Liver stiffness regression after sustained virological response by direct-acting antivirals reduces the risk of outcomes

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The role of liver stiffness measurement (LSM) after sustained virological response (SVR) in HCV patients treated by direct-acting antivirals (DAAs) remains unclear. We aimed to evaluate LSM regression value after SVR and to identify risk factors associated with liver related complications (LRC) or death. This retrospective study analyzed patients with LSM ≥ 10 kPa with LSM by transient elastography pre-DAAs and post-SVR. Patients with previous hepatic decompensation were excluded. Medical records were reviewed to identify primary outcomes. Kaplan–Meier curves and time-to-event Cox proportional-hazard models were performed. 456 patients [65% female, 62 years (IQR 57–68)] were included. During a follow-up of 2.3 years (IQR 1.6–2.7), 28 patients developed 37 outcomes [rate = 29.0 (95% CI 20.0–42.0) per 1000 person-years]. The cumulative incidence of outcomes was significantly lower in patients who regressed LSM ≥ 20% [3.4% (95% CI 1.8–7.0) vs. 9.0% (5.5–14.5), p = 0.028]. In a multivariate Cox-model [HR(95% CI)], male gender [HR = 3.00 (1.30–6.95), p = 0.010], baseline albumin < 3.5 mg/dL [HR = 4.49 (1.95–10.34), p < 0.001] and baseline unfavorable Baveno-VI [HR = 4.72 (1.32–16.83), p = 0.017] were independently associated and LSM regression ≥ 20% after SVR had a trend to reduce the risk of LRC or death [HR = 0.45 (0.21–1.02), p = 0.058]. The use of simple parameters before DAAs and repetition of LSM post-SVR can identify patients with different risks for severe outcome after HCV eradication.

Direct-acting antivirals (DAA) have changed the natural history of chronic hepatitis C in the last decade. These regimens are safe and highly effective, leading to rates of sustained virological response (SVR) higher than 90%. Therefore, international guidelines were updated to recommend DAA treatment to all individuals with chronic hepatitis C. A recent large cohort study demonstrated that treatment with DAAs is associated with reduced risk for mortality and hepatocellular carcinoma (HCC), reinforcing the long-term impact of SVR. However, the burden of liver-related events in patients with hepatitis C virus (HCV) that have successfully cleared the virus persists, particularly in individuals with cirrhosis previous to treatment.

Historically, liver biopsy remains the reference for liver fibrosis staging and detection of cirrhosis. However, this invasive method has been challenged by several limitations and performing a liver biopsy in SVR patients without abnormal liver tests might not be justified. Liver stiffness measurement (LSM) by transient elastography (TE) is the most validated non-invasive method to assess liver fibrosis in patients with ongoing HCV infection as an alternative to liver biopsy. LSM by TE can be used to predict severe outcomes in chronic hepatitis C: patients with LSM ≥ 9.5 kPa (advanced fibrosis/cirrhosis) had significantly lower overall survival compared to those in the lower range. Additionally, combining LSM values with platelet count in HCV-infected patients was extensively validated to identify high-risk esophageal varices.

Despite the broad validation of the diagnostic and prognostic value of LSM in patients with HCV, the correct interpretation of LSM in patients after SVR remains unclear. Several recent studies have reported a significant regression of LSM after SVR in patients with HCV treated by DAAs. However, it is still controversial whether the decrease of LSM after HCV eradication is related to suppression of viral necro-inflammatory activity rather than a loss of fibrous tissue.
than regression of liver fibrosis. Moreover, the extent of the long-term clinical impact of LSM reduction after SVR in patients with HCV treated by DAAs remains unclear because most trials had short follow-up periods and such severe outcomes are rare. Establishing an accurate longitudinal non-invasive strategy which could be used to predict severe outcomes after SVR would be clinically and epidemiologically important. The primary aim of this study was to evaluate the prognostic value of LSM regression after SVR by DAAs. The secondary aim was to identify risk factors associated with liver related complications (LRC) or death in patients with HCV after SVR.

Methods

Population and study design. This retrospective observational study was conducted at two tertiary centers for management of patients with viral hepatitis in Rio de Janeiro, Brazil. All adult patients with chronic hepatitis C treated by DAAs between October 2015 and November 2019 were eligible. The exclusion criteria were: (1) HCV treatment failure, loss of follow-up or missing data of SVR; (2) presence of hepatitis B coinfection, liver transplantation or hepatic decompensation (ascites, hepatic encephalopathy, variceal bleeding or HCC) before HCV treatment. (3) non-available LSM before or after DAA and (4) LSM < 10 kPa before HCV treatment. SVR was defined as undetectable HCV RNA at 12 weeks following the conclusion of treatment. LSM was assessed by TE by experienced operators using FIBROSCAN (Echosens, Paris, France). HCC screening and esophageal varices surveillance were performed according to the current international guidelines

Data collection. Demographic, clinical and biological data were collected by trained investigators using an electronic case-report form at Research Electronic Data Capture (REDCap, https://projectredcap.org/). Clinical features included comorbidities, data of HCV history/treatment and liver stiffness by TE before and after DAA therapy. Laboratory parameters (including liver enzymes, albumin, creatinine, fasting glucose and platelet count) performed at least three months before or after each LSM were collected. All medical records from included patients were reviewed by trained investigators to identify the prospective incidence of the following primary outcomes after SVR: LRC (ascites, hepatic encephalopathy, variceal bleeding, or HCC), liver transplantation or death.

Liver stiffness measurement. TE examinations were performed before and after HCV treatment in patients with a 3-h fasting by experienced operators. Liver fibrosis staging by TE was considered as reliable for analysis if the following criteria had been met: (1) at least 10 valid measurements; (2) an interquartile range (IQR) lower than 30% of the median of LSM and (3) a success rate of more than 60%. TE examination obtained closest to the date of start and the end of DAA treatment was considered for LSM before treatment and after SVR, respectively. Patients with LSM ≥ 10 kPa before HCV treatment were classified as compensated advanced chronic liver disease (c-ACLD). Additionally, those patients with LSM ≥ 20 kPa or platelet count < 150 × 10^9/mm3 before HCV treatment were considered as having an unfavorable Baveno VI status.

Statistical analysis. Continuous variables were reported as median (interquartile range, IQR) and discrete variables were reported as absolute (n) and relative frequency (%). The duration of follow-up was calculated from the end of treatment to the date of clinical outcome or last visit at one of the two centers until November 30, 2019. If several LRC occurred during the follow-up, the first one was considered for the analysis. The incidence rates of the primary outcomes (per 1,000 person-years) were calculated. The relative risk of incidence of outcomes was evaluated according to surrogate markers of liver dysfunction/fibrosis before HCV treatment and LSM regression post-SVR. A receiver operator characteristic (ROC) curve analysis was performed and the optimal threshold of LSM regression (%) to predict LRC or death was identified using the point nearest to the upper left corner of the ROC curve. Kaplan–Meier curves were plotted and the log-rank test was calculated for univariate analysis. We used the time to event Cox proportional-hazard model to identify factors associated with LRC or death (hazard-ratio, HR) after checking that the main variables verified the proportional-hazard assumption using the Schoenfeld residuals. Variables found to be associated (p value ≤ 0.10) with LRC or death in the univariate analysis were entered into the multivariate Cox models adjusted for age and gender. The variable Baveno VI status (favorable vs. unfavorable) replaced the variables LSM and platelet count at baseline in the models due to collinearity. The analysis was performed using STATA package, version 15, 2017 (StataCorp LP, College Station, TX, USA). Significance level was determined when p ≤ 0.05 assuming two-tailed tests.

Ethics approval and consent to participate. The study was conducted according to the good clinical practice guidelines and the Declaration of Helsinki and was approved with waiver of informed consent by the Ethical Committees from the Federal Hospital of Bonsucesso (IRB number 51736815.3.3001.5253) and the National Institute of Infectious Diseases Evandro Chagas (IRB number 51736815.3.0000.5262).
1131 patients with chronic hepatitis C treated by direct-acting agents (DAAs) in two tertiary centers from October-2015 to January-2020

144 patients excluded
n=26 treatment failure (non-SVR)
n=5 death before SVR evaluation
n=87 lost of follow-up after SVR
n=26 missing data of SVR

987 patients with sustained virological response (SVR) by DAAs prospective followed in two tertiary centers

120 patients excluded due to presence of liver-related complications before HCV treatment by DAAs

867 patients with SVR by DAAs without previous hepatic decompensation

110 patients excluded due to absence of liver stiffness measurement (LSM) by transient elastography before treatment or during follow-up
301 patients excluded due to LSM < 10 kPa before HCV treatment

456 patients with SVR by DAAs without previous hepatic decompensation who had liver stiffness measurement higher or equal to 10 kPa before HCV treatment and were prospective followed in two tertiary centers

Follow-up and incidence of clinical outcomes. Baseline LSM was performed in a median of 13.7 months (IQR 7.6–24.7) before the start of HCV treatment by DAA and LSM post-SVR in a median of 4.8 months (IQR 3.4–9.8) after the conclusion of treatment. During a median follow-up of 2.3 years (IQR 1.6–2.7) after SVR, 28 patients developed a total of 37 primary outcomes (LRC or death); HCC [n = 10 (27%), hepatic encephalopathy [n = 9 (24%)], ascites [n = 9 (24%)], variceal bleeding [n = 5 (14%)] or death [n = 4 (11%)]. No patient underwent liver transplantation. The overall rate of incidence of clinical outcomes was 29.0 (95% CI 20.0–42.0) per 1,000 person-years. Surrogate markers of liver dysfunction (albumin levels and platelet count) and/or liver fibrosis (LSM) before HCV treatment and LSM regression post-SVR were evaluated to stratify the risk of primary outcomes (LRC or death). Considering the baseline assessment before HCV treatment, the relative risk [RR (95%CI)] of incidence of outcomes was significantly higher in patients with serum albumin level less than 30 g/L (p = 0.002). The 17% decrease from before HCV treatment to after SVR was the optimal threshold of LSM regression to predict the incidence of LRC or death defined as the point nearest to the upper left corner of the ROC curve. This threshold yielded an AUROC of 0.60 (0.51–0.69) as well as sensitivity, specificity, positive and negative predictive values of 32% (16–52), 88% (85–91), 15% (7–27) and 95% (93–97), respectively.

Abbreviations: DAAs, direct-acting agents; HCV, hepatitis C virus; LSM, liver stiffness measurement; SVR, sustained virological response

Figure 1. Study flowchart for inclusion of patients.
els < 3.5 mg/dL [vs. albumin ≥ 3.5 mg/dL; RR = 4.51 (2.06–9.89), p < 0.001] and those individuals with unfavorable Baveno VI status [vs. favorable; RR = 6.58 (1.99–21.78), p < 0.001]. On the other hand, the risk of incidence of outcomes was significantly lower in those patients who had at least 20% regression of LSM post-SVR [19.3 per 1,000 person-years (95%CI 10.7–34.9)] compared with those without [43.0 per 1,000 person-years (95% CI 26.7–69.1)] [RR = 0.45 (0.21–0.96), p = 0.034] (Table 2). Patients with serum albumin level lower than 3.5 mg/dL before treatment had a higher cumulative incidence of clinical outcomes (95%CI) at 2 years compared with those with albumin ≥ 3.5 mg/dL [17.8% (10.5–29.3) vs 3.6% (2.0–6.7), log-rank p < 0.001] (Fig. 2A). Additionally, patients with unfavorable Baveno VI status at baseline had significantly higher incidence of LRC or death compared with those with favorable Baveno VI status [9.6% (6.3–14.4) vs 1.1% (0.3–4.2), log-rank p < 0.001] (Fig. 2B). In contrast, those patients who obtained a regression of at least 20% of LSM post-SVR had significantly lower cumulative incidence of clinical outcomes at 2-year follow-up compared with those without LSM regression [3.4% (1.8–7.0) vs. 9.0% (5.5–14.5), p = 0.028] (Fig. 3).

Table 1. Characteristics of patients included in the study before HCV treatment (at baseline). Data expressed as *n (%) or b median [IQR]. ALT alanine aminotransferase, AST aspartate aminotransferase, INR international normalized ratio, PEG-IFN pegylated interferon, RBV ribavirin. Missing data (n): type-2 diabetes (2), hypertension (2), dyslipidemia (9), HCV genotype (1), previous HCV treatment (1), duration of HCV treatment (1), ALT (19), AST (15), total bilirubin (35), albumin (60), INR (68), fasting glucose (34), creatinine (32), platelet count (4).
The presence of male gender, higher baseline AST levels (per 10 U/L), higher LSM values, baseline serum albumin < 3.5 mg/dL, baseline platelet count < 150 × 10^9/mm³, presence of baseline unfavorable Baveno VI status and regression of LSM post-SVR were associated (p < 0.10) with the incidence of outcomes in the univariate Cox proportional-hazard analysis (Table 3). Those variables were inserted in the multivariate Cox proportional-hazard models adjusted for age and gender. The variables LSM and platelet count before HCV treatment were replaced by the variable "Baveno VI status" in the multivariate models due to strong collinearity. The following factors were independently associated with the incidence of outcomes in the univariate Cox proportional-hazard analysis (Table 3). Those variables independently associated with an increased risk [HR = 5.56 (95% CI 2.22–13.97)] and LSM regression after SVR with a decreased risk of LRC or death [HR = 0.31 (95% CI 0.12–0.80)] (Supplementary Table 1). Moreover, we repeated the analysis considering the LSM more distantly performed from the end of the HCV treatment (median time = 12.5 months [IQR 10.3–14.9]) to evaluate the impact of LSM regression on incidence of LRC or death. LSM post-SVR remained stable, since the median value (IQR) for the first and the last LSM post-SVR were 12.4 kPa (8.5–19.9) and 12.0 kPa (7.9–19.2), respectively. Male gender [vs female, HR = 2.96 (95% CI 1.29–6.82), p = 0.011], baseline serum albumin < 3.5 mg/dL [vs ≥ 3.5 mg/dL, HR = 4.33 (1.89–9.94), p = 0.001], baseline unfavorable Baveno VI [vs favorable, HR = 4.72 (1.32–16.83), p = 0.017] and baseline unfavorable Baveno VI [vs favorable, HR = 4.72 (1.32–16.83), p = 0.017] (Table 3). In this multivariate model LSM regression of at least 20% after SVR had a trend to reduce the risk of LRC or death [HR = 0.45 (95%CI 0.21–1.02), p = 0.058]. Additionally, LSM regression post-SVR of at least 20% significantly reduced the risk of LRC or death in multivariate Cox models adjusted for age-and-gender [HR = 0.43 (0.20–0.92), p = 0.029] and those adjusted for age, gender and baseline albumin level [HR = 0.44 (0.20–0.99), p = 0.049]. We observed similar results in a sensitivity analysis considering only patients with unfavorable Baveno VI status before HCV treatment (n = 250 and incidence of primary outcomes in 25 patients): low albumin level was independently associated with an increased risk [HR = 5.56 (95% CI 2.22–13.97)] and LSM regression after SVR with a decreased risk of LRC or death [HR = 0.31 (95% CI 0.12–0.80)] (Supplementary Table 1). Moreover, we repeated the analysis considering the LSM more distantly performed from the end of the HCV treatment (median time = 12.5 months [IQR 10.3–14.9]) to evaluate the impact of LSM regression on incidence of LRC or death. LSM post-SVR remained stable, since the median value (IQR) for the first and the last LSM post-SVR were 12.4 kPa (8.5–19.9) and 12.0 kPa (7.9–19.2), respectively. Male gender [vs female, HR = 2.96 (95% CI 1.29–6.82), p = 0.011], baseline serum albumin < 3.5 mg/dL [vs ≥ 3.5 mg/dL, HR = 4.33 (1.89–9.94), p = 0.001], baseline unfavorable Baveno VI [vs favorable, HR = 4.72 (1.32–16.83), p = 0.017] and baseline unfavorable Baveno VI [vs favorable, HR = 4.72 (1.32–16.83), p = 0.017] (Table 3). In this multivariate model LSM regression of at least 20% after SVR had a trend to reduce the risk of LRC or death [HR = 0.45 (95%CI 0.21–1.02), p = 0.058]. Additionally, LSM regression post-SVR of at least 20% significantly reduced the risk of LRC or death in multivariate Cox models adjusted for age-and-gender [HR = 0.43 (0.20–0.92), p = 0.029] and those adjusted for age, gender and baseline albumin level [HR = 0.44 (0.20–0.99), p = 0.049]. We observed similar results in a sensitivity analysis considering only patients with unfavorable Baveno VI status before HCV treatment (n = 250 and incidence of primary outcomes in 25 patients): low albumin level was indepen-
dently associated with an increased risk [HR = 5.56 (95% CI 2.22–13.97)] and LSM regression after SVR with a decreased risk of LRC or death [HR = 0.31 (95% CI 0.12–0.80)] (Supplementary Table 1). Moreover, we repeated the analysis considering the LSM more distantly performed from the end of the HCV treatment (median time = 12.5 months [IQR 10.3–14.9]) to evaluate the impact of LSM regression on incidence of LRC or death. LSM post-SVR remained stable, since the median value (IQR) for the first and the last LSM post-SVR were 12.4 kPa (8.5–19.9) and 12.0 kPa (7.9–19.2), respectively. Male gender [vs female, HR = 2.96 (95% CI 1.29–6.82), p = 0.011], baseline serum albumin < 3.5 mg/dL [vs ≥ 3.5 mg/dL, HR = 4.33 (1.89–9.94), p = 0.001], baseline unfavorable Baveno VI [vs favorable, HR = 4.72 (1.32–16.83), p = 0.017] and baseline unfavorable Baveno VI [vs favorable, HR = 4.72 (1.32–16.83), p = 0.017] (Table 3). In this multivariate model LSM regression of at least 20% after SVR had a trend to reduce the risk of LRC or death [HR = 0.45 (95%CI 0.21–1.02), p = 0.058]. Additionally, LSM regression post-SVR of at least 20% significantly reduced the risk of LRC or death in multivariate Cox models adjusted for age-and-gender [HR = 0.43 (0.20–0.92), p = 0.029] and those adjusted for age, gender and baseline albumin level [HR = 0.44 (0.20–0.99), p = 0.049]. We observed similar results in a sensitivity analysis considering only patients with unfavorable Baveno VI status before HCV treatment (n = 250 and incidence of primary outcomes in 25 patients): low albumin level was inde-

Table 2. Incidence rate [per 1,000 person-years (95% confidence interval)] of liver related complications or death in patients with hepatitis C after sustained virological response by direct-acting agents. "n = 396 patients and 25 clinical outcomes; n = 452 patients and 28 clinical outcomes. Baveno VI status was defined as unfavorable if LSM ≥ 20 kPa or platelet count < 150 × 10^9/mm³. LRC liver-related complications (ascites, hepatic encephalopathy, variceal bleeding or hepatocellular carcinoma)."
On the other hand, the regression of LSM ≥ 20% during follow-up after HCV eradication seems to be a significant protective factor. The threshold of 20% for LSM regression after SVR compared to before treatment was chosen as the rounded point nearest to the upper left corner of the ROC curve for LSM decrease to predict LRC or death. Large sample size studies, such as the ERCHIVES (n = 12,467) and ANRS CO22 Hepather (n = 9,895) cohorts, confirmed that interferon-free HCV treatment significantly decreases all-cause mortality. However, the evaluation of the impact of LSM regression after SVR compared to LSM before HCV treatment was not feasible in both studies. Early regression in LSM after SVR might be related to changes in hepatic inflammation rather than fibrosis. However, a large Canadian cohort of HIV-HCV coinfected study that prospectively examined long-term changes in LSM before and after SVR due to DAAs suggested that LSM following SVR likely reflects true reversal of fibrosis.

We acknowledge that the prognostic value of post-SVR LSM remains controversial. Pons et al. reported that albumin level at baseline and regression of LSM after SVR during follow-up were independently associated with the risk of HCC in 572 patients with LSM ≥ 10 kPa before treatment during a median follow-up of 2.9 years. In addition, the regression of LSM at SVR compared to 30-days before the start of HCV treatment was related to lower incidence of LRC in the 640 patients from the HEPAVIR Spanish cohort during a median follow-up of 31 months. However, this study exclusively included HIV-HCV coinfected patients with LSM ≥ 9.5 kPa and the authors did not exclude those patients with prior hepatic decompensation. Semmler et al. reported that LSM determined after HCV-eradication and its combination with von Willebrand factor antigen and platelet count) could predict post-SVR hepatic decompensation in patients with c-ACLD. On the other hand, regression of LSM post-SVR were not associated with an increased risk of hepatic decompensation. However, this

Figure 2. Cumulative incidence of liver related complications or death according to characteristics before HCV treatment (A) albumin levels (< 3.5 mg/dL vs. ≥ 3.5 mg/dL) and (B) Baveno VI status (unfavorable vs favorable) [all log-rank tests]; SVR sustained virological response.
study included patients with previous hepatic decompensation and the impact of LSM regression after SVR was evaluated using a lower threshold compared to our study (≥ 20%).

Hepatic venous pressure gradient (HVPG) remains the most robust predictor of clinical decompensation in patients with cirrhosis22. However, HVPG measurement is an invasive method not available worldwide. The role of LSM regression after SVR remains unclear. Despite a markedly decrease in LSM after SVR, this regression seems not to be correlated with HVPG changes in patients with HCV-related cirrhosis and clinically significant portal hypertension before HCV treatment23. On the other hand, Mandorfer et al. have shown that SVR by DAAs ameliorated portal hypertension and the relative change in LSM (per %; HR = 0.972; 95%CI: 0.945–0.999; p = 0.044) was a predictor of a HVPG decrease ≥ 10%24. Additionally, Thabut et al. analyzed data from 891 patients

Figure 3. Cumulative incidence of liver related complications or death according to regression of liver stiffness measurement (LSM) after sustained virological response (SVR) [all log-rank tests].

Table 3. Factors associated with incidence of liver related complications or death during follow-up after sustained virological response. A time-dependent Cox model was used for the analysis. Variables found to be associated (p value ≤ 0.10) with LRC or death in the univariate analysis were entered into the multivariate Cox models adjusted for age and gender. The variable Baveno VI status (favorable vs. unfavorable) replaced the variables LSM and platelet count at baseline in the models due to collinearity. Baveno VI status was defined as unfavorable if LSM ≥ 20 kPa or platelet count < 150 × 10⁹/ mm³. Liver-related complications were ascites, hepatic encephalopathy, variceal bleeding or hepatocellular carcinoma. ALT alanine aminotransferase, AST aspartate aminotransferase, CI confidence interval, DCV daclatasvir, HR hazard ratio, LSM liver stiffness measurement, SOF sofosbuvir, SVR sustained virological response.

| Univariate analysis | Multivariate analysis |
|---------------------|-----------------------|
|                      | HR [95% CI] | p value | HR [95% CI] | p value |
| Male gender (vs female) | 2.28 [1.08–4.79] | 0.030 | 3.00 [1.30–6.95] | 0.010 |
| Age (per 10 years) | 1.01 [0.94–1.09] | 0.841 | 1.00 [0.99–1.01] | 0.704 |
| Type-2 diabetes (yes vs no) | 0.98 [0.88–1.09] | 0.761 |
| Hypertension (yes vs no) | 0.96 [0.74–1.26] | 0.778 |
| Dyslipidemia (yes vs no) | 1.00 [0.98–1.02] | 0.779 |
| HIV infection (yes vs no) | 0.51 [0.07–3.77] | 0.511 |
| HCV genotype-1 (vs other) | 0.54 [0.20–1.41] | 0.208 |
| Experimented patients (vs naive) | 1.55 [0.73–3.29] | 0.250 |
| SOF/DCV regimen (vs others) | 2.10 [0.80–5.53] | 0.132 |
| ALT (per 10 UI/L) | 1.01 [0.97–1.07] | 0.666 |
| AST (per 10 UI/L) | 1.05 [0.99–1.11] | 0.100 | 0.97 [0.90–1.05] | 0.519 |
| Albumin level < 3.5 mg/dL (vs ≥ 3.5 mg/dL) | 4.44 [2.02–9.75] | < 0.001 | 4.49 [1.95–10.34] | < 0.001 |
| Platelet count < 150 × 10⁹/ mm³ (vs ≥ 150 × 10⁹/ mm³) | 4.71 [1.91–11.63] | 0.001 |
| LSM before HCV treatment (per kPa) | 1.04 [1.01–1.06] | 0.001 |
| Baveno VI status (unfavorable vs favorable) | 6.48 [1.95–21.5] | 0.002 | 4.72 [1.32–16.83] | 0.017 |
| Regression of LSM ≥ 20% after SVR (vs < 20%) | 0.44 [0.20–0.93] | 0.033 | 0.45 [0.21–1.02] | 0.058 |
with HCV with biopsy-proven cirrhosis from the French ANRS CirVir cohort\(^25\). The authors reported that unfavorable Baveno VI status at baseline was associated with a significantly lower 5-year overall survival compared to those with favorable Baveno VI at baseline. Moreover, progression of portal hypertension in patients who had HCV suppression at inclusion or during follow-up was more frequent in patients with unfavorable Baveno VI status compared to those with favorable Baveno VI. Our findings reinforced the relationship between the presence of portal hypertension before HCV treatment and clinical outcomes post-SVR: unfavorable Baveno VI status before HCV treatment was associated with a worst prognosis leading to an increased risk of clinical outcomes after SVR [adjusted HR = 4.72 (1.32–16.83)] in a multivariate model adjusted for confounding factors. In the current study, similar results were observed when considering the LSM performed most distantly from the end of treatment [median of 12.5 months (IQR, 10.3–14.9)] instead of the closest TE examination [median of 4.8 months (IQR, 3.4–9.8)] following HCV treatment (Supplementary Table 2).

The main limitation of this study remains the lack of paired liver biopsy to assess the relationship of histological regression of liver fibrosis and lower incidence of clinical outcomes after SVR. However, the accuracy of TE to stage liver fibrosis was extensively validated\(^26\) and international guidelines have been recommending liver fibrosis staging using LSM by TE in HCV-infected patients\(^27\). Additionally, the performance of liver biopsy in SVR patients should be reserved in cases of known or suspected mixed etiologies (e.g. metabolic syndrome, alcoholism or autoimmunity) and for patients who persist with elevated aminotransferases after SVR. We acknowledge that the retrospective study design might be considered a limitation. However, we performed an analysis of a large real-life cohort of patients with HCV treated by DAAs whose medical records for identification of outcomes were reviewed by two trained investigators (JP and CI). We are aware of the relative short follow-up (median time of 2.3 years) in the current analysis. However, in July 2015, first generations of DAAs became available in Brazil and were made available free of charge by the Public Health System (SUS, Sistema Único de Saúde), but only for patients with advanced liver fibrosis/cirrhosis. Universal access to DAAs was implemented by the Brazilian Ministry of Health in mid-2017. Nevertheless, our median follow-up time was similar to previous studies and our study identified 37 outcomes after SVR in 28 patients with c-ACLD (LSM ≥ 10 kPa) before HCV treatment.

Other potential criticisms might be the absence of sensitivity analyses for factors associated with all-cause mortality or liver neoplasm and the fact that TE examinations before and after SVR were not performed in pre-determined time-points by the same operator. The performance of stratified analyses considering all outcomes or sensitivity time-dependent analyses for single events separately were not feasible due to the relatively small number of clinical outcomes during follow-up. The limited number of events might be explained by the exclusion of patients with previous hepatic decompensation and the relatively short follow-up period. We acknowledge that the 14 months median time from baseline LSM to initiation of DAAs was longer than desirable. The HCV care pathway in Brazil remains difficult to navigate due to the need of multiple visits before accessing DAA regimens. This complex process can lead to a relatively long time from liver fibrosis assessment by LSM to start of HCV treatment. However, a period of approximately 1 year is a relative short time within the natural history of chronic hepatitis C in which significant changes in liver fibrosis might take decades to occur. Additionally, we are aware of the lack of spleen stiffness measurement before and after HCV treatment, the absence of evaluation of portal hypertension progression/regression after SVR and the odds of interobserver variability in liver fibrosis staging using LSM by TE in patients with chronic hepatitis C\(^28\).

The major strength of our study relies on the large sample size of a real-world cohort of patients with HCV treated by DAAs with reliable paired LSM by TE before HCV treatment and after SVR. To the best of our knowledge, this is the first study which evaluated the impact of LSM regression in the incidence of severe clinical outcomes after HCV eradication in Latin America. Additionally, we included patients with c-ACLD before HCV treatment but without previous hepatic decompensation and the analysis of paired TE examinations were performed considering the same probe to avoid issues related to technical differences between M and XL probes. We acknowledge that TE is not recommended by the current guidelines to monitor fibrosis changes after SVR because the cutoffs and the interval for identifying cirrhosis regression are yet to be defined\(^29\). However, this study suggested that the comparison of LSM performed after SVR [at median times of 4.8 and 12.5 months after end of treatment] with that performed before HCV treatment may provide important information regarding the risk of severe outcomes in the following years following SVR.

In conclusion, the findings of this study showed that low albumin and unfavorable Baveno VI status (LSM ≥ 20 kPa or platelet count < 150 × 10⁹/mm³) before HCV treatment were associated with an increased risk of clinical outcome during follow-up of patients with c-ACLD. Additionally, our results also suggested a significant decrease in the incidence of LRC or death associated with regression of LSM after SVR in this population. Non-invasive methods are low-cost alternative for monitoring patients with advanced fibrosis/cirrhosis. The use of simple parameters before HCV treatment and repetition of LSM post-SVR can help to identify patients with different risks of developing severe outcome after HCV eradication.

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Author contributions

J.P.: data collection, drafting, interpretation of data and critical revision of the manuscript; G.P.: study concept and design; analysis and interpretation of data and critical revision of the manuscript; L.G., J.D., L.V., C.B.: interpretation of data and critical revision of the manuscript; C.I., R.S., U.C.: data collection; E.P.N., B.G., V.G.V.: interpretation of data and critical revision of the manuscript; F.E.: study concept and design; study supervision; interpretation of data; statistical analysis; critical revision of the manuscript.

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Competing interests

The authors declare no competing interests.
