Treatment of chronic HCV infection in people co-infected with HIV in a Spanish tertiary hospital and in relation to real life

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Abstract

Background: Nowadays the consequences of hepatitis C virus (HCV) in patients co-infected with Human Virus Immunodeficiency (HIV) are significant and include accelerated progression of liver disease, high rates of end-stage liver disease and a less survival after liver decompensation. Thanks to direct-acting antiviral drugs (DAA) in non-interferon regimens, cure rates above 95% have been achieved with fewer side effects, allowing for effective short treatments. In real life, transitional elastography allows for effective and close monitoring without the need for invasive procedures such as liver biopsy.

Objectives: To analyse the characteristics of HIV/HCV co-infected patients treated with DAAs in Interferon-free regimens in relation to real life in a Spanish tertiary Hospital.

Methods: Retrospective descriptive study. Review of the clinical history of the first 100 treatments with DAAs interferon-free in HIV/HCV co-infected patients treated in our centre.

Results: A total of 100 different treatments corresponding to 98 HIV/HCV co-infected patients on follow-up were included in the analysis. Seventy-three (73%) were male. Forty-six (46.9%) had previously been treated with pegylated interferon and ribavirin (PR), of which only 11% had had a partial response and the rest were null responders. Only two patients had HBsAg positive and 35% had an AIDS defining disease. The mean stiffness at the beginning of treatment was 14.35 kPa (SD ± 12.3 kPa). At the end of treatment, the mean stiffness obtained by transitional elastography was 12.5 kPa (SD ± 12 kPa).

Conclusions: In our series, DAA interferon-free regimes are quite effective in HIV/HCV co-infected patients (ITT: 96% of sustained viral response (SVR); DOT: 98% of SVR), regardless of fibrosis degree, cirrhosis, opioid substitution therapy, gender, PR pre-treatment or CD4 count. Stratifying by treatment, we found a decrease in the mean values measured by elastography pre and post-treatment. New studies will be needed to determine the usefulness of transitional elastography as a tool to predict the evolution of liver disease in treated patients who achieved SVR.

Introduction

Hepatitis C Virus (HCV) infection is a ubiquitous disease with a variable prevalence according to different geographical areas, focusing in some countries on specific population groups, such as in the Mediterranean basin with people with a history of parenteral substance abuse [1]. Most infected (55-85%) have a chronic disease over the years, many of whom develop cirrhosis and / or hepatocellular carcinoma. It is estimated that 400,000 people will die annually in connection with a terminal liver disease and there are about 71 million infected people worldwide, many of whom are unaware of their serological status [2]. This is the reason why it involves a Public Health problem of the first magnitude. Currently, the standard of care for therapy against HCV infection includes direct-acting antiviral (DAAs) interferon-free regimens [3] thereby the cure of more than 95% of treated patients is achieved and the progression of liver fibrosis can be stopped or delayed, reducing the risk of death from cirrhosis and hepatocarcinoma. This leads to greater survival in patients without liver cirrhosis. These treatments are also easy to perform, totally oral, with a limited duration in time, they are generally very well tolerated and mainly with “pangenotypic activity”. However, accessibility to these treatments and even the diagnosis of chronic infection is complicated and limited, especially in developing countries. In this regard, the World Health Organization (WHO) has prepared the document: “Global strategy of the health sector against viral hepatitis, 2016-2021” to plan a horizon towards the elimination of viral hepatitis, including HCV [4]. In recent years, liver stiffness measured by transition elastography, a non-invasive technique, has become the election procedure to determine the degree of fibrosis in patients with chronic HCV infection [5] and sometimes to plan the duration of treatment. It remains to be determined as a tool to predict the evolution of liver disease in treated
patients and who achieved SVR [6]. The cure of the infection or SVR is defined as the negativization of viremia at week 12 after treatment. This SVR is associated with a normalization of liver function tests and an improvement or disappearance of necroinflammation and liver fibrosis in patients without cirrhosis. In patients with cirrhosis, healing is associated with a decrease, but not elimination, of the risk of clinical events related to chronic liver disease [7].

Material and methods

Retrospective descriptive study. Review of the clinical history of the first 100 treatments with DAAs Interferon-free regimens in HIV/HCV co-infected patients treated in our centre. Statistical analysis of variables such gender, age, risk factors, HCV-genotype, transitional elastography value and CD4 counts was performed using the SPSS statistical software package release 22.0 (IBM, Chicago, IL, USA).

Results

A total of 100 different treatments corresponding to 98 HIV/HCV co-infected patients on follow-up at a tertiary reference hospital in Almeria (Andalusia, Southern Spain) who received DAAs therapy were included in the analysis. Seventy-three (73%) were male. Forty-six (46.9%) had been previously treated with pegylated interferon and ribavirin (PR), of which only 11% had had a partial response and the rest were null responders. Seventy percent (70%) were HCV genotype GT-1 (1a: 27%, 1b: 10%, 1 without subtype: 33%) GT-3 14% and genotype GT-4 16%. Fourteen (14%) used opioid substitution therapy (mainly methadone). Only 2 patients had HBsAg positive and 35% had an AIDS defining disease. The mean CD4 count at the start of treatment was 550/mm^3 (typical deviation (SD) of 312.8/mm^3), after 12 weeks of SVR at 602/mm^3 (SD of 312.6/mm^3). Baseline median of 507/mm^3 (after 12 weeks: 535/mm^3).

All patients underwent pre-treatment transitional elastography concluding that 17.5% had a grade of fibrosis F0-F1 (< 6 kPa), 29.9% F2 (6.1-9.4 kPa), 29.9% F3 (9.5-14.5 kPa) and 22.7% F4 (≥ 14.6 kPa). The mean stiffness at the beginning of treatment was 14.35 kPa (SD ± 12.3 kPa). At the end of treatment, the mean stiffness obtained by transitional elastography was 12.5 kPa (SD ± 12 kPa). Forty-two (42%) were underwent treatments that included RBV.

If we stratify for the different treatments, we found that the mean value pre-treatment measured by elastography in those patients who received a regimen based on Sofosbuvir / Daclatasvir ± Ribavirin (SOF / DCC ± RBV) was 12.55 kPa (SD ± 8.04 kPa). At the end of the treatment, an average value of 8.6 kPa (DS ± 4.6 kPa) was obtained in this group, mostly HIV / HCV GT-3a coinfected patients. On the other hand, in the group that received treatment with Sofosbuvir / Simeprevir ± Ribavirin (SOF / SMV ± RBV), mainly with HIV / HCV GT-1, an average reference value of 19.64 kPa (SD ± 14.6 kPa) was obtained and after finishing the treatment, a decrease of the average value to 14.7 kPa (SD ± 13.5 kPa). The group that received treatment with paritaprevir / omibitasvir / dasabuvir ± ribavirin (3D ± RBV), the mean value at baseline measured by elastography was 14.19 kPa (SD ± 13.4 kPa) and post - treatment: 12.1 kPa (SD ± 10.6 kPa). In this group, most of the coinfected patients were in the GT-1a. When analysing the treatment with simeprevir / daclatasvir with a baseline elastography value of 2.3 kPa and post-treatment of 4.5 kPa. Analysing both patients, we observed that the first of them, treated with SOF + VEL had a non-alcoholic steatohepatitis in the last abdominal ultrasound performed after HCV treatment with normal lipid profile, so the worsening in the values measured by transitional elastography could be related to non-alcoholic steatohepatitis. As for the other patient (treated with SMV + DCV), she had relapsed to alcohol consumption, currently under treatment with antipsychotics and poor lipid control.

The most used combination was sofosbuvir/ledipasvir (SOF/LDV) ± RBV by 34%, followed by 3D ± RBV in 25% of cases. Of the series, 95.7% had successfully completed the treatment, with 2 premature discontinuations [none of them about adverse events (AE); one reached SVR]. One patient died due to no hepatic causes. Sixty-eight (68%) were undetectable at week 4 of treatment, 97.8% at week 8 of treatment and 98.9% at end of treatment. On intention to treat analysis (ITT) the rate of SVR was 96% [98% on directly observed therapy (DOT)]. Only 2 patients had a relapse (2%: a non-cirrhotic GT-1a patient treated with sofosbuvir + daclatasvir × 12w and a non-cirrhotic GT-3a patient treated with sofosbuvir + daclatasvir × 12w) and none had virological failure during treatment. No patient had any severe adverse event (SAE) and none had stopped therapy from any AE.

Conclusions

In our series, DAAs interferon-free regimes are quite effective in HIV/HCV co-infected patients (ITT: 96% of SVR; DOT: 98% of SVR), regardless of fibrosis degree, cirrhosis, opioid substitution therapy, gender, PR pre-treatment or CD4 count. Both relapsed patients were initially treated with the same AAD interferon-free regimen but achieved SVR after being re-treated. No SAE was observed, and no premature discontinuation was observed because any kind of AE. Stratifying by treatment, we found a decrease in the mean values measured by elastography pre and post-treatment. New studies will be needed to determine the usefulness of transitional elastography as a tool to predict the evolution of liver disease in co-infected HIV/HCV patients treated with DAAs who achieved SVR.

References
