 98/382 (26%) vs. 45/189 (24%) vs. 46/191 (24%); A vs. C p=NS; B vs. D p=NS Arthralgia: 89/383 (23%) vs. 88/382 (23%) vs. 49/189 (26%) vs. 50/191 (26%); A vs. C p=NS; B vs. D p=NS Depression: 58/383 (15%) vs. 72/382 (19%) vs. 36/189 (19%) vs. 32/191 (17%); A vs. C p=NS; B vs. D p=NS Hemoglobin <8.5 g/dL: 22/383 (6%) vs. 9/382 (2%) vs. 12/189 (6%) vs. 6/191 (3%); A vs. C p=NS; B vs. D p=NS	Roche

Author, Year Country	Duration						
Study Name	of	0.4	Calaman Analama	Colonia Analysis	Histologic	Alama Familia	Funding
Quality	Followup	Outcome	Subgroup Analyses	Subgroup Analyses	Response	Adverse Events	Source
Reddy, 2010 ⁵⁹						Neutrophils <500/mL3: 26/383 (7%) vs. 25/382	
International, 14						(7%) vs. 10/189 (5%) vs. 9/191 (5%); A vs. C p=NS;	
countries						B vs. D p=NS	
						Platelets < 20K/mL3: 3/383 (1%) vs. 0/382 (0%) vs.	
Continued						0/189 (0%) vs. 3/191 (2%); A vs. C p=NS; B vs. D	
U						p=NS	

Author, Year Country Study Name Quality	Interferon Regimen	Ribavirin Regimen	Protease Inhibitor Regimen	Eligibility		Number Screened/ Eligible/ Enrolled/ Analyzed	Age Sex Race	Genotype Severity of Liver Disease Proportion Treatment-Naïve
Roberts, 2009 ⁶⁰	A. Pegylated	A. 1000-1200	NA	Treatment naïve	HBV	NR/NR/	A vs. B	A vs. B
Australia	interferon alfa-	mg/day for 48		Ages 18 -75 years		896/871	Age mean:	genotype 1: all
	2a 360 mcg	weeks		HCV genotype 1	History of decompensated liver		44 vs. 43	Treatment-naïve: all
Impact of high-	weekly for 12	B. 1000-1200		infection	disease		Female: 31%	Fibrosis stage 3 or 4: 14% vs.
dose Pegylated	weeks followed			HCV RNA >600 IU/mL	Evidence of hepatocellular		vs. 35%	16%
interferon alfa-2a	by 180 mcg for	weeks		Elevated ALT	carcinoma		non White:	HCV RNA ≥ 800K: 70% vs. 67%
on virologic	36 weeks (48			Compensated liver	Liver disease of other origin		18% vs. 17%	
response rates in	weeks total)			disease (Child-Pugh	Therapy with systemic antiviral,			
patients with	B. Pegylated			score <7)	antineoplastic, or			
hepatitis C	interferon alfa-			Histologic findings	immunomodulatory agents			
genotype 1: a	2a 180 mcg			consistent with chronic	within 6 months			
randomized	weekly for 48			hepatitis on liver biopsy	Pregnancy or breast feeding and			
controlled trial	weeks			within last 36 months	male partner of women			
					Neutrophils <1500/mL ³			
Overall Quality:				*Protocol modified	Hemoglobin <12 g/dL in women			
Fair				during study to remove	and <13 g/dL in men			
				ALT, pretreatment	Creatinine >1.5 times the upper			
				biopsy, and	limit of normal			
				compensated cirrhosis	Active severe psychiatric disease			
				inclusion/exclusion	Any severe chronic or			
				requirements	uncontrolled disease			
					Current or recent drug or alcohol			
					abuse			
					Cirrhosis			

Author, Year Country Study Name Quality	Duration of Followup	Outcome	Subgroup Analyses	Subgroup Analyses	Histologic Response	Adverse Events	Funding Source
Roberts, 2009 ⁶⁰	24 weeks	A vs. B	A vs. B	A vs. B	NR	A vs. B	Roche
Australia	after end of	ETR: 70% vs. 66%;	White: 183/355	HCV RNA <800K:		Overall withdrawals: 113/433 (26%) vs. 136/438	
	treatment	p=0.18	(52%) vs. 167/365	81/125 (65%) vs.		(31%); p=NS	
Continued	(week 72)		(46%); p=NS	84/138 (61%); p=NS		Withdrawals due to adverse events: 44/433 (10%)	
		SVR: (230/433) 53%	Asian: 40/61 (66%)	HCV RNA ≥800K:		vs. 36/438 (8%); p=NS	
		vs. (219/438) 50%;	vs. 40/55 (73%);	147/302 (49%) vs.		Deaths: NR	
		p=0.29	p=NS	132/293 (45%); p=NS		Serious adverse events: 46/433 (11%) vs. 45/438	
			Other: 7/17 (41%) vs.	Eiler i METAVID		(10%); p=NS	
			12/18 (67%); p=NS	Fibrosis METAVIR stage 3 or 4: 17/60		Headache: 227/433 (52%) vs. 208/438 (47%); p=NS	
			Male: 149/298 (50%)	(28%) vs. 16/67 (24%);		Influenza like illness: 180/443 (42%) vs. 183/438	
			vs. 134/285 (47%);	p=NS		(42%); p=NS	
			p=NS	Fibrosis METAVIR		Nausea: 179/433 (41%) vs. 169/438 (39%); p=NS	
			Female: 81/135	stage 0,1,or 2: 148/256		Fatigue: 159/433 (37%) vs. 174/438 (40%); p=NS	
			(60%) vs. 85/153	(58%) vs. 134/242		Myalgia: 114/433 (26%) vs. 97/438 (22%); p=NS	
			(56%); p=NS	(55%); p=NS		Rash: 110/433 (25%) vs. 116/438 (26%); p=NS Depression: 84/433 (19%) vs. 85/438 (19%); p=NS	
			<40 years: 104/146			Arthralgia: 82/433 (19%) vs. 76/438 (17%); p=NS	
			(71%) vs. 97/141			Pyrexia: 66/433 (15%) vs. 47/438 (11%); p=NS	
			(69%); p=NS			Chills: 64/433 (15%) vs. 34/438 (8%); p<0.001	
			>40 years: 126/287			Neutropenia: 76/433 (21%) vs. 55/438 (13%);	
			(44%) vs. 122/297			p=0.05	
			(41%); p=NS			Thrombocytopenia: 17 (4%) vs. 6 (1%); p=0.02 Anemia: 5 (1%) vs. 3 (1%); p=NS	
			Weight <85 kg:			(, (, r	
			167/294 (57%) vs.				
			156/297 (53%);				
			p=NS				
			Weight >85 kg:				
			63/139 (45%) vs.				
			63/141 (45%); p=NS				

						Number		
Author, Year						Screened/		
Country			Protease			Eligible/	Age	Genotype
Study Name	Interferon	Ribavirin	Inhibitor			Enrolled/	Sex	Severity of Liver Disease
Quality	Regimen	Regimen	Regimen	Eligibility	Exclusion	Analyzed	Race	Proportion Treatment-Naïve
Sood, 2008 ⁶¹	A: Pegylated-	A: Ribavirin 10-12	None	Aged between 16-70-	Chronic HCV patients with	NR/103/	A vs. B	A vs. B
India	interferon	mg.kg/day/24		years-old	genotypes other than Genotype 3	103/103	Age - Mean:	Genotype 3: 100%
	alpha-2b 1.0	weeks		HCV-RNA positive	Total leukocyte count < 3000 per		43 vs. 37	
Comparison of	μg/kg/week/24	B: Ribavirin 10-12		with genotype 3	cubic millimeter		years	(Knodell)
low-dose	weeks	mg.kg/day/24		Treatment naïve	Platelet count < 70 000 per cubic			HAI score - Mean (SD): 7.2
pegylated	B: Pegylated-	weeks		ALT >1.2 x Upper limit				(3.15) vs. 4.68(2.12)
interferon vs.	interferon			of Normal (ULN) at	Hemoglobin level lower than 10		vs. 22%	Fibrosis score - Mean(SD):
standard high-	alpha-2B 1.5			screening and for at	g per deciliter			2.34(1.27) vs. 1.64(1.29)
dose pegylated	μg/kg/week/24			least the previous 6	co infection with hepatitis B		Race: NR	Cirrhosis: NR
interferon in	weeks			months	virus or human			
combination with				Liver biopsy-proven	immunodeficiency			100% Treatment naïve
ribavirin in				chronic HCV within 6	virus,			
patients with				months prior to	Alcohol intake exceeding 20			
chronic hepatitis				inclusion	g/day			
C with genotype					Presence of drug abuse,			
3: An Indian					psychiatric illness, or thyroid			
Experience					dysfunction			
					Pregnancy and lactation			
Overall Quality:					Decompensated liver disease			
Fair					Evidence of liver disease due to			
					other etiology such as			
					autoimmune or drug-induced			
					hepatitis			
					Serious concurrent medical			
					illnesses (such as malignancy,			
					severe cardiopulmonary disease,			
					or uncontrolled diabetes mellitus)			
					Inability to give an informed			
					written consent			

Author, Year Country Study Name Quality	Duration of Followup	Outcome	Subgroup Analyses	Subgroup Analyses	Histologic Response	Adverse Events	Funding Source
Sood, 2008 ⁶¹	Followup	A vs. B	NR	NR	NR	A vs. B	NR
India	visits at 24	ETR: 72/76(94.7%)				Overall withdrawals: 1/76 (1.3%) vs. 2/27 (7.4%);	
Continued	weeks post-	vs. 24/27(88.9%),				p=NS Withdrawala (due to AE): 0/76 vs 1/27 (40/): n=NS	
Continued	treatment	p=0.375				Withdrawals (due to AE): 0/76 vs. 1/27 (4%); p=NS Deaths: NR	
		SVR: 60/76(78.9%)				Severe Adverse Events: NR	
		vs. 25/27(926%),					
		p=0.145				Adverse events:	
						Influenza-like symptoms - 20/27(74.0 %%) vs.	
						44/76(57.9%); p=NS	
						Malaise or fatigue -10/27(37.0%) vs. 22/76(29.0%); p=NS	
						Nausea or vomiting - 5/27(18.5%) vs. 11/76(14.5%) p=NS	
						Headache 4/27 (14.8%) vs. 8/76(10.5%); p=NS	
						Abdominal discomfort - 4/27(14.8%) vs. 8/76	
						(10.5%); p=NS	
						Diarrhea 4/27(14.8%) vs. 9 /76(11.8%); p=NS	
						Grade III or IV laboratory abnormalities	
						Neutrophils - 3/27(11.1%) vs. 1/76(1.3%); p=0.02	
						Platelets - 4/27(14.8%) vs. 2/76(2.6%); p=0.02	

Evidence Table 8. Quality rating: Trials of dual therapy with pegylated interferon plus ribavirin: dose effects

Author, Year	Randomization adequate?	Allocation concealment adequate?	Groups similar at baseline?	Eligibility criteria specified?	Outcome assessors masked?	Care provider masked?	Patient masked?	Attrition and withdrawals reported?	Loss to followup: differential/high?	Intention- to-treat analysis	Quality	Funding
Abergel, 2006 ⁴⁷	Unclear	Unclear	Unclear	Yes	Unclear	Unclear	Unclear	No	Unclear	Yes	Fair	NR
Brady, 2010 ⁴⁸	Yes	Unclear	Yes	Yes	No, open label	No, open label	No, open label	Yes	Yes	Yes	Fair	Schering Plough
Fried, 2008 ⁵⁰	Unclear	Unclear	Yes	Yes	Unclear	Yes	Yes	Yes	Yes	Yes	Fair	NR
Hadziyannis, 2004 ²⁹	Yes	Yes	No	Yes	Yes	Unclear	Unclear	Yes	Unclear	Yes	Fair	Roche, Basel, Switzerland
Helbling, 2006 ⁵¹	Yes	Yes	Yes	Yes	No, open label	No, open label	No, open label	Yes	No	Yes	Fair	NR
Jacobson, 2007a ⁵²	Yes	Yes	Yes	Yes	No, open label	No, open label	No, open label	Yes	Yes	Yes	Fair	Schering- Plough Corp. , Kenilworth, NJ
Jacobson, 2007b ⁵³	Yes	Yes	Yes	Yes	No, open label	No, open label	No, open label	Yes	Yes	Yes	Fair	Schering- Plough Corp. , Kenilworth, NJ
Kawaoka, 2009 ⁵⁴	Unclear	Unclear	Unclear	Yes	No, open label	No, open label	No, open label	Yes	No	Yes	Fair	NR
Krawitt, 2006 ⁵⁵	Unclear	Unclear	Yes	Yes	Unclear	Unclear	Unclear	Unclear	Unclear	Yes	Fair	Integrated Therapeutics Group (Schering- Plough)
McHutichson, 2009 ⁶²	Unclear	Unclear	Yes	Yes	Unclear	Unclear	Unclear	Yes	Unclear	Yes	Fair	Schering- Plough
Meyer-Wyss, 2006 ⁵⁷	Unclear	Yes	No	Yes	No, open label	No, open label	No, open label	Yes	Yes	Yes	Poor	Essex Chemie AG, Lucerne
Mimidis, 2006 ⁵⁸	Unclear	Unclear	Unclear	Yes	No (not described)	No (not described)	No (not described)	No	No	No	Poor	NR
Reddy, 2010 ⁵⁹	Yes	Yes	Yes	Yes	Unclear	Yes	Yes	Yes	Yes	Yes	Fair	Roche

Author, Year	Randomization adequate?	Allocation concealment adequate?	similar at	Eligibility criteria specified?	Outcome assessors masked?	Care provider masked?	Patient masked?		Loss to followup: differential/high?		Quality	Funding
Roberts, 2009 ⁶⁰	Unclear	Unclear	Yes	Yes	/ I	No, open label	No, open label	Yes	Yes	Yes	Fair	NR
Sood, 2008 ⁶¹	Unclear	Unclear	Unclear	Yes	Unclear	Unclear	Unclear	Yes	No	Yes	Fair	NR

Key Question 4

Evidence Table 9. Studies on sustained virologic response and clinical outcomes

Author, Year	Study Type Duration of	Comparison Definition of Sustained			Number analyzed Number meeting inclusion criteria excluded due to	
Country Quality	Followup	Virological Response	Inclusion Criteria	Exclusion Criteria	missing data or lost to followup	Population Characteristics
Arase, 2007 ⁶³ Japan Overall Quality: Fair	Retrospective cohort study Duration of followup: Mean 7.4 years	SVR vs. no SVR SVR=Undetectable HCV RNA 6 months after completion of long-term IFN therapy	>=60 years of age; ALT elevation greater than double upper limits within 6 months (ALT normal range 12-50IU/l); no corticosteroid immunosuppressive agents or antiviral agents used in last 6 months; no hepatitis B surface antigens, antinuclear antibodies, or antimitochondrial antibodies detectable in serum; leukocytes>3000/mm³, platelet count >80,000/mm³, and bilirubin <2.0 mg/ml; IFN therapy >4 weeks	History of alcohol abuse or advanced liver cirrhosis, encephalopathy, bleeding esophageal varices, or ascites	Number analyzed: 500 Excluded due to missing data or lost to followup: Unclear	SVR (n=140) vs. no SVR (n=360) Mean age (years): 63 vs. 64 (p=0.07) Female: 41% vs. 53% (p=0.01) Race: Not reported Genotype 1b: 34% vs. 71% (p<0.0001) Viral load (kIU/ml): 172 vs. 661 (p<0.0001) Cirrhosis (Knodell F4): 9% vs. 16% (p=0.009)

Author, Year Country		Variables Assessed as	Variables Included in		
Quality	Treatments	Univariate Predictors	Multivariate Models	Results	Funding Source
Arase, 2007 ⁶³	Interferon alpha-2a or -	Age, sex, liver fibrosis, liver	Hepatocellular cancer: Sex,	SVR vs. no SVR	Okinaka Memorial
Japan	Interferon alpha-2b	activity, viral load, genotype,	liver fibrosis	Hepatocellular cancer:	Institute for Medical
	monotherapy: 94%	AST, ALT	All-cause and liver-related	Adjusted HR 0.19 (0.08-0.45)	Research and Japanese
Continued	Interferon plus ribavirin		mortality: Sex, liver fibrosis	All-cause mortality: Adjusted	Ministry of Health, Labor
	combination therapy: 6%			HR 0.39 (0.16-0.93)	and Welfare
				Liver-related mortality:	
				Adjusted HR 0.13 (0.03-0.59)	

Author, Year Country	Study Type Duration of	Comparison Definition of Sustained			Number analyzed Number meeting inclusion criteria excluded due to missing data or lost to	
Quality Backus, 2011 ⁶⁴	Followup Retrospective	Virological Response SVR vs. no SVR	Inclusion Criteria HCV genotype 1, 2, or 3;	Exclusion Criteria HIV infection,	followup Number analyzed: 16,864	Population Characteristics SVR vs. no SVR
USA Overall Quality: Fair	cohort study Duration of followup: Median 3.8 years	SVR=Undetectable HCV RNA 6 months after completion of antiviral therapy	started pegylated interferon + ribavirin between January 2001 and June 2007; stopped treatment by June 2008; HCV RNA test after end of treatment	hepatocellular cancer prior to treatment	Excluded due to missing data or lost to followup: 5365	(genotypes 1 [n=12,166], 2 [n=2904], and 3 [n=1794]) Mean age (years): 51 vs. 52, 53 vs. 53. and 51 vs. 51 Female: 5% vs. 4%, 4% vs. 3%, and 4% vs. 3% Non white: 40% vs. 51%, 33% vs. 31%, and 30% vs. 29% Genotype: Results stratified by genotype Viral load >=500,000 IU/mL: 70% vs. 82%, 78% vs. 83%, and 64% vs. 68%
						vs. 83%, and 64% vs. 68% Cirrhosis: 9% vs. 15%, 7% vs. 12%, and 12% vs. 20%

Author, Year					
Country		Variables Assessed as	Variables Included in		
Quality	Treatments	Univariate Predictors	Multivariate Models	Results	Funding Source
Backus, 2011 ⁶⁴	Pegylated interferon (alfa-2aa	Age, sex, albumin, AST,	Age, sex, albumin, AST,	SVR vs. no SVR (genotypes	US Department of
USA	or 2b) plus ribavirin	AST/ALT ratio, creatinine	AST/ALT ratio, creatinine	1, 2, and 3, respectively)	Veterans Affairs, Veterans
		clearance, platelets, sodium,	clearance, platelets, sodium,	All-cause mortality: Adjusted	Health Administration,
Continued		cirrhosis, Chronic obstructive	cirrhosis, Chronic obstructive	HR 0.71 (0.60-0.86), 0.62	Office of Public Health
		pulmonary disease (COPD),	pulmonary disease (COPD),	(0.44-0.87), and 0.51 (0.35-	and Environmental
		diabetes, HTN, tobacco use,	diabetes, HTN, tobacco use,	0.75)	Hazards
		treatment duration <60%	treatment duration <60%		
		recommended, bilirubin, body	recommended, bilirubin, body		
		mass index, HBV co-infection,	mass index, HBV co-infection,		
		viral load, hemoglobin, CAD,	viral load, hemoglobin,		
		cancer, congestive heart	coronary artery disease,		
		failure, cerebrovascular	cancer, congestive heart		
		disease, schizophrenia, recent	failure, cerebrovascular		
		alcohol abuse, anxiety	disease, schizophrenia, recent		
		disorder, depression, hard drug	alcohol abuse, anxiety		
		use, post-traumatic stress	disorder, depression, hard drug		
		disorder (PTSD),	use, post-traumatic stress		
		socioeconomic status	disorder (PTSD),		
		instability, multiple treatment	socioeconomic status		
		course, erythropoiesis	instability, multiple treatment		
		stimulating agent use,	course, erythropoiesis		
		granulocyte colony stimulating	stimulating agent use,		
		factor use, year of treatment	granulocyte colony stimulating		
		start	factor use, year of treatment		
			start		

Author, Year	Study Type	Comparison			Number analyzed Number meeting inclusion criteria excluded due to	
Country	Duration of	Definition of Sustained		F 1 . G	missing data or lost to	B 14: Cl 4:4:
Quality	Followup	Virological Response	Inclusion Criteria	Exclusion Criteria	followup	Population Characteristics
Bruno, 2007 ⁶⁵	Retrospective	SVR vs. no SVR	Anti-HCV and HCV-RNA	Over 70 years of age; lack	Number analyzed: 883	SVR (n=124) vs. no SVR
Italy	cohort study	SVR=Undetectable HCV	positive and diagnosis of	of histological diagnosis of	Excluded due to missing	(n=759)
Overall Quality: Fair	Duration of	RNA 6 months after	complete cirrhosis by	cirrhosis, gastroesophageal	data or lost to followup:	Mean age (years): 53 vs. 44
	followup: Mean 8	completion of antiviral	histological criteria (Ishak	varices; previous episodes	Unclear	(p=0.004)
	years	therapy	score of 6 or Knodell score of	of decompensation or		Female: 27% vs. 38%
			4); liver biopsy within 18	bleeding; Child class B or		(p<0.001)
			months of start of IFN	C, concurrent		Non White: 0 (0%) vs. 0
			treatment	Hepatocellular carcinoma		(0%)
				or extra hepatic tumors;		Race: Not reported
				subjects co-infected with		Genotypes 1 and 4: 37% vs.
				hepatitis B or HIV		63% (p<0.001)
						Viral load: Not reported
						Cirrhosis: All (inclusion
						criterion)

Author, Year Country Quality	Treatments	Variables Assessed as Univariate Predictors	Variables Included in Multivariate Models	Results	Funding Source
Bruno, 2007 ⁶⁵	Interferon monotherapy	Age, sex, platelet count,	Hepatocellular carcinoma:	SVR vs. no SVR	Associazione per la Ricera
Italy Continued		genotype	Age, sex, platelet count Liver-related mortality: Age, platelet count	Ascites, encephalopathy, or gastrointestinal bleeding: Not calculated, 0 events/1061 person-years vs. 107 events/5703 person-years (1.88 events/100 person-years) Hepatocellular carcinoma: Adjusted HR 0.39 (0.17-0.88) Liver-related mortality: 0.14 (0.04-0.59)	sulle Malattie Epatiche (ARME), Bologna, Italy

Author, Year Country Quality	Study Type Duration of Followup	Comparison Definition of Sustained Virological Response	Inclusion Criteria	Exclusion Criteria	Number analyzed Number meeting inclusion criteria excluded due to missing data or lost to followup	Population Characteristics
Cardoso, 2010 ⁶⁶ France Overall Quality: Fair	Retrospective cohort study (of patients originally enrolled in clinical trials) Duration of followup: Median 3.5 years	SVR vs. no SVR SVR=Undetectable HCV RNA 6 months after completion of antiviral therapy	Anti-HCV and HCV RNA positive, documented chronic hepatitis C, biopsy-proven bridging fibrosis or cirrhosis, treated with interferon-based therapy in clinical trials between 1987 and 2007	HBV, hepatitis D virus, or HIV infection co-infection; history of hepatic decompensation	Number analyzed: 307 Excluded due to missing data or lost to followup: Unclear	SVR (n=103) vs. no-SVR (n=204) Mean age (years): 55 vs. 55 (p=0.93) Female: 30% vs. 34% (p=0.51) Race: Not reported Genotype 1: 36% vs. 72% (p<0.001) Viral load (log ₁₀ l/ml): 5.5 vs. 5.7 (p=0.08) Cirrhosis (METAVIR F4): 53% vs. 61% (p=0.19)

Author, Year Country	T	Variables Assessed as	Variables Included in	D. I	T. P. G
Quality	Treatments	Univariate Predictors	Multivariate Models	Results	Funding Source
Cardoso, 2010 ⁶⁶	Pegylated interferon and	Age, sex, BMI, alcohol	Hepatocellular carcinoma:	SVR vs. no SVR	Schering Plough
France	ribavirin: 252 (82%)	consumption, diabetes, ALT,	Age, bilirubin, albumin,	Hepatocellular carcinoma:	
	Pegylated interferon	bilirubin, albumin, platelets,	platelet count	Adjusted HR 0.33 (0.23-0.89)	
Continued	monotherapy: 22 (7%)	genotype, viral load,	Ascites/variceal bleeding and	Ascites or variceal bleeding:	
	Conventional interferon with or	inflammation, fibrosis and	liver-related mortality:	Adjusted HR 0.21 (0.05-0.92)	
	without ribavirin: 33 (11%)	steatosis scores	Bilirubin, albumin, platelets	Liver-related mortality:	
				Adjusted HR 0.27 (0.08-0.95)	

Author, Year Country Quality	Study Type Duration of Followup	Comparison Definition of Sustained Virological Response	Inclusion Criteria	Exclusion Criteria	Number analyzed Number meeting inclusion criteria excluded due to missing data or lost to followup	Population Characteristics
Coverdale, 2004 ⁶⁷	Prospective cohort	SVR vs. response relapse	Virologically and	Clinical or imaging	Number analyzed: 343	Demographics for all
Australia	study (some	vs. nonresponse	histologically proven chronic	evidence of liver-related	Excluded due to missing	treated patients (not
Overall Quality:	patients originally	SVR=Undetectable HCV	hepatitis C	complications	data or lost to followup:	reported by SVR status)
Poor	enrolled in	RNA on at least 2			Unclear	Median age (years): 37
	randomized trials)	occasions at least 2 years				Female: 33%
	Duration of	after completion of				Race: Not reported
	followup: Median	therapy				Genotype 1: 38%
	9 years					Viral load: Not reported
						Median fibrosis score
						(Scheuer): 2

Author, Year Country	Treatments	Variables Assessed as Univariate Predictors	Variables Included in Multivariate Models	Results	Funding Source
Quality					
Coverdale, 2004 ⁶⁷	Interferon alpha-2a or	Statistically significant	Age, duration, place of birth,	SVR vs. response-relapse vs.	National Institutes of
Australia	Interferon alpha-2b	predictors of outcomes in	mode of transmission,	nonresponse	Health
		univariate analyses were age,	genotype, fibrosis score,	Liver-related complications	
Continued		duration, place of birth, mode	albumin, bilirubin,	(hepatic decompensation,	
		of transmission, genotype,	prothrombin time	complications of portal	
		fibrosis score, albumin,		hypertension, hepatocellular	
		bilirubin, prothrombin time.		carcinoma, liver	
		Other tested variables not		transplantation, and liver-	
		reported.		related mortality) at 10 years:	
				Not statistically significant in	
				multivariate analysis, adjusted	
				HR not reported (p=0.06)	
				Hepatocellular carcinoma at	
				10 years: Not statistically	
				significant in multivariate	
				analysis, adjusted HR and p	
				value not reported	
				Liver transplant or liver-	
				related death at 10 years: Not	
				statistically significant in	
				multivariate analysis, adjusted	
				HR not reported (p=0.20)	

Author, Year Country Quality	Study Type Duration of Followup	Comparison Definition of Sustained Virological Response	Inclusion Criteria	Exclusion Criteria	Number analyzed Number meeting inclusion criteria excluded due to missing data or lost to followup	Population Characteristics
El Braks, 2007 ⁶⁸ France Overall Quality: Poor	Retrospective cohort study Duration of followup: Mean 7.7 years	SVR vs. no SVR SVR=Undetectable HCV RNA 6 months after completion of antiviral therapy	HCV-related cirrhosis defined by association of positive serum HCV antibodies and RNA, with typical liver histology; absence of complication before or at inclusion; daily alcohol consumption <50 g; at least 3 month course of antiviral treatment using standard or pegylated interferon with or without ribavirin, according to therapeutic advance over time and initial guidelines; a regular followup >=30 months after the starting of first treatment; residence in France allowing regular followup	HBV or HIV co-infection; contraindication to antiviral treatment, particularly platelet and polymorphonuclear counts ≤80,000/mm3 and 1500/mm3, respectively; Hepatocellular carcinoma or suspicious findings such as liver nodule or serum level of alpha-fetoprotein above 50 ng/mL	Number analyzed: 113 Excluded due to missing data or lost to followup: Unclear	SVR (n=37) vs. no SVR (n=76) Mean age (years): 51 vs. 56 (p=0.02) Female: 16% vs. 50% (p=0.0005) Race: Not reported HCV genotype 1: 36% vs. 73% (p=0.0001) Viral load: Not reported Cirrhosis: All (inclusion criterion)

Author, Year Country Quality	Treatments	Variables Assessed as Univariate Predictors	Variables Included in Multivariate Models	Results	Funding Source
El Braks, 2007 ⁶⁸	Interferon monotherapy: 35/113	Age, sex, genotype, duration	Duration of treatment	SVR (n=37) vs. no SVR	Not reported
France	(31%)	of treatment		(n=76)	
	Interferon + ribavirin: 40/113			Clinical events	
Continued	(35%)			(hepatocellular cancer,	
	Pegylated interferon + ribavirin:			ascites, hepatic	
	38/113 (34%)			encephalopathy, or death):	
				Adjusted HR 0.14 (0.04-0.45)	

Author, Year Country	Study Type Duration of Followup	Comparison Definition of Sustained	Inclusion Criteria	Exclusion Criteria	Number analyzed Number meeting inclusion criteria excluded due to missing data or lost to followup	Paralation Characteristics
Quality Fernandez- Rodriguez, 2010 ⁶⁹ Spain Overall Quality: Poor	Retrospective cohort study Duration of followup: Median 35 months	SVR vs. no SVR SVR=Undetectable HCV RNA 6 months after completion of antiviral therapy	HCV-associated cirrhosis	Child-Pugh-Turcotte's score (CPT) >6; HIV or HBV co infection; alcohol intake >40 g per day in males or >20 g per day in females; present or past psychosis or severe depression; neutropenia <1500 per ml and/or thrombocytopenia <100,000 platelets per ml; organ transplantation; severe heart disease; uncontrolled seizures; uncontrolled diabetes; autoimmune disorders; end-stage renal failure; anemia; hemoglobinopathies; severe heart disease; pregnancy; no reliable method of contraception; uncontrolled arterial hypertension; age older than 70 years	Number analyzed: 509 Excluded due to missing data or lost to followup: 59	Population Characteristics SVR (n=174) vs. no SVR (n=394) Mean age (years): 51 vs. 52 (p=0.31) Female: 69% vs. 73%, p=0.37 Genotype 1: 24% vs. 55% (p=0.001) Race: Not reported Viral load (10 ⁶ IU/ml): 1.7 vs. 3.1 (p=0.001) Cirrhosis: All (inclusion criterion)

Author, Year Country		Variables Assessed as	Variables Included in		
Quality	Treatments	Univariate Predictors	Multivariate Models	Results	Funding Source
Fernandez-Rodriguez, 2010 ⁶⁹	Pegylated interferon-2a or 2b	Statistically significant	Age, albumin, esophageal	SVR vs. no SVR	Study conducted on behalf
Spain		predictors of outcomes in	varices, ultrasonographic signs	Combined clinical endpoint	of the Group for the
		univariate analyses were age,	of portal hypertension, platelet	(hepatic decompensation,	Assessment of Prevention
Continued		albumin, esophageal varices,	count, bilirubin, prothrombin	upper gastrointestinal	of Cirrhosis Complications
		ultrasonographic signs of	activity	bleeding secondary to rupture	and Virological Response
		portal hypertension, platelet		of esophageal or gastric	(APREVIR). No
		count, bilirubin, prothrombin		varices, hepatocellular	additional funding
		activity. Other tested variables		carcinoma, liver	sources.
		not reported.		transplantation, and liver-	
				related or liver-unrelated	
				mortality): Adjusted HR 0.38	
				(0.18-0.76)	

Author, Year Country Quality	Study Type Duration of Followup	Comparison Definition of Sustained Virological Response	Inclusion Criteria	Exclusion Criteria	Number analyzed Number meeting inclusion criteria excluded due to missing data or lost to followup	Population Characteristics
Hasegawa, 2007 ⁷⁰ Japan Overall Quality: Fair	Retrospective cohort study Duration of followup: Median 4.6 years	SVR vs. no SVR SVR=Sustained undetectable HCV RNA after completion of antiviral therapy (duration of undetectability not specified)	HCV-associated cirrhosis	HBV co-infection	Number analyzed: 105 Excluded due to missing data or lost to followup: Unclear	SVR (n=48) vs. no SVR (n=58) Age >56 years: 60% vs. 55% (p>0.05) Male: 65% vs. 66% (p>0.05) Race: Not reported Genotype 1b: 19% vs. 21% (p>0.05) Viral load >=100 KIU/ml or >=1 Meq/mL: 25% vs. 62% (p<0.001) Cirrhosis: All (inclusion criterion)

Author, Year Country		Variables Assessed as	Variables Included in		
Quality	Treatments	Univariate Predictors	Multivariate Models	Results	Funding Source
Hasegawa, 2007 ⁷⁰	Natural or recombinant	Age, sex, BMI, albumin,	Choline esterase, alpha-	SVR vs. no SVR	Not reported
Japan	Interferon alpha: 67%	cholinesterase, platelet count,	fetoprotein, viral load, daily	Hepatocellular carcinoma:	
	Natural Interferon-beta: 31%	alpha-fetoprotein, indocyanine	dose of interferon, duration of	Adjusted HR 0.18 (0.04-0.81)	
Continued	Both: 1.6%	green retention rate at 15	interferon, use of induction		
		minutes, fasting blood glucose,	therapy		
		AST, ALT, viral load,			
		genotype, use of combination			
		therapy, total dose of			
		interferon, daily dose of			
		interferon, use of induction			
		therapy, type of interferon			

Country	Study Type Duration of Followup	Comparison Definition of Sustained Virological Response	Inclusion Criteria	Exclusion Criteria	Number analyzed Number meeting inclusion criteria excluded due to missing data or lost to followup	Population Characteristics
Hung, 2006 ⁷¹ Taiwan Overall Quality: Fair	Cohort study (unclear if retrospective or prospective) Duration of followup: Median 37 months	SVR vs. no SVR SVR=Undetectable HCV RNA 6 months after completion of antiviral therapy	Anti-HCV positive; elevated ALT values for at least 6 months; Child-Pugh score A	HIV or HBV co-infection; alcoholism; autoimmune hepatitis; major contraindications to IFN or ribavirin therapy; severe thrombocytopenia or a history of hepatic encephalopathy, bleeding esophageal varices and ascites	Number analyzed: 132 Excluded due to missing data or lost to followup: Unclear	SVR (n=73) vs. no SVR (n=59) Mean age (years): 55 vs. 58 (p=0.07) Female: 43% vs. 54% (p=0.12) Race: Not reported Genotype 1b: 27% vs. 78% (p<0.001) Viral load >=2 x 10 ⁶ copies/ml: 21% vs. 51% (p<0.001) Cirrhosis: 100% (inclusion criterion)

Author, Year					
Country		Variables Assessed as	Variables Included in		
Quality	Treatments	Univariate Predictors	Multivariate Models	Results	Funding Source
Hung, 2006 ⁷¹	Interferon-2b plus ribavirin	Age, sex, body weight, viral	Age, sex, body weight, viral	SVR vs. no SVR	Chang Gung Memorial
Taiwan		load, platelet count, ALT,	load, platelet count, ALT,	Hepatocellular carcinoma:	Hospital and Department
		Histological Activity Index	Histological Activity Index	Adjusted HR 0.28 (0.09-0.92)	of Health of Taiwan
Continued		score, genotype	score, genotype		

Author, Year Country Quality	Study Type Duration of Followup	Comparison Definition of Sustained Virological Response	Inclusion Criteria	Exclusion Criteria	Number analyzed Number meeting inclusion criteria excluded due to missing data or lost to followup	Population Characteristics
Imazeki, 2003 ⁷² Japan Overall Quality: Fair	Retrospective cohort study Duration of followup: Mean 8.2 years	SVR vs. no SVR SVR=Undetectable HCV RNA 6 months after completion of antiviral therapy	HCV RNA positive who underwent liver biopsy	Hepatocellular carcinoma detected within six months of liver biopsy	Number analyzed: 459 Excluded due to missing data or lost to followup: 9	Demographics for all treated patients (not reported by SVR status) Mean age (years): 49 Female: 36% Race: Not reported Genotype 1: 74% Viral load: Not reported Cirrhosis (Desmet F4): 13%

Author, Year Country Quality	Treatments	Variables Assessed as Univariate Predictors	Variables Included in Multivariate Models	Results	Funding Source
Imazeki, 2003 ⁷²	Interferon-2a: 84%	Age, sex, fibrosis stage, AST,	Age, sex, fibrosis stage, AST,	SVR vs. untreated and no	Not reported
Japan	Interferon-2b: 12% Both: 4%	ALT, albumin, platelet count, viral load, genotype, alcohol	ALT, albumin, platelet count, alcohol consumption, duration	SVR vs. untreated Liver-related mortality:	
Continued	Both. 170	consumption, duration of disease, BMI, co morbidities, diabetes mellitus, hypertension, fatty liver, cardiopulmonary disease	of disease	Adjusted HR 0.06 (0.007- 0.43) and 0.55 (0.27-1.1) All-cause mortality: Adjusted HR 0.030 (0.003-0.27) and 0.26 (0.11-0.61)	
				SVR vs. no SVR# Liver-related mortality: Adjusted HR 0.11 (0.01-0.96) All-cause mortality: Adjusted HR 0.12 (0.01-1.3)	

Author, Year Country Quality	Study Type Duration of Followup	Comparison Definition of Sustained Virological Response	Inclusion Criteria	Exclusion Criteria	Number analyzed Number meeting inclusion criteria excluded due to missing data or lost to followup	Population Characteristics
Innes, 2011 ⁷³ UK Overall Quality: Fair	Retrospective cohort study Duration of followup: Mean 5.3 years	SVR vs. no SVR SVR=Undetectable HCV RNA >6 months after completion of antiviral therapy	Initial course of antiviral therapy	Unsustained SVR (presence of viremia subsequent to meeting definition for SVR), HIV- positive, unknown treatment response	Number analyzed: 1215 Number excluded: 48	SVR (560) vs. no SVR (655) Mean age (years): 42 overall Female: 34% vs. 28% Non white: 10% vs. 6% Genotype 1: 19% vs. 50% Viral load: Not reported Cirrhosis: 10% vs. 18%

Author, Year Country		Variables Assessed as	Variables Included in		
Quality	Treatments	Univariate Predictors	Multivariate Models	Results	Funding Source
Innes, 2011 ⁷³	Pegylated interferon plus	Sex, age, race, injection drug	Age, race (liver-related	SVR vs. no SVR	Scottish government
UK	ribavirin: 61%	use, genotype, cirrhosis,	hospitalizations only),	Liver-related mortality:	
	Pegylated interferon	alcohol-related hospitalization,	injection drug use (liver-	Adjusted HR 0.22 (0.09-0.58)	
Continued	monotherapy: 1%	elevated ALT	related hospitalizations only),	Liver-related hospital	
	Interferon plus ribavirin: 21%		cirrhosis, alcohol-related	episode: Adjusted HR 0.22	
	Interferon monotherapy: 18%		hospitalization, elevated ALT	(0.15-0.34)	

Author, Year Country	Study Type Duration of	Comparison Definition of Sustained			Number analyzed Number meeting inclusion criteria excluded due to missing data or lost to	
Quality Izumi, 2005 ⁷⁴	Followup	Virological Response SVR vs. no SVR	Inclusion Criteria	Exclusion Criteria	followup	Population Characteristics
Japan Overall Quality: Fair	Cohort study, appears retrospective Duration of followup: Not reported	SVR=Undetectable HCV RNA 6 months after completion of antiviral therapy	Chronic HCV infection, underwent antiviral therapy	Not reported	Number analyzed: 495 Excluded due to missing data or lost to followup: Unclear	Demographics for patients treated with interferon monotherapy and interferon plus ribavirin combination therapy, respectively (not reported by SVR status) Mean age (years): 52 and 58 Female: 43% and 44% Race: Not reported Genotype 1b: 71% and 80% Median viral load (kIU/ml): 470 and 680 Cirrhosis: 35% and 2%

Author, Year Country		Variables Assessed as	Variables Included in		
Quality	Treatments	Univariate Predictors	Multivariate Models	Results	Funding Source
Izumi, 2005 ⁷⁴	Interferon monotherapy:69%	Not reported	Unclear; age, sex, and fibrosis	SVR vs. no SVR	Japanese Ministry of
Japan	Interferon-2b plus ribavirin		stage reported as statistically	Hepatocellular carcinoma:	Health Labor and Welfare
	combination therapy: 34%		significant predictors of	Adjusted HR 0.36 (0.04-0.83)	
Continued			outcomes in multivariate		
			model		

Author, Year Country	Study Type Duration of	Comparison Definition of Sustained			Number analyzed Number meeting inclusion criteria excluded due to missing data or lost to	
Quality	Followup	Virological Response	Inclusion Criteria	Exclusion Criteria	followup	Population Characteristics
Kasahara, 2004 ⁷⁵	Retrospective	SVR vs. no SVR	Histological diagnosis of	History of clinical signs at	Number analyzed: 2698	SVR (n=738) vs. no-SVR
Japan	cohort	SVR=Undetectable HCV	chronic hepatitis or cirrhosis	entry into the study of	Excluded due to missing	(n=1930)
Overall Quality:	Duration of	RNA 6 months after		complications of cirrhosis,	data or lost to followup:	Median age (years): 51 vs.
Poor	followup: Mean 6	completion of antiviral		i.e. ascites, jaundice,	Unclear	54 (p=0.12)
	years	therapy		encephalopathy, or variceal		Female: 31% vs. 37%
				bleeding; evidence of		(p=0.32)
				Hepatocellular carcinoma		Race: Not reported
				at entry as assessed by		Genotype 1: Not reported
				ultrasonography and/or		Viral load: Not reported
				computed tomography;		Cirrhosis (Desmet F4):
				HBV co-infection; co-		3.0% vs. 5.4% (p=0.34)
				existing liver diseases such		_
				as autoimmune hepatitis or		
				primary biliary cirrhosis;		
				excessive alcohol		
				consumption (>80 g/day);		
				HIV co-infection		

Author, Year Country Quality	Treatments	Variables Assessed as Univariate Predictors	Variables Included in Multivariate Models	Results	Funding Source
Kasahara, 2004 ⁷⁵	Interferon	Univariate analyses not	Age, sex, fibrosis score, time	SVR vs. no SVR	Not reported
Japan		performed	at liver biopsy	Liver-related mortality:	
				Adjusted HR 0.04 (0.005-	
Continued				0.30)	
				All-cause mortality: Adjusted	
				HR 0.14 (0.06-0.35)	

Author, Year Country Quality	Study Type Duration of Followup	Comparison Definition of Sustained Virological Response	Inclusion Criteria	Exclusion Criteria	Number analyzed Number meeting inclusion criteria excluded due to missing data or lost to followup	Population Characteristics
Maruoka, 2012 ⁷⁶	Retrospective	SVR vs. no SVR	HCV positive, underwent	Other causes of chronic	Number analyzed: 577	For all treated patients (not
Japan Overall Quality: Fair	cohort study Duration of followup: Mean 9.9 years	SVR=Undetectable HCV RNA >6 months after completion of antiviral therapy	liver biopsy	liver disease, HIV-positive, detection of hepatocellular cancer within 1 year of antiviral therapy, dropout within one year	(received antiviral therapy) Excluded due to missing data or loss to followup: Unclear for those treated with antiviral therapy, including persons untreated 114/835 lost to followup within 1 year	reported by SVR status) Mean age (years): 50 Female: 36% Non white: Not reported Genotype 1: 73% Viral load high (≥100 KIU, 100 kc, 1.0 Meq, 10 ⁴ /50 mcL, or 30 core antigens): 69% Cirrhosis: 10%

Author, Year					
Country		Variables Assessed as	Variables Included in		
Quality	Treatments	Univariate Predictors	Multivariate Models	Results	Funding Source
Maruoka, 2012 ⁷⁶	Interferon- Or-	Sex, age, fibrosis stage,	Sex (mortality only), age	SVR vs. untreated patients	Not reported
Japan	monotherapy: 83%	inflammatory grade, genotype,	(hepatocellular cancer only),	and no SVR vs. untreated	
	Interferon- or sequential	high viral load, genotype 1 and	fibrosis stage, inflammatory	patients	
Continued	therapy: 3.3%	high viral load, elevated ALT,	grade, genotype 1 and high	All-cause mortality: Adjusted	
	Interferon- ribavirin	low platelets, low albumin	viral load (hepatocellular	HR 0.17 (0.08-0.40) and 0.84	
	combination therapy: 14%		cancer only), elevated ALT,	(0.50-1.4)	
			low platelets, low albumin	Hepatocellular carcinoma:	
				Adjusted HR: 0.14 (0.05-	
				0.42) and 1.2 (0.69-2.0)	
				SVR vs. no SVR#	
				All-cause mortality: Adjusted	
				HR 0.20 (0.08-0.54)	
				Hepatocellular carcinoma:	
				Adjusted HR 0.12 (0.04-0.40)	

Author, Year Country Quality	Study Type Duration of Followup	Comparison Definition of Sustained Virological Response	Inclusion Criteria	Exclusion Criteria	Number analyzed Number meeting inclusion criteria excluded due to missing data or lost to followup	Population Characteristics
Morgan, 2010 ⁷⁷ USA Overall Quality: Fair	Prospective cohort study of patient enrolled in a randomized trial Duration of followup: Median 79 to 86 months	SVR vs. no SVR SVR=Undetectable HCV RNA 6 months after completion of antiviral therapy	Advanced hepatic fibrosis (Ishak fibrosis score 3) according to liver biopsy performed within 12 months; lack of SVR to previous treatment for at least 24 weeks with standard interferon with or without ribavirin; no history of hepatic decompensation or Hepatocellular carcinoma	Not reported	Number analyzed: 526 Excluded due to missing data or lost to followup: 30 of 180 patients with SVR, not reported for breakthrough/relapse and nonresponder groups	SVR (n=140) vs. breakthrough/relapse (n=77) vs. no SVR (n=309) Mean age (years): 49 vs. 49 vs. 50 (p=0.23) Female: 24% vs. 26% vs. 30% (p=0.30) Non white: 20% vs. 20% vs. 32% (p=0.001) Genotype 1: 72% vs. 86% vs. 94% (p<0.0001) Viral load: Not reported Cirrhosis (Ishak 5 or 6): 21% vs. 31% vs. 43% (p<0.0001)

Author, Year Country Quality	Treatments	Variables Assessed as Univariate Predictors	Variables Included in Multivariate Models	Results	Funding Source
Morgan, 2010 ⁷⁷ USA Continued	Pegylated interferon-2a-180 μg/week + ribavirin 1000-12000 mg/day for 24weeks	Not reported	Age, race, platelet count, AST/ALT ratio, albumin, alkaline phosphatase, alpha- fetoprotein	SVR vs. no SVR All-cause mortality or liver transplantation: Adjusted HR 0.17 (0.06-0.46) Any liver-related outcome (decompensated liver disease [ascites, variceal bleeding, hepatic encephalopathy, spontaneous bacterial peritonitis], hepatocellular carcinoma, liver transplantation, liver-related mortality): Adjusted HR 0.15 (0.06-0.38) Decompensated liver disease: Adjusted HR 0.13 (0.03-0.53) Hepatocellular carcinoma: Adjusted HR 0.19 (0.04-0.80) Liver-related mortality or liver transplantation: Adjusted HR 0.12 (0.03-0.48)	National Institute of Diabetes and Digestive and Kidney Diseases, National Institute of Allergy and Infectious Diseases, the National Cancer Institute, the National Institutes of Health, and Hoffmann-La Roche, Inc

Author, Year Country Quality	Study Type Duration of Followup	Comparison Definition of Sustained Virological Response	Inclusion Criteria	Exclusion Criteria	Number analyzed Number meeting inclusion criteria excluded due to missing data or lost to followup	Population Characteristics
Shiratori, 2005 ⁷⁸ Japan Overall Quality: Poor	Prospective cohort study of patients enrolled in randomized trials Duration of followup: Median 6.8 years	SVR vs. no SVR SVR=Undetectable HCV RNA 6 months after completion of antiviral therapy	HCV positive, elevated ALT levels for more than 6 months, abnormal histologic findings on liver biopsy specimens, indicating fibrotic state F4, platelet county greater than 3 x 10 ⁹ cells/L and Child-Pugh A classification	HBV infection, autoimmune hepatitis, primary biliary cirrhosis, drug-induced liver disease, hepatocellular carcinoma on imaging prior to enrollment	Number analyzed: 271 Excluded due to missing data or lost to followup: 30 at 3 years, 86 at 7 years	For all treated patients (not reported by SVR status) Mean age (years): 57 Female: 62% Race: Not reported Genotype 1: 75% Viral load (log ₁₀ copies/ml): 5.8 Cirrhosis: 100% (inclusion criterion)

Author, Year Country Quality	Treatments	Variables Assessed as Univariate Predictors	Variables Included in Multivariate Models	Results	Funding Source
Shiratori, 2005 ⁷⁸	Interferon α-2a: 58%	Univariate analyses not	Age	SVR vs. untreated patients	None declared
Japan	Natural interferon α: 42%	performed		and no SVR vs. untreated	
				patients	
Continued				Hepatocellular carcinoma:	
				Adjusted HR 0.31 (0.16-0.61)	
				and 0.77 (0.51-1.2)	
				All-cause mortality: Adjusted	
				HR 0.05 (0.006-0.34) and	
				0.71 (0.43-1.2)	
				SVR vs. no SVR#	
				Hepatocellular carcinoma:	
				Adjusted HR 0.40 (0.18-0.89)	
				All-cause mortality: Adjusted	
				HR 0.07 (0.01-0.56)	

Author, Year Country Quality	Study Type Duration of Followup	Comparison Definition of Sustained Virological Response	Inclusion Criteria	Exclusion Criteria	Number analyzed Number meeting inclusion criteria excluded due to missing data or lost to followup	Population Characteristics
Veldt, 2007 ⁷⁹ Europe and Canada Overall Quality: Poor	Retrospective cohort Duration of followup: Median 2.1 years	SVR vs. no SVR SVR=Undetectable HCV RNA 6 months after completion of antiviral therapy	Biopsy-proven advanced fibrosis or cirrhosis (Ishak score, 4 to 6) treated with interferon-based regimen	HIV or HBV co-infection; decompensated liver disease	Number analyzed: 479 Excluded due to missing data or lost to followup: Unclear	SVR (n=142) vs. no-SVR (n=337) Mean age (years): 48 vs. 49 (p=0.45) Female: 27% vs. 32% (p=0.23) Race: Not reported Genotype 1: 39% vs. 67% (p<0.001) Viral load (x10 ⁵ IU/mL): 8.5 vs. 8.0 (p=0.75) Cirrhosis (Ishak 5 or 6): 71% vs. 77% (p=0.45)

Author, Year Country		Variables Assessed as	Variables Included in		
Quality	Treatments	Univariate Predictors	Multivariate Models	Results	Funding Source
Veldt, 2007 ⁷⁹	Interferon monotherapy: 27%	Univariate analyses not	All outcomes: Age, sex,	SVR vs. no SVR	Netherlands Organisation
Europe and Canada	Interferon and ribavirin: 27%	performed	previous non response,	Any event (death, liver	for Health Research and
	Pegylated interferon		bilirubin level, albumin level,	failure, and hepatocellular	Development
Continued	monotherapy: 2.1%		platelet count, treatment	cancer): Adjusted HR 0.20	
	Pegylated interferon and		center, treatment period	(0.07-0.58)	
	ribavirin: 43%		Hepatocellular carcinoma:	All-cause mortality: Adjusted	
			Also adjusted for anti-hepatitis	HR 0.31 (0.07-1.4)	
			B core antigen positivity	Liver-related mortality:	
				Adjusted HR 0.19 (0.02-1.4)	
				Hepatocellular carcinoma:	
				Adjusted HR 0.46 (0.12-1.70)	

Author, Year Country Quality	Study Type Duration of Followup	Comparison Definition of Sustained Virological Response	Inclusion Criteria	Exclusion Criteria	Number analyzed Number meeting inclusion criteria excluded due to missing data or lost to followup	Population Characteristics
Yoshida, 2002 ⁸⁰ Japan Overall Quality: Poor	Retrospective cohort Duration of followup: Mean 5.4 years	SVR vs. no SVR SVR=Undetectable HCV RNA 6 months after completion of antiviral therapy	HCV antibody positive; received liver biopsy	HBV co-infection, alcoholic liver disease, autoimmune hepatitis, or primary biliary cirrhosis.	Number analyzed: 2889 Excluded due to missing data or lost to followup: Unclear	SVR (817) vs. non SVR (1613) Mean age (years): 48 vs. 51 Female: 30% vs. 40% Race: Not reported Genotype: Not reported Viral load: Not reported Cirrhosis (Desmet F4): 6.5% vs. 11%

Author, Year Country Quality	Treatments	Variables Assessed as Univariate Predictors	Variables Included in Multivariate Models	Results	Funding Source
Yoshida, 2002 ⁸⁰	Interferon-alpha: 84%	Univariate analyses not	Age, sex	SVR vs. untreated and no	Ministry of Health,
Japan	Interferon-beta: 14%	performed		SVR vs. untreated	Labour, and Welfare of
	Both: 2%			Liver-related mortality:	Japan and Ministry of
Continued				Adjusted HR 0.050 (0.01-	Education, Culture,
				0.22) and 0.39 (0.22-0.68)	Sports, Science, and
				All-cause mortality: Adjusted	Technology of Japan
				HR 0.15 (0.06-0.34) and 0.47	
				(0.29-0.76)	
				SVR vs. no SVR#	
				Liver-related mortality:	
				Adjusted HR 0.13 (0.02-0.66)	
				All-cause mortality Adjusted	
				HR 0.32 (0.12-0.86)	

Author, Year Country	Study Type Duration of	Comparison Definition of Sustained			Number analyzed Number meeting inclusion criteria excluded due to missing data or lost to	
Quality	Followup	Virological Response	Inclusion Criteria	Exclusion Criteria	followup	Population Characteristics
Yu, 2006 ⁸¹	Retrospective	SVR vs. no SVR	Seropositive for anti-HCV	Concurrent HBV infection,	Number analyzed: 1057	For all treated patients (not
Taiwan	cohort	SVR=Undetectable HCV	antibody and HCV RNA and	HIV infection, autoimmune	Excluded due to missing	reported by SVR status)
Overall Quality:	Duration of	RNA 6 months after	biopsy-proven chronic	hepatitis, heavy ETOH use	data or lost to followup:	Mean age (years): 47
Poor	followup: Mean	completion of antiviral	hepatitis with or without	(>80g/day), or evidence of	Unclear	Female: 40%
	5.2 years	therapy	cirrhosis	Hepatocellular carcinoma		Race: Not reported
				-		Genotype 1: 46%
						Viral load: Not reported
						Cirrhosis (criteria not
						reported): 16%

Author, Year Country Quality	Treatments	Variables Assessed as Univariate Predictors	Variables Included in Multivariate Models	Results	Funding Source
Yu, 2006 ⁸¹	Interferon monotherapy: 28%	Univariate analyses not	Age, sex, ALT, genotype,	SVR vs. untreated and no	Department of Health,
Taiwan	Interferon plus ribavirin combination therapy: 72%	reported	interferon monotherapy or interferon plus ribavirin	SVR vs. untreated Hepatocellular carcinoma:	Taiwan and Taiwan Liver Research Foundation
Continued			combination therapy	Adjusted HR 0.25 (0.13-0.46) and 0.99 (0.64-1.5) All-cause mortality: Adjusted HR 0.37 (0.14-0.99) and 1.3 (0.56-3.1)	
				SVR vs. no SVR# Hepatocellular carcinoma: Adjusted HR 0.25 (0.13-0.50) All-cause mortality: 0.28 (0.08-1.0)	

Evidence Table 10. Quality rating: Studies on sustained virologic response and clinical outcomes

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 use accurate methods for ascertaining exposures, potential confounders, and outcomes?	(4) Were outcome assessors and/or data analysts blinded to treatment?	(5) Did the article the number of patients who met inclusion criteria excluded due to missing data or loss to followup?	(6) Did the study perform appropriate statistical analyses on potential confounders (should evaluate at least age, sex, genotype, fibrosis stage, viral load)?	(7) Is there important (overall or differential) exclusion of patients due to missing data or loss to followup?	(8) Were outcomes prespecified and defined, and ascertained using accurate methods?	Overall Quality (good, fair, poor)
Arase, 2007 ⁶³	Yes	No	Yes	Unclear	No	Yes	Unclear	Yes	Fair
Backus, 2011 ⁶⁴	Yes	No	Yes	No	Yes	Yes	Yes	Yes	Fair
Bruno, 2007 ⁶⁵	Yes	No	Yes	Unclear	No	No	Unclear	Yes	Fair
Cardoso, 2010 ⁶⁶	Yes	No	Yes	Unclear	No	Yes	Unclear	Yes	Fair
Coverdale, 2004 ⁶⁷	Unclear	No	Unclear	No	No	Unclear	Unclear	Yes	Poor
El Braks, 2007 ⁶⁸	Yes	No	Yes	Unclear	No	No	Unclear	Yes	Poor
Fernandez-Rodriguez, 2010 ⁶⁹	Unclear	No	Yes	No	Yes	Unclear	No	Yes	Poor
Hasegawa, 2007 ⁷⁰	Unclear	Unclear	Yes	Unclear	No	Yes	Unclear	Yes	Fair
Hung, 2006 ⁷¹	Yes	No	Yes	Unclear	No	Yes	Unclear	Yes	Fair
Imazeki, 2003 ⁷²	Yes	No	Yes	Unclear	Yes	Yes	No	Yes	Fair
Innes, 2011 ⁷³	Yes	No	Yes	Unclear	Yes	Yes	No	Yes	Fair
Izumi, 2005 ⁷⁴	Yes	Unclear	Yes	Unclear	No	Unclear	Unclear	Yes	Fair
Kasahara, 2004 ⁷⁵	No	Yes	Yes	Unclear	No	No	Unclear	Yes	Poor
Maruoka, 2012 ⁷⁶	Yes	Unclear	Yes	Unclear	Yes	Yes	Yes	Yes	Fair
Morgan, 2010 ⁷⁷	Yes	No	Yes	Unclear	No	Unclear	Unclear	Yes	Fair
Shiratori, 2005 ⁷⁸	Unclear	Yes	Yes	Unclear	Yes	No	Yes	Yes	Poor
Veldt, 2007 ⁷⁹	Yes	No	Yes	No	No	No	Unclear	Yes	Poor
Yoshida, 2002 ⁸⁰	Yes	No	Yes	No	No	No	Unclear	Yes	Poor
Yu, 2006 ⁸¹	Yes	No	Yes	No	No	No	Unclear	Yes	Poor

Evidence Table 11. Sustained virologic response and clinical outcomes summary results

Author, Year Country Quality	Study Type Number Analyzed Duration of Followup Proportion with Cirrhosis: SVR vs. no SVR	Hepatocellular Carcinoma: Adjusted Hazards Ratio (95% CI)	Liver-Related Mortality: Adjusted Hazard Ratio (95% CI)	All-Cause Mortality: Adjusted Hazard Ratio (95% CI)	Other Clinical Outcomes: Adjusted Hazard Ratio (95% CI)	Results Adjusted for at Least Age, Sex, Viral Load, Genotype, and Fibrosis Stage, or no Association Found in Univariate Analyses
Studies of general populations of treated patients with HCV infection						
Arase, 2007 ⁶³ Japan Overall Quality: Fair	Retrospective cohort n=500 Mean 7.4 years Cirrhosis: 9% vs. 16%	SVR vs. no SVR: 0.19 (0.08-0.45)	SVR vs. no SVR: 0.13 (0.03-0.59)	SVR vs. no SVR: 0.39 (0.16-0.93)	NR	Yes
Backus, 2011 ⁶⁴ # USA Overall Quality: Fair	Retrospective cohort n=16,864 Median 3.8 years Cirrhosis: 9-12% vs. 12- 20%	NR	NR	SVR vs. no SVR (genotypes 1, 2, and 3, respectively): 0.71 (0.60-0.86), 0.62 (0.44-0.87), and 0.51 (0.35-0.75)	NR	Yes
Coverdale, 2004 ⁶⁷ * Australia Overall Quality: Poor	Prospective cohort (some patients originally enrolled in randomized trials) n=343 Median 9 years Cirrhosis: Not reported, median fibrosis score F2 (Scheuer)	SVR vs. response- relapse vs. nonresponse Adjusted HR not reported (p>0.05)	SVR vs. response- relapse vs. nonresponse Liver transplant or liver-related death: Adjusted HR not reported (p=0.20)	NR	SVR vs. response- relapse vs. nonresponse Liver-related complications:** Adjusted HR not reported (p=0.06)	Unclear
Imazeki, 2003 ⁷² Japan Overall Quality: Fair	Retrospective cohort n=459 Mean 8.2 years Cirrhosis: 13% overall	NR	SVR vs. no SVR: 0.11 (0.01-0.96)##	SVR vs. no SVR: 0.12 (0.01-1.3)##	NR	Yes
Innes, 2011 ⁷³ UK Overall Quality: Fair	Retrospective cohort n=1215 Mean 5.3 years Cirrhosis: 10% vs. 18%	NR	SVR vs. no SVR: 0.22 (0.09-0.58)	NR	SVR vs. no SVR Liver-related hospital episode: 0.22 (0.15- 0.34)	Yes

Author, Year Country Quality	Study Type Number Analyzed Duration of Followup Proportion with Cirrhosis: SVR vs. no SVR	Hepatocellular Carcinoma: Adjusted Hazards Ratio (95% CI)	Liver-Related Mortality: Adjusted Hazard Ratio (95% CI)	All-Cause Mortality: Adjusted Hazard Ratio (95% CI)	Other Clinical Outcomes: Adjusted Hazard Ratio (95% CI)	Results Adjusted for at Least Age, Sex, Viral Load, Genotype, and Fibrosis Stage, or no Association Found in Univariate Analyses
Izumi, 2005 ⁷⁴ Japan Overall Quality: Fair	Cohort study, appears retrospective n=495 Duration of followup: Not reported Cirrhosis: 5.1% overall	SVR vs. no SVR: 0.36 (0.04-0.83)	NR	NR	NR	Unclear
Kasahara, 2004 ⁷⁵ Japan Overall Quality: Poor	Retrospective cohort n=2698 Mean 6 years Cirrhosis: 3.0% vs. 5.4%	NR	SVR vs. no SVR: 0.04 (0.005-0.30)	SVR vs. no SVR: 0.14 (0.06-0.35)	NR	No
Maruoka, 2012 ⁷⁶ Japan Overall Quality: Fair	Retrospective cohort n=577 Mean 9.9 years Cirrhosis: 10% overall	SVR vs. no SVR: 0.12 (0.04-0.40)##	NR	SVR vs. no SVR: 0.20 (0.08-0.54)##	NR	Yes
Yoshida, 2002 ⁸⁰ Japan Overall Quality: Poor	Retrospective cohort n=2889 Mean 5.4 years Cirrhosis: 6.5% vs. 11%	NR	SVR vs. no SVR: 0.13 (0.02-0.66)##	SVR vs. no SVR: 0.32 (0.12-0.86)##	NR	No
Yu, 2006 ⁴² Taiwan Overall Quality: Poor	Retrospective cohort n=1057 Mean 5.2 years Cirrhosis: 16% overall	S SVR vs. no SVR: 0.25 (0.13-0.54)##	NR	SVR vs. no SVR: 0.28 (0.08-1.0)##	NR	No
Studies of populations with advanced fibrosis and cirrhosis			1			
Bruno, 2007 ⁶⁵ Italy Overall Quality: Fair	Retrospective cohort study n=883 Mean 8 years Cirrhosis: All	SVR vs. no SVR: 0.39 (0.17-0.88)	SVR vs. no SVR: 0.14 (0.04-0.59)	NR	SVR vs. no SVR Ascites, encephalopathy, or gastrointestinal bleeding: Not calculated, 0 events/1061 person- years vs. 107 events/5703 person- years (1.88 events/100 person-years)	No

Author, Year Country Quality	Study Type Number Analyzed Duration of Followup Proportion with Cirrhosis: SVR vs. no SVR	Hepatocellular Carcinoma: Adjusted Hazards Ratio (95% CI)	Liver-Related Mortality: Adjusted Hazard Ratio (95% CI)	All-Cause Mortality: Adjusted Hazard Ratio (95% CI)	Other Clinical Outcomes: Adjusted Hazard Ratio (95% CI)	Results Adjusted for at Least Age, Sex, Viral Load, Genotype, and Fibrosis Stage, or no Association Found in Univariate Analyses
Cardoso, 2010 ⁶⁶ France Overall Quality: Fair	Retrospective cohort study (of patients originally enrolled in clinical trials) n=307 Median 3.5 years Cirrhosis: 53% vs. 61%	SVR vs. no SVR: 0.33 (0.23-0.89)	SVR vs. no SVR: 0.27 (0.08-0.95)	NR	SVR vs. no SVR Ascites or variceal bleeding: 0.21 (0.05- 0.92)	Yes
El Braks, 2007 ⁶⁸ France Overall Quality: Poor	Retrospective cohort study n=113 Mean 7.7 years Cirrhosis: All	NR	NR	NR	SVR vs. no SVR Clinical events (hepatocellular cancer, ascites, hepatic encephalopathy, or death): 0.14 (0.04-0.45)	No
Fernandez-Rodriguez, 2010 ⁶⁹ # Spain Overall Quality: Poor	Retrospective cohort study n=509 Median 35 months Cirrhosis: All	NR	NR	NR	SVR vs. no SVR Combined clinical endpoint:*** 0.38 (0.18- 0.76)	Unclear
Hasegawa, 2007 ⁷⁰ ^ Japan Overall Quality: Fair	Retrospective cohort study n=105 Median 4.6 years Cirrhosis: All	SVR vs. no SVR: 0.18 (0.04-0.81)	NR	NR	NR	Yes
Hung, 2006 ⁷¹ Taiwan Overall Quality: Fair	Cohort study (unclear if retrospective or prospective) n=132 Median 37 months Cirrhosis: All	SVR vs. no SVR: 0.28 (0.09-0.92)	NR	NR	NR	Yes
Morgan, 2010 ⁷⁷ # USA Overall Quality: Fair	Prospective cohort study of patient enrolled in a randomized trial n=526 Median 79 to 86 months Cirrhosis: 21% vs. 43%	SVR vs. no SVR: 0.19 (0.04-0.80)	SVR vs. no SVR Liver-related mortality or liver transplantation: 0.12 (0.03-0.48)	SVR vs. no SVR All-cause mortality or liver transplantation: 0.17 (0.06-0.46)	SVR vs. no SVR Any liver-related outcome:^^ 0.15 (0.06- 0.38) Decompensated liver disease: 0.13 (0.03- 0.53)	Unclear

Author, Year Country Quality	Study Type Number Analyzed Duration of Followup Proportion with Cirrhosis: SVR vs. no SVR	Hepatocellular Carcinoma: Adjusted Hazards Ratio (95% CI)	Liver-Related Mortality: Adjusted Hazard Ratio (95% CI)	All-Cause Mortality: Adjusted Hazard Ratio (95% CI)	Other Clinical Outcomes: Adjusted Hazard Ratio (95% CI)	Results Adjusted for at Least Age, Sex, Viral Load, Genotype, and Fibrosis Stage, or no Association Found in Univariate Analyses
Shiratori, 2005 ⁷⁸	Prospective cohort study		NR		NR	No
Japan	of patients enrolled in	SVR vs. no SVR:		SVR vs. no SVR: 0.07		
Overall Quality Deer	randomized trials	0.40 (0.18-0.89)##		(0.01-0.56)##		
Overall Quality: Poor	n=271 Median 6.8 years					
	Cirrhosis: All					
Veldt, 2007 ⁷⁹	Retrospective cohort	SVR vs. no SVR:	SVR vs. no SVR:	SVR vs. no SVR: 0.31	SVR vs. no SVR	No
Europe and Canada	n=479	0.46 (0.12-1.7)	0.19 (0.02-1.4)	(0.07-1.4)	Any event (death, liver	
	Median 2.1 years				failure, and	
Overall Quality: Fair	Cirrhosis: 71% vs. 77%				hepatocellular cancer):	
					0.20 (0.07-0.58)	

Abbreviations: HCV, hepatitis C virus; NR, not reported; SVR, sustained virologic response.

Note: SVR defined in all studies as undetectable HCV RNA in serum 6 months after the end of antiviral therapy, except as noted.

^{*} SVR defined as undetectable HCV RNA on at least 2 occasions at least 2 years after completion of therapy.

[^] Duration of undetectability to meet criteria for SVR not reported.

[#] Study primarily evaluated patients who received pegylated interferon plus ribavirin.

^{**} Hepatic decompensation, complications of portal hypertension, hepatocellular carcinoma, liver transplantation, and liver-related mortality.

^{***} Hepatic decompensation, upper gastrointestinal bleeding secondary to rupture of esophageal or gastric varices, hepatocellular carcinoma, liver transplantation, and liver-related or liver-unrelated mortality.

^{^^} Decompensated liver disease (ascites, variceal bleeding, hepatic encephalopathy, spontaneous bacterial peritonitis), hepatocellular carcinoma, liver transplantation, and liver-related mortality.

^{##} Calculated from estimates for SVR vs. untreated and no SVR vs. untreated

Evidence Table 12. Studies on sustained virologic response and quality of life

Author, Year Country			Duration of	Inclusion		Number Screened/ Eligible/
Quality	Study Type	Comparison	Followup	Criteria	Exclusion Criteria	Enrolled/ Analyzed
Arora, 2006 ⁸² Australia, Europe, New Zealand, North America, and South America Overall Quality: Poor	Cohort study (patients enrolled in an randomized trial)	SVR vs. no SVR SVR=No detectable HCV RNA at end of followup (72 weeks)	72 weeks	No prior treatment for chronic hepatitis C infection, positive HCV RNA, normal ALT	Cirrhosis, other chronic liver disease, HIV infection, other serious chronic illness, pregnancy	Number screened: Not reported Number eligible: Not reported Number enrolled: 440 (randomized to an antiviral treatment arm) Number analyzed: Unclear

Author, Year Country Quality	Demographic Characteristics of Study Population (Age, Race, Mean Viral Load)	Genotype HCV Viral Load HIV Infection IV Drug Use	Treatments	Confounders Assessed in Analysis	Results (by Clinical Outcome)	Funding Source
Arora, 2006 ⁸² Australia, Europe, New Zealand, North America, and South America Continued	Not reported by SVR status Mean age: 43 years Female: 60% Non white: 14%	Not reported by SVR status Advanced fibrosis: 10% Genotype 1: 68% Viral load: 1.1-1.2 x 10 ⁶ copies/ml IVDU: 30% HIV positive: excluded	Pegylated interferon alfa-2a (24 or 48 weeks)	Genotype, country, treatment, fibrosis stage, baseline score	SVR vs. no SVR, mean difference in change from baseline SF-36 physical function: +4.7 (p<0.05) SF-36 role limitations-physical: +13 (p<0.05) SF-36 bodily pain: +11 (p<0.0001) SF-36 general health: +10 (p<0.0001) SF-36 vitality: +9.3 (p<0.0001) SF-36 social function: +5.1 (p>0.05) SF-36 role limitations-emotional: +7.3 (p>0.05) SF-36 mental health: +3.1 (p>0.05) SF-36 physical component summary: +4.9 (p<0.0001) SF-36 mental component summary: +2.0 (p>0.05) Fatigue Severity Scale, total score: -4.4 (p<0.01) Fatigue Severity Scale, VAS: -10 (p<0.001)	Roche Pharmaceuticals

Author, Year Country Quality	Study Type	Comparison	Duration of Followup	Inclusion Criteria	Exclusion Criteria	Number Screened/ Eligible/ Enrolled/ Analyzed
Bernstein, 2002 ⁸³ Australia, North America, Europe, Taiwan, New Zealand Overall Quality: Poor	Cohort study (patients originally enrolled in 3 randomized trials)	SVR vs. no SVR SVR=No detectable HCV RNA 24 weeks after completion of antiviral therapy	72 weeks	Not previously treated with interferon- based therapies, positive HCV antibody, elevated serum ALT level,	Other chronic liver disease, significant co morbid conditions, pregnancy, evidence of substance abuse within 1 year	Number screened: Not reported Number eligible: Not reported Number enrolled: 1441 Number analyzed: 983 (275 SVR, 708 no SVR)
				positive HCV RNA		

Author, Year Country Quality	Demographic Characteristics of Study Population (Age, Race, Mean Viral Load)	Genotype HCV Viral Load HIV Infection IV Drug Use	Treatments	Confounders Assessed in Analysis	Results (by Clinical Outcome)	Funding Source
Bernstein, 2002 ⁸³ Australia, North America, Europe, Taiwan, New Zealand Continued	Not reported by SVR status Mean age <=40 years: 41% Female: 32% Non white: 14%	Not reported by SVR status Cirrhosis: 32% Genotype, viral load, HIV infection, IV drug use not reported	Pegylated interferon alfa-2a or interferon alfa-2a 2a	None	SVR vs. no SVR, mean difference in change from baseline SF-36 physical function: +4.6 (p<0.001) SF-36 role limitations-physical: +9.8 (p<0.001) SF-36 bodily pain: +2.9 (p<0.01) SF-36 general health: +9.1 (p<0.001) SF-36 vitality: +9.6 (p<0.001) SF-36 social function: +6.2 (p<0.001) SF-36 role limitations-emotional: +8.4 (p<0.01) SF-36 mental health: +4.6 (p<0.001) SF-36 physical component summary: +2.8 (p<0.001) SF-36 mental component summary: +3.0 (p<0.001) Fatigue Severity Scale, total score: -0.5 (p<0.001) Fatigue Severity Scale, VAS: -11.5 (p<0.001)	F. Hoffman-La Roche

Author, Year Country Quality	Study Type	Comparison	Duration of Followup	Inclusion Criteria	Exclusion Criteria	Number Screened/ Eligible/ Enrolled/ Analyzed
Bini, 2006 ⁸⁴	Prospective cohort	SVR vs. no SVR	48 or 72 weeks	No prior	HBV infection, HIV infection,	Number screened:
USA	study	SVR=No detectable HCV	(24 weeks after	treatment for	neutropenia, anemia,	Not reported
		RNA 24 weeks after	end of treatment)	chronic	thrombocytopenia, renal insufficiency,	Number eligible: Not
Overall Quality: Poor		completion of antiviral		hepatitis C	AFP >50 ng/ml, decompensated	reported
		therapy		infection,	cirrhosis, prior organ transplantation,	Number enrolled: 138
				positive HCV	cancer, severe co morbid condition,	(46 normal ALT, 92
				antibody,	poorly controlled diabetes or thyroid	elevated ALT)
				positive HCV RNA, liver	disease, autoimmune disease, seizure disorder, concurrent	Number analyzed: 138
				biopsy	immunosuppressive therapy, more	130
				consistent with	than 10 g alcohol/day or illicit drugs	
				chronic HCV	within 6 months	
				infection	Within 6 months	
				Each patient		
				with normal		
				ALT matched		
				with 2 patients		
				with elevated		
				ALT on		
				genotype, HCV		
				viral load, and		
				presence of		
				cirrhosis		

Author, Year Country Quality	Demographic Characteristics of Study Population (Age, Race, Mean Viral Load)	Genotype HCV Viral Load HIV Infection IV Drug Use	Treatments	Confounders Assessed in Analysis	Results (by Clinical Outcome)	Funding Source
Bini, 2006 ⁸⁴ USA Continued	Normal ALT and elevated ALT groups, respectively (not reported by SVR status)	Normal ALT and elevated ALT groups, respectively (not reported by SVR	Interferon alfa-2b + ribavirin	None	SVR vs. no SVR, mean difference in change from baseline (normal ALT and elevated ALT subgroups, respectively; p values not reported)	No external funding
Continued	Mean age: 50 and 49 years Female: 11% and 8% Non white: 59% and 66%	status) Cirrhosis: 11% and 11% Genotype 1: 78% and 78% Viral load >2 x 10 ⁶ copies/ml: 44% and 44% IVDU: 67% and 65% HIV positive: excluded			SF-36 physical function: +18 and +15 SF-36 role limitations-physical: +22 and +27 SF-36 bodily pain: +3.4 and +9.3 SF-36 general health: +3.0 and +9.9 SF-36 vitality: +12 and +12 SF-36 social function: +9.5 and +11 SF-36 role limitations-emotional: +20 and +18 SF-36 mental health: +14 and +18 SF-36 physical component summary: +3.8 and +7.1 SF-36 mental component summary: +6.0 and +2.1 Positive well being: +14 and -3.1 Sleep somnolence: +11 and +5.4 Health distress: +9.3 and +11 Hepatitis-specific health distress: +5.4 and +2.6 Hepatitis-specific limitations: +13 and +3.8	

Author, Year Country	0. 1. 7		Duration of	Inclusion		Number Screened/ Eligible/
Quality	Study Type	Comparison	Followup	Criteria	Exclusion Criteria	Enrolled/ Analyzed
Bonkovsky, 1999 ⁸⁵	Cohort study	SVR vs. no SVR	72 weeks	Positive HCV	Malignancy, depressive illness, HIV	Number screened:
USA and Canada	(patients enrolled in	SVR=No detectable HCV		antibody,	infection, decompensated liver	Not reported
	a randomized trial)	RNA 24 weeks after		positive HCV	disease, previous use of interferon,	Number eligible: Not
Overall Quality: Poor`		completion of antiviral		RNA, ALT >1.5	previous use of chemotherapeutic of	reported
		therapy		times upper	other agents, thyroid abnormality	Number enrolled: 704
				limit of normal,		Number analyzed:
				liver biopsy		437 (41 SVR, 396 no
				confirming		SVR)
				diagnoses of		
				chronic		
				hepatitis		

Author, Year Country Quality	Demographic Characteristics of Study Population (Age, Race, Mean Viral Load)	Genotype HCV Viral Load HIV Infection IV Drug Use	Treatments	Confounders Assessed in Analysis	Results (by Clinical Outcome)	Funding Source
Bonkovsky, 1999 ⁸⁵	Not reported by SVR	Not reported by SVR	Consensus	None	SVR vs. no SVR, mean difference in change	Amgen Inc.;
USA and Canada	status	status	interferon or		from baseline (values estimated from graph)	United States
	Mean age: 43 years	Cirrhosis: 16%	interferon alfa-2b		SF-36 physical function: +6.0 (p<0.05)	Public Health
Continued	Female: 27%	Genotype 1: 68%			SF-36 role limitations-physical: +22 (p<0.01)	Service
	Non white: 23%	Viral load: Not			SF-36 bodily pain: -0.5 (p>0.05)	
		reported			SF-36 general health: +7.5 (p<0.01)	
		IVDU: 41%			SF-36 vitality: +9.5 (p<0.05)	
		HIV positive:			SF-36 social function: +10 (p<0.05)	
		excluded			SF-36 role limitations-emotional: +11	
					(p>0.05)	
					SF-36 mental health: +4.0 (p>0.05)	

Author, Year Country Quality	Study Type	Comparison	Duration of Followup	Inclusion Criteria	Exclusion Criteria	Number Screened/ Eligible/ Enrolled/ Analyzed
Hassanein, 2004 ⁸⁶	Cohort study	SVR vs. no SVR	72 weeks	No prior	Neutrophils <1500 per cubic	Number screened:
Australia, North	(patients enrolled in	SVR=No detectable HCV		interferon,	millimeter, platelets <90.000 per cubic	1459
America, Europe,	a randomized trial)	RNA 24 weeks after		HCV RNA	millimeter, hemoglobin <12 g/dl in	Number eligible: Not
Taiwan, Brazil, Mexico		completion of antiviral		>=2000	women or <13 g/dl in men, HIV	reported
		therapy		copies/ml, ALT	infection, decompensated liver	Number enrolled:
Overall Quality: Poor				>upper limit of	disease, serum creatinine >1.5 times	1149
-				normal, liver	upper limit of normal, poorly controlled	Number analyzed:
				biopsy	psychiatric disease, alcohol or drug	649
				consistent with	dependence within one year before	
				chronic	study entry, substantial coexisting	
				hepatitis C	medical conditions	

Author, Year Country Quality	Demographic Characteristics of Study Population (Age, Race, Mean Viral Load)	Genotype HCV Viral Load HIV Infection IV Drug Use	Treatments	Confounders Assessed in Analysis	Results (by Clinical Outcome)	Funding Source
Hassanein, 2004 ⁸⁶ Australia, North America, Europe, Taiwan, Brazil, Mexico Continued	Not reported by SVR status Mean age: 43 years Female: 29% Non white: 16%	Not reported by SVR status Cirrhosis: 13% Genotype 1: 63% Viral load: 5.9 to 6.0 x 10 ⁶ copies/ml IVDU: Not reported HIV positive: excluded	Pegylated interferon alfa-2a, pegylated interferon alf-2a +ribavirin, or interferon alfa-2b + ribavirin	None	SVR vs. no SVR, mean difference in change from baseline SF-36 physical function: +5.5 (p<0.01) SF-36 role limitations-physical: +5.7 (p<0.05) SF-36 bodily pain: +4.1 (p<0.05) SF-36 general health: +8.6 (p<0.01) SF-36 vitality: +6.3 (p >0.05) SF-36 social function: +5.8 (p<0.01) SF-36 role limitations-emotional: +9.3 (p<0.01) SF-36 mental health: +5.0 (p<0.01) SF-36 physical component summary: +2.2 (p<0.01) SF-36 mental component summary: +2.6 (p<0.01) Total fatigue: +3.3 (p<0.01) Fatigue severity: +7.4 (p<0.01)	Roche Pharmaceuticals

Author, Year			-			Number Screened/
Country			Duration of	Inclusion		Eligible/
Quality	Study Type	Comparison	Followup	Criteria	Exclusion Criteria	Enrolled/ Analyzed
McHutchison, 200187	Cohort study	SVR vs. relapse vs. non	72 weeks	Positive HCV	Decompensated cirrhosis, AFP >50	Number screened:
USA	(patients enrolled in	responder		RNA, liver	ng/ml, anemia (hemoglobin <12 g/dl in	1337
	a randomized trial)	SVR=No detectable HCV		biopsy	women and <13 g/dl in men), HIV	Number eligible: 933
Overall Quality: Poor		RNA 24 weeks after		consistent with	infection, psychiatric conditions,	Number enrolled: 933
		completion of antiviral		chronic	seizure disorders, cardiovascular	Number analyzed:
		therapy		hepatitis,	disease, hemophilia, poorly controlled	824 (195 SVR, 150
		Relapse: Not defined		elevated serum	diabetes mellitus, autoimmune	relapse, 478 non
				ALT	diseases, s/p organ transplantation,	responder)
					unable to practice contraception	

Author, Year Country Quality	Demographic Characteristics of Study Population (Age, Race, Mean Viral Load)	Genotype HCV Viral Load HIV Infection IV Drug Use	Treatments	Confounders Assessed in Analysis	Results (by Clinical Outcome)	Funding Source
McHutchison, 2001 ⁸⁷	Mean age: 43 vs. 44	Cirrhosis: Not	Interferon alfa-2a	None	SVR and relapse, mean difference in	Schering-
USA	years Female: 42% vs. 32%	reported Genotype 1: 43% vs.	for 24 or 48 weeks, with or		change from baseline vs. non responder (p not reported, values estimated from graph)	Plough and Scripps Clinic
Continued	Non white: 8% vs. 12%	81% Viral load >2 million copies/ml: 58% vs. 74% IVDU: Not reported HIV positive: excluded	without ribavirin		SF-36 physical function: +2.4 and +0.8 SF-36 role limitations-physical: +5.2 and +3.2 SF-36 bodily pain: +1.6 and +1.7 SF-36 general health: +5.2 and +1.5 SF-36 vitality: +4.7 and +2.0 SF-36 social function: +3.1 and +0.4 SF-36 role limitations-emotional: +3.0 and +1.2 SF-36 mental health: +2.0 and 0.0 Sleep somnolence: +3.4 and +2.3 Health distress: +5.4 and +1.2 Hepatitis-related health distress: +5.7 and +1.1 Hepatitis-related limitations: +4.6 and +2.1	

Author, Year Country Quality	Study Type	Comparison	Duration of Followup	Inclusion Criteria	Exclusion Criteria	Number Screened/ Eligible/ Enrolled/ Analyzed
Neary, 1999 ⁸⁸	Cohort study	SVR vs. no SVR and	72 weeks (24	Chronic HCV	Women not using effective birth	Number screened:
USA, Europe, Australia	(patients enrolled in	overall response vs. no	weeks after end	infection,	control, decompensated cirrhosis,	495
	a randomized trial)	overall response	of treatment)	previously	anemia (hemoglobin <12 g/dl in	Number eligible:
Overall Quality: Poor		SVR=No detectable HCV		treated with	women and <13 g/dl in men), white	Unclear
		RNA 24 weeks after		one or two	blood cell count <3000 per cubic mm,	Number enrolled: 349
		completion of antiviral		courses of	neutrophil count <1500 per cubic mm,	Number analyzed:
		therapy		interferon	platelet count less than 100,000 per	Unclear (257 with
		Overall response=SVR		alpha with	cubic mm, HIV infection, prior organ	"complete data"
		plus >=2-point		relapse on	transplantation, severe psychiatric	
		improvement in Knodell		most recent	conditions, seizure disorder,	
		HAI score		course, liver	cardiovascular disease, renal	
				biopsy showing	insufficiency, hemoglobinopathy,	
				chronic	hemophilia, poorly controlled diabetes	
				hepatitis after	mellitus, immunologically mediated	
				relapse	diseases	

Author, Year Country Quality	Demographic Characteristics of Study Population (Age, Race, Mean Viral Load)	Genotype HCV Viral Load HIV Infection IV Drug Use	Treatments	Confounders Assessed in Analysis	Results (by Clinical Outcome)	Funding Source
	Race, Mean Viral Load) Not reported by SVR or overall response status Mean age: 43 years Female: 35% Non white: 6.4%	Not reported by SVR or overall response status Bridging fibrosis or cirrhosis: 17% Genotype 1: 56% Viral load >2 million copies/ml: 75% IVDU: 40% HIV positive: excluded	Interferon alfa-2b with or without ribavirin	None	SVR and relapse. mean difference in change from baseline vs. non responder (estimated from graph) (p values not reported) SF-36 physical function: +8.0 and +3.8 SF-36 role limitations-physical: +7.6 and +4.9 SF-36 bodily pain: +2.4 and +2.7 SF-36 general health: +9.4 and +5.6 SF-36 vitality: +7.8 and +5.6 SF-36 social function: +9.4 and +4.1 SF-36 role limitations-emotional: +6.0 and +12 SF-36 mental health: +2.8 and +1.8 Sleep somnolence: +2.1 and +3.8 Health distress: +8.9 and +1.6 Hepatitis-related health distress: +11 and -0.8 Hepatitis-related limitations: +6.7 and +2.6	Source Schering- Plough
					Mental health-18: +3.4 and +2.3 Overall response vs. no response (estimated from graph) SF-36 physical function: +8.3 (p<0.05) SF-36 role limitations-physical: +10 (p>0.05) SF-36 bodily pain: +3.7 (p>005) SF-36 general health: +6.9 (p<0.05) SF-36 vitality: +5.8 (p<0.05) SF-36 social function: +9.2 (p<0.05) SF-36 role limitations-emotional: +3.6 (p>0.05) SF-36 mental health: +1.3 (p>0.05) Sleep somnolence: +1.5 (p>0.05) Health distress: +6.4 (p<0.05) Hepatitis-related health distress: +12 (p<0.05) Hepatitis-related limitations: +7.8 (p<0.05) Mental health-18: +1.5 (p>0.05)	

Author, Year						Number Screened/
Country			Duration of	Inclusion		Eligible/
Quality	Study Type	Comparison	Followup	Criteria	Exclusion Criteria	Enrolled/ Analyzed

Author, Year Country			Duration of	Inclusion		Number Screened/ Eligible/
Quality	Study Type	Comparison	Followup	Criteria	Exclusion Criteria	Enrolled/ Analyzed
Rasenack, 200389	Cohort study	SVR vs. no SVR	72 weeks (24	Positive HCV	Prior interferon therapy, other disease	Number screened:
Germany, Canada,	(patients enrolled in	SVR=No detectable HCV	weeks after end	antibody,	of the liver or other major diseases,	Not reported
New Zealand, Spain	a randomized trial)	RNA 24 weeks after	of treatment)	positive HCV	pregnant, substance abuse within the	Number eligible: Not
		completion of antiviral		RNA,	last year	reported
Overall Quality: Poor		therapy		persistently		Number enrolled: 531
				elevated ALT,		Number analyzed:
				liver biopsies		Unclear
				consistent with		
				chronic		
				hepatitis C		

Author, Year Country Quality	Demographic Characteristics of Study Population (Age, Race, Mean Viral Load)	Genotype HCV Viral Load HIV Infection IV Drug Use	Treatments	Confounders Assessed in Analysis	Results (by Clinical Outcome)	Funding Source
Rasenack, 2003 ⁸⁹	Not reported by SVR	Not reported by SVR	Pegylated	None	SVR vs. no SVR, mean difference in change	F. Hoffman-La
Germany, Canada,	status	status	interferon alfa-2a		from baseline	Roche
New Zealand, Spain	Mean age: 41 years	Bridging	or interferon alfa-		SF-36 physical function: +5.0 (p=0.001)	
	Female: 33%	fibrosis/cirrhosis: 13%	2a		SF-36 role limitations-physical: +14	
Continued	Non white: 15%	Injection drug use:			(p<0.001)	
		37%			SF-36 bodily pain: +5.2 (p=0.014)	
		Viral load: 7.4 to 8.2 x			SF-36 general health: 12 (p<0.001)	
		10 ⁶ copies/ml			SF-36 vitality: +9.4 (p<0.001)	
		HIV positive: Not			SF-36 social function: +5.8 (p=0.005)	
		reported			SF-36 role limitations-emotional: +8.4	
		Genotype: Not			(p=0.02)	
		reported			SF-36 mental health: +5.3 (p=0.001)	
					SF-36 physical component summary: +3.2	
					(p<0.001)	
					SF-36 mental component summary: +2.9	
					(p=0.005)	
					Fatigue Severity Scale, total score: -0.5	
					(p=0.001)	
					Fatigue Severity Scale, VAS: -8.4 (p<0.001)	

Author, Year Country Quality	Study Type	Comparison	Duration of Followup	Inclusion Criteria	Exclusion Criteria	Number Screened/ Eligible/ Enrolled/ Analyzed
Ware, 1999 ⁹⁰ Australia, North America, and Europe Overall Quality: Poor	Cohort study (patients enrolled in a randomized trial)	SVR vs. no SVR SVR=No detectable HCV RNA 24 weeks after completion of antiviral therapy Overall response vs. no overall response Overall response=SVR + Knodell histology activity index inflammation score improved by 2 U or more	72 weeks (24 weeks after end of treatment)	Chronic HCV infection, relapsed after response to interferon treatment,	Decompensated cirrhosis, hemoglobin <12 g/dl in women and <13 g/dl in men, WBC <3000 per cubic millimeter, neutrophil count <1500 per cubic millimeter, platelet count <100,000 per cubic millimeter, HIV infection, prior organ transplantation, severe psychiatric conditions, seizure disorder, cardiovascular disease, renal insufficiency, hemoglobinopathy, hemophilia, poorly controlled diabetes mellitus, immunologically mediated	Number screened: 495 Number eligible: 349 Number enrolled: 349 Number analyzed: 250 (66 SVR and 184 no SVR)
		improved by 2 U or more			mellitus, immunologically mediated diseases	

Author, Year Country Quality	Demographic Characteristics of Study Population (Age, Race, Mean Viral Load)	Genotype HCV Viral Load HIV Infection IV Drug Use	Treatments	Confounders Assessed in Analysis	Results (by Clinical Outcome)	Funding Source
Ware, 1999 ⁹⁰ Australia, North America, and Europe Continued	Not reported by response status Mean age: 43 years Female: 35% Non white: 6.4%	Not reported by response status Bridging fibrosis/cirrhosis: 18% Injection drug use: 40% Viral load: 4.8 to 5.2 x 10 ⁶ copies/ml HIV positive: Excluded Genotype 1: 56%	Interferon alfa-2b or interferon alfa- 2b + ribavirin	None	SVR vs. no SVR and overall response vs. no overall response, mean difference in change from baseline (p values not reported) SF-36 physical function: +2.6 and +3.5 SF-36 role limitations-physical: +1.5 and +3.1 SF-36 bodily pain: +0.45 and +1.6 SF-36 general health: +3.3 and +3.5 SF-36 vitality: +2.2 and +2.8 SF-36 social function: +3.4 and +4.3 SF-36 role limitations-emotional: -0.02 and +1.1 SF-36 mental health: +1.3 and +0.62 Sleep: +0.02 and +1.2 Health distress: +7.6 and +6.2 Chronic hepatitis C health distress: +11.5 and +11.3 Chronic hepatitis C limitations: +5.3 and +7.5	Integrated Therapeutics Group, Inc (subsidiary of Schering- Plough)

Evidence Table 13. Quality rating: Studies on sustained virologic response and quality of life

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 use accurate methods for ascertaining exposures, potential confounders, and outcomes?	(4) Were outcome assessors and/or data analysts blinded to treatment?	(5) Did the article report attrition?	(6) Did the study perform appropriate statistical analyses on potential confounders (should adjust for at least age, sex, genotype, fibrosis stage)?	(7) Is there important differential loss to followup or overall high loss to followup?	(8) Were outcomes prespecified and defined, and ascertained using accurate methods?	Overall Quality
Arora, 2006 ⁸²	Yes	Unclear	Yes	No (patients aware of SVR status)	No	Yes	Unclear	Yes	Poor
Bernstein 2002 ⁸³	Yes	Unclear	Yes	No (patients aware of SVR status)	No	No	Unclear	Yes	Poor
Bini 2006 ⁸⁴	Unclear	Unclear	Yes	No (patients aware of SVR status)	No	No	Unclear	Yes	Poor
Bonkovsky 1999 ⁸⁵	Yes	Unclear	Yes	Yes (blinded to virological status, though not histological status)	Yes	No	Yes (high)	Yes	Poor
Hassanein 2004 ⁸⁶	Yes	Unclear	Yes	No (patients aware of SVR status)	Yes	No	Yes (high)	Yes	Poor
McHutchison 2001 ⁸⁷	Yes	No	Yes	Unclear	Yes	No	No	Yes	Poor
Neary 1999 ⁸⁸	Yes	Unclear	Yes	Unclear	Yes	No	Yes (high)	Yes	Poor
Rasenack 2003 ⁸⁹	Yes	Unclear	Yes	No (patient aware of SVR status)	Yes	No	Yes (high)	Yes	Poor
Ware 1999 ⁹⁰	Yes	Unclear	Yes	No (patient aware of SVR status)	Yes	No	Yes (high)	Yes	Poor

Evidence Table 14. Sustained virologic response and quality of life summary table scores

Author, Year Country	SF-36 Physical Function	SF-36 Role Limitations- Physical	SF-36 Bodily Pain	SF-36 General Health	SF-36 Vitality	SF-36 Social Function	SF-36 Role Limitations- Emotional	SF-36 Mental Health
Arora, 2006 ⁸² Australia, Europe, New Zealand, North America, and South America	+4.7 (p<0.05)	+13 (p<0.05)	+11 (p<0.0001)	+10 (p<0.0001)	+9.3 (p<0.0001)	+5.1 (p>0.05)	+7.3 (p>0.05)	+3.1 (p>0.05)
Bernstein, 2002 ⁸³ Australia, North America, Europe, Taiwan, New Zealand	+4.6 (p<0.001)	+9.8 (p<0.001)	+2.9 (p<0.01)	+9.1 (p<0.001)	+9.1 (p<0.001)	+6.2 (p<0.001)	+8.4 (p<0.01)	+4.6 (p<0.001)
Bini 2006 ⁸⁴ * USA	+18 and +15	+22 and +27	+3.4 and +9.3	+3.0 and +9.9	+12 and +12	+9.5 and +11	+20 and +18	+14 and +18
Bonkovsky 1999 ⁸⁵ USA and Canada	+6.0 (p<0.05)	+22 (p<0.01)	-0.5 (p>0.05)	+7.5 (p<0.01)	+9.5 (p<0.05)	+10 (p<0.05)	+11 (p>0.05)	+4.0 (p>0.05)
Hassanein, 2004 ⁸⁶ Australia, North America, Europe, Taiwan, Brazil, Mexico	+5.5 (p<0.01)	+5.7 (p<0.05)	+4.1 (p<0.5)	+8.6 (p<0.01)	+6.3 (p>0.05)	+5.8 (p<0.01)	+9.3 (p<0.01)	=5.0 (p<0.01)
McHutchison, 2001 ⁸⁷ ^ USA	+2.4	+5.2	+1.6	+5.2	+4.7	+3.1	+3.0	+2.0
Neary, 1999 ⁸⁸ ^# USA, Europe, Australia	+8.0	+7.6	+2.4	+9.4	+7.8	+9.4	+6.0	+2.8
Rasenack, 2003 ⁸⁹ ** Germany, Canada, New Zealand, Spain	+5.0 (p=0.001)	+14 (p<0.001)	+5.2 (p=0.014)	+12 (p<0.001)	+9.4 (p<0.001)	+5.8 (p=0.005)	+8.4 (p=0.02)	+5.3 (p=0.001)
Ware, 1999 ⁹⁰ ^ Australia, North America, and Europe	+2.6	+1.5	+0.45	+3.3	+2.2	+3.4	-0.02	+1.3

Author, Year Country Study Name	SF-36 Physical Component Summary	SF-36 Mental Component Summary	Sleep Somnolence	Fatigue Severity Scale, Total Score	Fatigue Severity Scale, Visual Analogue Scale		Hepatitis- Specific Health Distress	Hepatitis- Specific Limitations
Arora, 2006 ⁸² Australia, Europe, New Zealand, North America, and South America	+4.9 (p<0.0001)	+2.0 (p>0.05)	NR	+4.4 (p<0.01)	-10 (p<0.01)	NR	NR	NR
Bernstein, 2002 ⁸³ Australia, North America, Europe, Taiwan, New Zealand	+2.8 (p<0.001)	+3.0 (p>0.001)	NR	-0.5 (p<0.001)	-12 (p<0.001)	NR	NR	NR
Bini 2006 ⁸⁴ * USA	+3.8 and +7.1	+6.0 and +2.1	+11 and +5.4	NR	NR	+9.3 and +11	+5.4 and +2.6	+13 and +3.8
Bonkovsky 1999 ⁸⁵ USA and Canada	NR	NR	NR	NR	NR	NR	NR	NR
Hassanein, 2004 ⁸⁶ Australia, North America, Europe, Taiwan, Brazil, Mexico	+5.0 (p<0.01)	+2.6 (p<0.01)	NR	+3.3 (p<0.01)	+7.4 (p<0.01)	NR	NR	NR
McHutchison, 2001 ⁸⁷ ^ USA	NR	NR	+3.4	NR	NR	+5.4	+5.7	+4.6
Neary, 1999 ⁸⁸ ^# USA, Europe, Australia	NR	NR	+2.1	NR	NR	+8.9	+11	+6.7
Rasenack, 2003 ⁸⁹ ** Germany, Canada, New Zealand, Spain	+3.2 (p<0.001)	+2.9 (p=0.005)	NR	-0.5 (p=0.001)	-8.4 (p<0.001)	NR	NR	NR
Ware, 1999 ⁹⁰ ^ Australia, North America, and Europe	+0.02	None	NR	NR	NR	+7.6	+12	+5.3

Abbreviations: NR, not reported.

Note: Absence of p values indicates that they were not reported.

* Results reported for normal alanine transaminase and elevated alanine transaminase subgroups, respectively

^ Results for relapsers reported separately and excluded from table.

Same cohort as Ware, 1999.

** Cohort included in Bernstein, 2002.

Appendix H References

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