Introduction

Chronic infection with the hepatitis C virus (HCV) is one of the most common causes of liver cirrhosis and its sequelae in the Western world. Worldwide approximately 130–150 million people are infected. Due to hepatic and extrahepatic diseases, the risk of mortality is significantly increased in patients with HCV infection compared to seronegative persons [1]. A sustained virologic response (SVR) 12–24 weeks after treatment is the goal of antiviral therapy in order to prevent cirrhosis, hepatocellular carcinoma (HCC), and death [2]. Especially in cirrhotic patients, an SVR is associated with a significantly reduced 5-year mortality [3]. However, patients with liver cirrhosis had lower chances of SVR compared to non-cirrhotic patients in the era of pegylated interferon (peginterferon)/ribavirin treatment. Since 2011, antiviral therapy has changed because of the approval of the first direct-acting antiviral agents (DAA), i.e. telaprevir and boceprevir. Administered in combination with peginterferon alpha and ribavirin, these substances increased SVR rates in comparison to the dual combination of peginterferon alpha and ribavirin. However, augmented known and new adverse effects during treatment as well as SVR rates between 19 and 71% in difficult-to-treat patient groups limited the benefit for patients with cirrhosis [4–7]. Moreover, due to the administration of interferon alpha and the long duration of 48 weeks, the majority of patients with liver cirrhosis had contraindications. The next milestone in antiviral drug development was the approval of sofosbuvir, a nucleotide inhibitor of the HCV polymerase, in 2013/2014. This agent has a high barrier to antiviral drug resistance, rarely causes drug-drug interactions, and has a pan-genotypic efficacy. Therefore, it represents an ideal corner stone as a partner for combination with other DAA which have been approved in 2014, such as the NS3 protease inhibitor simeprevir or the NS5A inhibitors daclatasvir and ledipasvir. These combinations significantly increased SVR rates above 90% in virtually all groups of patients [8–12]. Comparable results can be achieved with a three-drug regimen containing the NS3 protease inhibitor paritaprevir/ritonavir, the NS5A inhibitor ombitasvir, and the non-nucleoside NS5B inhibitor dasabuvir, which was approved in 2014/2015 [13–16].

Keywords
Hepatitis C virus · Direct-acting antiviral agents · Liver cirrhosis · Sofosbuvir · Hepatocellular carcinoma

Summary

Background: Treatment of chronic hepatitis C infection is most urgent in patients with severe liver fibrosis and cirrhosis because of the high risk of decompensation, hepatocellular carcinoma, and consecutively death. The development and approval of several direct-acting antiviral drugs (DAA) in the past years has revolutionized antiviral therapy especially for patients with liver cirrhosis. Methods: This review will focus on recent data from clinical trials and recommendations for the therapy of hepatitis C-infected patients with compensated cirrhosis. Results: Clinical data for cirrhotic patients mainly exist for a combination of the nucleotide analog sofosbuvir with either a protease inhibitor (simeprevir) or an NS5A inhibitor (daclatasvir, ledipasvir) or a three-DAA combination consisting of an NS3 protease inhibitor, an NS5A inhibitor, and a non-nucleoside NS5B inhibitor (paritaprevir/ritonavir, ombitasvir, and dasabuvir). Rates of sustained virologic response in patients with compensated cirrhosis are comparable to patients without cirrhosis; however, the addition of ribavirin and/or longer treatment durations are especially recommended when other negative predictors are present, such as prior treatment failure, features of advanced cirrhosis, or the presence of baseline resistance. Conclusion: Nowadays, a highly active, short, and safe interferon-free treatment regimen is available for almost all patients.

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HCV treatment in the Western world are shown in table 1.

In general, the chance of achieving SVR with DAA combination therapies for patients with compensated cirrhosis is comparable to non-cirrhotic patients; however, there is a remaining risk for decompensation and acute-on-chronic liver failure during and after treatment [22]. Patients with advanced and decompensated cirrhosis should be treated in experienced centers, and the need for liver transplantation should be evaluated. In patients with end-stage liver disease and high short-term mortality, no clinical improvement might be visible despite viral eradication [23–25]. These patients may have a greater benefit from antiviral therapy after liver transplantation.

### Antiviral Therapy

The decision for or against a certain antiviral treatment regimen depends on several factors. Firstly, the HCV genotype has to be determined. The majority of available DAA combination regimens are active against HCV genotype 1 but differ in their activity against other HCV genotypes and subtypes. Secondly, prior antiviral therapies have to be considered. Patients with a relapse or non-response after treatment with peginterferon and ribavirin still have high chances of viral eradication. However, prior treatments with DAA may be associated with the selection of resistance-associated variants (RAV) which may influence treatment outcome of subsequent DAA combination regimens. Here, resistance analysis is recommended for the selection of an effective DAA combination, if available. Thirdly, the viral load in international units quantified via real-time polymerase chain reaction (PCR) is of importance prior to the beginning of treatment, when a short therapy over 8 weeks in patients without cirrhosis is considered. Forthly, concomitant medications have to be checked for potential drug-drug interactions ([www.hep-druginteractions.org](http://www.hep-druginteractions.org)).

Due to the short duration, improved tolerability and high efficacy as well as potential long-lasting side effects, interferon-free HCV treatment is the preferred recommendation [20, 21]. Nevertheless, in countries with restricted access to new antivirals, interferon-containing regimens can still be an option.

The indication for HCV treatment in HCV/HIV co-infected patients is identical to those in patients with HCV monoinfection. In general, the fibrosis progression in patients co-infected with HIV is accelerated. Several trials were conducted in co-infected patients, and the treatment outcomes were comparable to patients with monoinfection [26–29]. Some antiretroviral agents and combinations were co-administered in clinical trials. Therefore, potential drug-drug interactions have to be considered prior to treatment.

Table 2 gives an overview about the recommended and approved treatment options for the different genotypes for patients with cirrhosis. The corresponding SVR rates are illustrated in table 3.

In the following part of the review, the recommended antiviral therapies for patients with compensated cirrhosis are presented. HCV therapy for patients with decompensated cirrhosis is discussed by Welker and Zeuzem [30] in a separate article in this special issue.

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**Table 1. Available and approved substances for HCV treatment in the Western world**

| Peglated interferon alpha 2a and 2b |
| Ribavirin |
| NS3/4A protease inhibitors |
| First wave (associated with numerous side effects and high pill burden) |
| Telaprevir |
| Boceprevir |
| Second wave (improved tolerability and once-daily dosing) |
| Simeprevir |
| Paritaprevir (ritonavir-boosted) |
| NS5A inhibitors |
| Daclatasvir |
| Ledipasvir |
| Ombitasvir |
| Nucleotide NS5B polymerase inhibitor |
| Sofosbuvir |
| Non-nucleoside NS5B polymerase inhibitor |
| Dasabuvir |

The substances which are currently available and approved for HCV treatment in the Western world are shown in table 1.
**Table 2.** Treatment recommendations according to the different genotypes for patients with cirrhosis

| Treatment combination | Duration, weeks | TN  | TE  |
|-----------------------|----------------|-----|-----|
| **Genotype 1**      |                |     |     |
| SOF + SMV ± R        | 12 or 24       | app | app |
| SOF + DCV ± R        | 12 or 24       | app | app |
| SOF + LDV            | 12             | app | nrec|
| SOF + LDV + R        | 12             | rec | rec |
| SOF + LDV + R        | 24             | rec | rec |
| PTV/r + OMV + DSV   | 12             | app (1b)| app (1b)|
| PTV/r + OMV + DSV + R| 12             | rec (1b)| rec (1b)|
| PTV/r + OMV + DSV + R| 24             | rec (1a)| rec (1a)|
| **Genotype 2**      |                |     |     |
| SOF + R              | 12             | rec | rec (16–24 weeks) |
| SOF + P/R            | 12             | app | app |
| SOF + DCV ± R        | 12             | nrec| app |
| **Genotype 3**      |                |     |     |
| SOF + R              | 24             | rec | rec |
| SOF + P/R            | 12             | app | app |
| SOF + DCV + R        | 24             | app | rec |
| **Genotype 4**      |                |     |     |
| SOF + LDV + R        | 12             | rec | rec |
| SOF + LDV ± R        | 24             | app | app |
| PTV/r + OMV + R      | 24             | app | app |
| SOF + SMV ± R        | 12             | app | app |

Rec = Recommended as first-line therapy; app = approved, but not recommended as first-line therapy; nrec = not recommended; TN = treatment-naive; TE = treatment-experienced; SOF = sofosbuvir; SMV = simeprevir; R = ribavirin; P = pegylated interferon alpha; DCV = daclatasvir; LDV = ledipasvir; PTV/r = paritaprevir/ritonavir; OMV = ombitasvir; DSV = dasabuvir.

**Table 3.** SVR rates for the different treatment options in hepatitis C patients with (mainly compensated) cirrhosis

| Treatment combination | Duration, weeks | Genotype |
|-----------------------|----------------|----------|
|                        |                | 1        | 2        | 3        | 4        |
| SOF + R               | 12             | 83–94 (TN) | 34      | 43      |
| SOF + R               | 16             | 87      |
| SOF + R               | 24             | 100     | 60 (TE)–92 (TN) | 100     |
| SOF + SMV ± R         | 12/24          | 83 (w/o R) | 100     |
| SOF + DCV ± R         | 12             | 92      | 83–89   |
| SOF + DCV ± R         | 24             | 97      |
| SOF + LDV + R         | 12             | 96      |
| SOF + LDV             | 24             | 98      | 93–96   |
| SOF + LDV + R         | 24             | 100     |
| PTV/r + OMV + DSV    | 12             | 100 (1b) |
| PTV/r + OMV + DSV + R| 12             | 92      |
| PTV/r + OMV + DSV + R| 24             | 96      |
| PTV/r + OMV + R       | 12/24          | 96–100  |

TN = Treatment-naive; TE = treatment-experienced; SOF = sofosbuvir; SMV = simeprevir; R = ribavirin; DCV = daclatasvir; LDV = ledipasvir; PTV/r = paritaprevir/ritonavir; OMV = ombitasvir; DSV = dasabuvir.

**Genotype 1**

**Ledipasvir + Sofosbuvir ± Ribavirin**

The combination of the nucleotide HCV polymerase inhibitor sofosbuvir and ledipasvir, an NS5A inhibitor, has been evaluated in a broad phase III study program (ION studies) and has shown consistently high rates of sustained response in all groups of patients, including those with failure to boceprevir or telaprevir pre-treatment (>90%). This combination treatment is available as a fixed-dose formulation (400 mg sofosbuvir + 90 mg ledipasvir). An anal-
ysis of all cirrhotic patients treated in phase II and III studies (n > 500) demonstrated SVR rates of 92, 96, 98, and 100% for ledipasvir/sofosbuvir 12 weeks, ledipasvir/sofosbuvir + ribavirin 12 weeks, ledipasvir/sofosbuvir 24 weeks, and ledipasvir/sofosbuvir + ribavirin, respectively. The overall SVR rates were significantly lower in treatment-experienced patients with a platelet count <75,000/μl (82%) [31]. The SIRIUS trial prospectively evaluated ledipasvir/sofosbuvir + ribavirin for 12 weeks and ledipasvir/sofosbuvir + placebo for 24 weeks and found similar rates of SVR (96 and 97%) [32]. Thus, in order to reduce costs and duration of antiviral therapy, the EASL and the German guidelines recommend ledipasvir/sofosbuvir + ribavirin for 12 weeks as the favored option in patients with compensated cirrhosis. Due to an SVR rate of 96% with 12 weeks of sofosbuvir and ledipasvir in treatment-naive patients with cirrhosis, a ribavirin-free treatment is also an option. A longer duration of therapy can be considered when contraindications, intolerance to ribavirin, or negative predictors of a successful treatment outcome exist [20, 21].

Paritaprevir/Ritonavir + Ombitasvir + Dasabuvir + Ribavirin

The TURQUOISE-II trial was a phase III study evaluating paritaprevir/ritonavir + ombitasvir + dasabuvir + ribavirin over 12 or 24 weeks in patients with compensated cirrhosis. SVR rates were 92 and 96% for 12 and 24 weeks, respectively. Throughout all phase III studies the regimen showed higher efficacy in patients infected with HCV-subtype 1b compared to subtype 1a. Retrospective analysis showed reduced SVR rates in the 12-week treatment arm only in a subgroup of HCV genotype 1a-infected patients. Treatment failures in HCV genotype 1a-infected patients in the 12-week group were associated with an alpha-fetoprotein level > 20 ng/ml, a platelet count < 90,000/μl, or an albumin level < 35 g/l [15]. Therefore, paritaprevir/ritonavir + ombitasvir + dasabuvir + ribavirin should be administered as a standard treatment for all HCV genotype 1-infected patients with cirrhosis for 12 weeks, with the exception of HCV genotype 1a-infected patients with one of the negative predictive laboratory values mentioned above. In this case a treatment duration of 24 weeks is recommended. More recently, paritaprevir/ritonavir + ombitasvir + dasabuvir were investigated without ribavirin in patients with compensated cirrhosis and genotype 1b infection in a single-arm non-controlled study (TURQUOISE III) for 12 weeks. Based on an SVR rate of 100% [33], the ribavirin-free regimen can be recommended as a new standard treatment in this population.

Generally, there are no differences in antiviral efficacy for treatment-naive versus experienced patients. However, patients with failure to boceprevir or telaprevir triple therapies have not been studied, and currently it is unknown whether selection of viral resistance to NS5 protease inhibitors may impact treatment outcome.

The combination of paritaprevir/ritonavir + ombitasvir + dasabuvir should not be administered to patients with decompensated cirrhosis (Child-Pugh B or C) due to an observed increased rate of hepatic failure and decomposition in these patients.

Sofosbuvir + Simeprevir + Ribavirin

In cohort 2 of the COSMOS study, the combination of simeprevir with sofosbuvir was evaluated in treatment-naive patients and previous non-responders to peginterferon and ribavirin with advanced fibrosis or cirrhosis (METAVIR F3–4). Patients were assigned to 12 or 24 weeks of treatment either with or without ribavirin. In cohort 2, 93–100% SVR rates were achieved, and there were no beneficial effects of additional ribavirin or longer treatment durations [11]. Cirrhotic patients (treatment-naive and peginterferon/ribavirin-experienced) had an SVR rate of 83% with sofosbuvir + simeprevir over 12 weeks in the OPTIMIST-2 study [25]. In both studies, patients with virological relapse after treatment were in the majority genotype 1a patients, and the RAV Q80K was found at baseline (SVR 74 vs. 92% with and without Q80K mutation, respectively). Therefore, a resistance analysis can be useful in genotype 1a patients when a simeprevir-containing treatment is considered, especially when a protease inhibitor was given earlier.

Sofosbuvir + Daclatasvir + Ribavirin

The ALLY-1 study evaluated the treatment of sofosbuvir + daclatasvir + ribavirin over 12 weeks in cirrhotic patients (Child–Pugh A–C) as well as after liver transplantation and HCV recurrence. Patients with compensated cirrhosis achieved an SVR in 92% of cases [24]. Before the approval of the two components, a compassionate use program treated mainly cirrhotics with sofosbuvir + daclatasvir with and without ribavirin for 24 weeks. In this difficult-to-treat population high SVR rates were found (97% for patients with liver cirrhosis), but the numbers of patients with only 12 weeks of therapy were low [34]. The EASL guidelines favor a 12-week therapy with ribavirin in cirrhotic patients. When ribavirin is contraindicated, a treatment duration of 24 weeks is recommended.

Genotype 2

Sofosbuvir + Ribavirin

This treatment option was evaluated during several phase III trials in patients with genotype 2 infection. Patients with cirrhosis had lower SVR rates than patients without cirrhosis after 12 weeks of treatment (92–97 vs. 83–94%), but the numbers of cirrhotic patients were low (n = 12 and 17) [35, 36]. In previous interferon relapsers and non-responders with genotype 2 and cirrhosis, only 6 of 10 patients achieved SVR with this combination [36]. Real-world studies after approval in cirrhotic genotype 2 patients showed similar results with a 12-week treatment course (SVR 76–82%) [37, 38]. Therefore, a longer treatment duration was evaluated in the BOSON study: Cirrhotic genotype 2 patients achieved 87 and 100% SVR rates with 16 and 24 weeks of sofosbuvir + ribavirin, respectively [39]. To date, this data has not been implemented in current guidelines.

The analysis of treatment failures in genotype 2 after sofosbuvir + ribavirin treatment illustrated the possible existence of intergenotypic recombinant hepatitis C viruses. In these cases, viral chimerism of genotype 1 and 2 are identified as genotype 2 via the com-
monly used INNO-LiPA 2.0 assay, but their treatment responses were comparable to genotype 1 patients [40].

Most recently, results of a phase III study with the combination of sofosbuvir and the new NS5A inhibitor velpatasvir with high antiviral activity in HCV genotype 2 have been presented. All (19/19) patients with cirrhosis and genotype 2 infection achieved an SVR. However, the compared treatment sofosbuvir + ribavirin over 24 weeks also showed an SVR in 18/19 patients in this study [41].

**Sofosbuvir + Daclatasvir ± Ribavirin**

No systematically evaluated data exist for genotype 2 patients with cirrhosis for this treatment option. However, daclatasvir showed high antiviral efficacy in HCV genotype 2 isolates in vitro [42, 43], and its theoretical potency was demonstrated in a small phase II trial in patients without cirrhosis (SVR 96%) [12]. In genotype 2, this treatment should be reserved for patients who failed the standard treatment, i.e. sofosbuvir + ribavirin.

**Genotype 3**

**Sofosbuvir + Ribavirin**

In genotype 3 patients, rates of SVR were disappointing with 12 weeks of treatment, especially in patients with cirrhosis (SVR 34%) [35]. A treatment expansion up to 24 weeks demonstrated an improvement with 82–92% SVR in treatment-naive cirrhotic patients, but only 60–76% SVR in patients with liver cirrhosis and failure to prior peginterferon/ribavirin [39, 44]. Similar results have been obtained from real-world studies [37].

**Sofosbuvir + Pegylated Interferon Alpha + Ribavirin**

In the BOSON study, the expanded treatment with sofosbuvir + ribavirin was compared to a conventional triple therapy including peginterferon alpha. This treatment was significantly superior compared to sofosbuvir + ribavirin over 24 weeks alone (SVR 88 vs. 79%) [39]. Thus, peginterferon in combination with sofosbuvir and ribavirin remains a treatment option especially in difficult-to-treat genotype 3 patients who are interferon-tolerant.

**Sofosbuvir + Daclatasvir ± Ribavirin**

In contrast to patients without cirrhosis, sofosbuvir + daclatasvir therapy over 12 weeks demonstrated an SVR in only 63% of patients with cirrhosis [45]. Therefore, the ALLY3+ study compared sofosbuvir + daclatasvir + ribavirin for 12 and 16 weeks in patients with advanced fibrosis and compensated cirrhosis. SVR rates were comparable: Patients with cirrhosis achieved an SVR in 15/18 patients (83%) and 16/18 patients (89%) in the 12-week and 16-week treatment arm, respectively. Both relapers in the 16-week group were sofosbuvir-experienced [46]. Meanwhile, data from a European compassionate use program are available: Genotype 3 patients with an urgent need for antiviral therapy were treated in most cases with sofosbuvir + daclatasvir with or without ribavirin for 24 weeks. SVR rates were 87%, with the highest rates being found in compensated cirrhotics. The use of ribavirin was not associated with higher SVR rates; however, SVR rates were lower in a minority of patients who received therapy for 12 weeks only [47]. Because of missing alternative treatment options guidelines recommend a treatment expansion with the addition of ribavirin and a duration of 24 weeks.

Also in genotype 3, velpatasvir + sofosbuvir over 12 weeks demonstrated significantly higher SVR rates compared to sofosbuvir + ribavirin over 24 weeks (cirrhotics 91 vs. 66%) and will expand the treatment options for genotype 3 patients after approval (expected in 2016) [41].

**Genotype 4**

**Ledipasvir + Sofosbuvir ± Ribavirin**

In a small trial (n = 21 patients), ledipasvir/sofosbuvir for 12 weeks was evaluated in genotype 4 and achieved an SVR in 7/7 patients with cirrhosis [48]. Another study included 44 genotype 4 patients and demonstrated high SVR rates (93%) irrespective of prior treatment or fibrosis status [49]. Similar to the treatment in genotype 1 patients, current guidelines recommend ledipasvir/sofosbuvir and ribavirin for 12 weeks in case of underlying cirrhosis until further data is available.

**Paritaprevir/Ritonavir + Ombitasvir ± Ribavirin**

Dasabuvir has no significant antiviral activity in genotype 4; therefore, treatment with paritaprevir/ritonavir and ombitasvir is sufficient. In a recently published study, paritaprevir/ritonavir + ombitasvir + ribavirin was compared for 12 and 16 weeks in patients with cirrhosis. In both groups high SVR rates were achieved (96 and 100% for 12 and 16 weeks, respectively) [50]. The combination is approved for a 24-week treatment with ribavirin in patients with cirrhosis.

**Simeprevir + Sofosbuvir**

Recently, data from an Egyptian study were published. Genotype 4 patients with cirrhosis (n = 23) were treated with simeprevir + sofosbuvir for 12 weeks and achieved an SVR in all cases [51].

**Genotype 5 & 6**

Compared to HCV genotype 1–4, patients infected with HCV genotypes 5 and 6 are uncommon in the Western world. In a small trial, 8/9 cirrhotic HCV genotype 5 patients achieved an SVR with ledipasvir/sofosbuvir over 12 weeks [49]. Similar results were found in a trial with HCV genotype 6 patients (SVR 96%, only 2 patients with cirrhosis) [52].

Based on data of patients infected with genotype 1, cirrhotic patients should be treated with ledipasvir/sofosbuvir for 12 weeks and additional ribavirin, although this regimen is not formally approved for the use in patients infected with HCV genotype 5 or 6.
**Safety Aspects and Adverse Effects of Antivirals**

**Sofosbuvir**

Sofosbuvir is generally well tolerated. Common side effects are nausea and sleep disturbances. Severe and life-threatening brady- cardias were observed in combination with amiodarone. As a result, these agents must not be co-administered.

**Sofosbuvir + Ledipasvir**

Ledipasvir was investigated in combination with sofosbuvir only. Adverse events were tiredness and headache. Less than 2% of cirrhotic patients had serious adverse events during phase III studies [31]. Interestingly, the TARGET study showed a difference in SVR rates in patients who had a co-medication with a proton pump inhibitor (98 without vs. 93% with proton pump inhibitor) [53]. When possible, a co-administration should be avoided.

**Simeprevir**

Beside nausea, headache, and tiredness, a photosensitivity reaction of the skin can occasionally be present during treatment. It is important to inform the patient about limiting sun exposure and using sun protection. A hyperbilirubinemia is a common finding. In most cases it is mild, and severe hyperbilirubinemia was seen in <1% of cases.

**Daclatasvir**

The safety profile of daclatasvir is comparable to the other new DAA. Headache, fatigue, and nausea are the most common side effects during treatment.

**Paritaprevir/Ritonavir + Ombitasvir + Dasabuvir**

The most common adverse events found during approval studies were sleep disturbances, nausea, and itching. Elevations of bilirubin to more than three times the upper limit of normal were more often seen in cirrhotics (9.7%); relevant elevations of alanine aminotransferase were less common. Due to these possible hepatotoxic effects and an observed worsening of liver function in several patients with decompensated cirrhosis, this treatment cannot be recommended for decompensated patients.

**Ribavirin**

In several treatment combinations for patients with cirrhosis ribavirin is still mandatory. Its adverse effect profile is mainly caused by hemolytic anemia. Dose adjustments during therapy can be required for the management of anemia. Common symptoms are dyspnea, fatigue, and cough.

**Surveillance and Follow-up after SVR**

An SVR 12 weeks after the end of therapy is associated with a long-lasting cure in more than 99% of patients. An additional PCR testing 24 or 48 weeks after therapy can be performed although upon detectable HCV RNA at this time point, further analysis for assessment of late relapse versus re-infection should always be initiated. In patients with cirrhosis despite SVR, a significant risk for the development of HCC, cholangiocarcinoma, and hepatic decompensation is still present, and long-term surveillance is mandatory for years. However, patients with viral eradication showed a significant risk reduction in comparison to those without SVR [54]. After SVR an improvement of liver stiffness measurement values was observed with transient elastography, indicating a regression of fibrosis in the long-term follow-up [55]. Co-factors which can cause fibrosis progression, such as alcohol consumption and diabetes, have to be avoided. In cirrhotic patients, a regular surveillance for HCC by abdominal ultrasound and an alpha-feto-protein measurement every 6 months is recommended. The occurrence of esophageal varices after SVR is rare if varices were not present at pre-treatment endoscopy. After SVR, an endoscopic control for varices is recommended every 2 years after SVR [56]. Invasive measurements of hepatic venous pressure gradients before and after antiviral treatment showed a significant reduction after SVR, indicating a regression and partial normalization in most patients with portal hypertension [57].

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