TRIPLE THERAPY IN CHRONIC HEPATITIS C: initial series in a public health program in the South of Brazil

Paulo R L Almeida¹, Carla Bortolin Fonseca¹, Vivian W Koch¹, Amanda M Souza¹, Alberi A Feltrin¹ and Cristiane Valle Tovo¹,²

ABSTRACT – Background – Chronic hepatitis C has great impact on world’s health. Current therapy for genotype 1 hepatitis C virus includes protease inhibitors boceprevir and telaprevir, associated to standard therapy – peginterferon alfa + ribavirin. There are no published data in Brazil on the results of this new therapy, and it is interesting an evaluation of what was accomplished up to this moment. Objectives – To evaluate virologic response to triple therapy, as well as the safety profile and tolerability. Method – This study is a clinical series of patients receiving triple therapy for C hepatitis in a single center of a Public Health System of South Brasil. Out of the 121 patients that initiated the triple therapy, the first patients that finished the treatment and evaluated the sustained virological response (24 weeks after the end of treatment) were included. Results – Twenty four genotype 1 chronic hepatitis C monoinfected patients were included. Nineteen (79.2%) patients had been previously treated. Thirteen (54.2%) patients were cirrhotic. Nineteen (79.2%) patients completed the planned therapy. By the end of the treatment, 14 (58.3%) out of 24 patients had undetectable viral load. Sustained virologic response occurred in 12 (50.0%) out of 24 patients, 07 (58.3%) in telaprevir group and 05 (41.7%) in boceprevir group. Out of 24 patients under triple therapy, 58% (n=14) presented anemia. Conclusion – In conclusion, despite the small number of patients treated with triple therapy evaluated in the current study, it possibly reflects the population under this therapy in real-life. HEADINGS – Hepatitis C. Protease inhibitors. Chronic C hepatitis, therapy.

INTRODUCTION

More than 170 million people worldwide suffer from chronic infection with the hepatitis C virus (HCV)¹⁰. It is a leading cause of end-stage liver disease and hepatocellular carcinoma worldwide, and the most common indication for orthotopic liver transplantation in the Western world.¹²

The current therapy to genotype 1 hepatitis C involves peginterferon alfa and ribavirin (PR) associated with a protease inhibitor (PI), boceprevir (BOC) or telaprevir (TVR). The triple therapy has shown an increase in the sustained virologic response (SVR), despite the increase of adverse effects.²,⁵,⁸,⁹,¹⁴

The efficacy of triple therapy was assessed in approximately 2,290 patients with chronic hepatitis C genotype 1 who had not been treated previously [ADVANCE²⁹ and ILLUMINATE²⁰] or who had failed previous treatment with PR. In these studies, adding BOC to the treatment containing PR has significantly increased the rates of SVR compared to the standard therapy (between 60% to 70% in triple therapy versus 30% in standard). The study PROVIDE²¹, a sub-analysis including prior null responders recruited from the studies SPRINT-2 and RESPOND-2, evaluated the response rates in the retreatment with BOC and PR in this population. The observed SVR was 38%.

Triple therapy was incorporated in Brazil by the National Committee for the Incorporation of Technologies (CONITEC) in 2012 and treatments initiated in 2013. This study aimed at evaluating the virologic response and adverse effects associated to the triple increase in the SVR rates compared to the ones from the double therapy with PR (between 70% to 80% in triple therapy versus 30% in the double therapy).

Triple therapy with BOC was assessed in approximately 1,700 patients who had not been treated before or who had failed the previous treatment with PR. In these studies, adding BOC to the treatment containing PR has significantly increased the rates of SVR compared to the standard therapy (between 60% to 70% in triple therapy versus 30% in standard).

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Correspondence: Prof. Cristiane Valle Tovo. Rua Cel. Aurelio Bitencourt, 115, 201. CEP: 90430-080 - Porto Alegre, RS, Brasil. E-mail: cris.tovo@terra.com.br
therapy in the first patients that were going through this treatment in real life in a tertiary attendance Hospital and reference for liver disease in South Brazil.

METHODS

A descriptive and retrospective observational study was carried out using information from patient’s records attended in a public center for hepatitis C attendance in a tertiary Public Hospital in the South of Brazil. It is an assessment of the first patients going through the triple therapy (PR and PI). There were 121 patients monoinfected with genotype 1 hepatitis C who went through triple therapy against HCV, and the present study analyse the data of the first patients who finished the treatment and evaluated the sustained virological response, defined as undetectable HCV-RNA 24 weeks after the end of the therapy. Data were acquired retrospectively and stored in a database charts.

Information about the sustained virological response, adverse effects and early withdraw and their reasons were assessed (7).

Anemia was graded according to the World Health Organization (WHO) scale: Grade I (9.5 – 10.9 g/dl), Grade II (8.0 – 9.4 g/dL), Grade III (6.5 – 7.9 g/dL) and Grade IV (<6.5 g/dL) (13).

RESULTS

Twenty-four patients monoinfected with genotype 1 hepatitis C who went through triple therapy with PR and PI were assessed. Twelve patients received BOC and 12 TVR. Fifteen patients were men (62.5%), 18 were white (75.0%) and the average age was 55.2 years old. The average body mass index (BMI) was 27.9 kg/m².

Most of the patients (14; 58.3%) presented comorbidities (systemic arterial hypertension was the most common). Fifteen (62.5%) patients used medication besides the ones that were part of the studied therapy – 09 (37.5%) of them used one or more drugs that could interact with the IP used.

Nineteen (79.0%) patients had already gone through previous treatments: 09 (47.3%) were null responders, 06 (31.6%) were relapsing and 04 (21%) did not have records of their previous results.

Thirteen (54.2%) patients had cirrhosis, according to clininc and/or laboratorial information or hepatic biopsy; 05 (20.8%) patients had biopsy with Metavir score of fibrosis F3 and 05 (20.8%) patients, fibrosis F2. One of the patients did not have liver biopsy or clinical documentation of cirrhosis.

The average viral load (VL) before the treatment was 4,213,040 UI/mL (63,500 to 26,920,077). Most of the patients (16; 66.7%) showed high VL (>600,000 UI/mL).

In the week 4, 11/12 (91.7%) patients using TVR reached undetectable VL, while in the BOC group 8/12 patients evaluated the viral load, but only one (12.5%) had an undetectable VL.

In the week 12, most of the patients (19; 79.2%) had an undetectable VL, with similar numbers in both groups. One patient did not evaluate the VL because of treatment withdraw in the seventh week due to a detectable VL in week 4.

Sixteen (66.7%) patients completed the whole planned treatment. Three patients did not complete the treatment but they also reached an undetectable VL. Thereby, 14/24 (58.3%) patients had undetectable VL at the end of the treatment.

When it was accessed the SVR, the VL was still undetectable in 12 (50%) patients – 07 (58.3%) from the TVR group and 05 (41.7%) in BOC group.

Summing up, the results were the following: 19 (79.2%) patients completed the treatment; 12 (50%) patients presented SVR; among those patients who interrupted the therapy, 04 (16.7%) were due to therapeutic failure and 01 (4.2%) because of adverse effects (Figure 1).

FIGURE 1. Flow diagram of study patients

Regarding adverse effects (AE), 09 (37.5%) patients presented itching (05 TVR and 04 BOC); 02 (8.3%) patients using TVR presented cutaneous rash; 09 (37.5%) patients presented anorectal symptoms (07 TVR); 02 (8.3%) referred dysgeusia.
One patient in the BOC interrupted the treatment due to exacerbation of cardiac failure. Other three patients (one in BOC group and two in TVR group) also presented adverse events that led to withdraw the treatment; however, because they had already presented undetectable VL, their treatments were considered completed by the response guided therapy. Among these patients, one presented thrombocytopenia (BOC), one had anal abscess and thrombocytopenia (TVR) and another patient had asthenia and incapacitating myalgia (TVR).

Fifty-eight percent of the 24 patients who went through triple therapy presented anemia. Among these, five patients presented grade I, six patients presented grade II and three others presented grade III. The BOC group presented anemia in 75% of the cases while the TVR group presented 41%.

Neutropenia with neutrophil count at 500 to 750/mm³ was evident in 50% of the patients using BOC and 16% of the patients using TVR during therapy. There were two cases of severe neutropenia (<500/mm³) – both patients belonged to the BOC group.

Eight patients had severe thrombocytopenia (platelets rate: 20,000 to 50,000/mm³); only one patient belonged to the BOC group while the other seven were in the TVR group.

**DISCUSSION**

This study presents the experience of the patients treated at a Public Health System center, and to our knowledge, this is the first publication referring to the Brazilian experience with the triple therapy.

In the present case series, the SVR rate of the patients who had gone through triple therapy was 50% (12 out of 24 patients). The result was lower than previous results that had been published in the trials of register, such as ADVANCE(8), ILLUMINATE(9), REALIZE(14), SPRINT-2(3) and RESPOND-2(2). The fact that the response was not so exuberant may suggest that these trials do not reflect the health care reality presenting selection biases. Because of the strict criteria for inclusion these patients distance themselves from other patients with common comorbidities who will go under triple therapy in real life. A previously published study in a setting of real-life had already shown lower results with double therapy than the ones described in the original trials(10).

A French study including cirrhotic patients previously experienced and now treated with triple therapy shows SVR rates varying from 19.4% (null responders) to 74.2% (relapsers) with TVR and from zero (null responder) to 53.9% (relapsers) with BOC, which is quite similar to the results found in the present study(11).

Half of all the patients involved in the present case series (12/24) presented rapid virologic response (RVR). Most of the patients who reached RVR were in the TVR group (11/12) once the patients in the BOC group were still in the lead-in phase.

The complete early virologic response (EVR) was distributed in a more homogenous way – 10 (83.3%) out of 12 patients using TVR and 09 (75.0%) out of 12 using BOC.

Among those patients who used TVR, SVR was observed in all naïve patients. In the ADVANCE(8) study, the SVR, in an intention-to-treat analysis, reached 75%. Among the patients who were previously treated with double therapy, 40% (4/10) presented SVR after the use of TVR. The REALIZE(14) study assessed patients previously treated, and presented a SVR of 80% in previous relapsing patients, although the response was lower regarding null responders (33%).

Boceprevir was indicated to 12 patients and 10 of these were previously experienced. SVR was observed in 41.7% (5/12) of the patients – 01 naïve and 04 previously treated. In the RESPOND-2(2) study, a SVR of 66% was obtained with the use of BOC to retreatment. Two patients using BOC were naïve, and only one presented SVR. The SPRINT-2(3) study, while submitting naïve patients to BOC presented response of 66%.

These studies show that adverse events have been more constant in patients treated with PI than the ones who were treated only with PR. The main adverse events associated with the use of BOC are anemia and dysgeusia(8,11). The average frequency of serious AE was only 12% among those who received BOC and 5% in those who received only PR. Concerning TVR, it was noticeable that cutaneous eruptions and itching were more frequent. Besides, adverse effects such as anemia, neutropenia and leukopenia occurred more frequently in the groups of TVR if compared to the control groups(5,8,14).

Although so far a small number of patients were included, the effects are consonant with such trials. Fifty-eighty percent of the patients presented anemia – among the patients using BOC this adverse effect was presented in 75% of the cases, while in the group of TVR, only in 41% of the patients.

In the group of patients with cirrhosis, 66% presented anemia, while among the patients who did not have cirrhosis, only 50% presented anemia. In the studies of phase III REALIZE(14) and RESPOND-2(2), as well as in the CUPIC(4), the rates of anemia were clearly higher in cirrhotic patients.

Dermatological lesions occurred up to 50% of the patients, including rash and itching, most prevalently in the group using TVR. According to some already published data, 50% of the patients treated with TVR presented some dermatological lesion comparing to 32% of patients who used BOC(3,4).

In conclusion, although the obtained data refers to a small number of cases, it possibly reflects the reality of clinical practical to be observed during triple therapy in chronic hepatitis C patients.

**Author contributions**

Almeida PRL conceptualized this observational study. Fonseca CB, Koch VW, Souza AM, Feltrin AA, Tovo CV, also reviewed the manuscript. Almeida PRL, Mattos AA and Tovo CV wrote the manuscript. Almeida PRL, Mattos AA and Tovo CV also reviewed the manuscript critically for important intellectual content. All authors approved the final version of the manuscript.
RESUMO – Contexto – A hepatite crônica pelo vírus C tem grande impacto na saúde mundial. A terapia atual do genótipo 1 inclui os inibidores de protease (IP) boceprevir e telaprevir, associados à terapia padrão – alfapeginterferona + ribavirina (PR). No Brasil ainda não há estudos publicados sobre os resultados dessa nova terapia, sendo de interesse a avaliação do que foi realizado até o momento. Objetivos – Avaliar a resposta virológica ao tratamento triplo, bem como o perfil de segurança e tolerabilidade. Métodos – O estudo consta de série de casos dos pacientes em uso de terapia tripla para o tratamento da hepatite C em um polo de tratamento da Secretaria Estadual da Saúde do Estado do Rio Grande do Sul, Brasil. Dentre os 121 pacientes que estão em uso de terapia tripla (PR e IP) foram apresentados os dados referentes aos primeiros que finalizaram o tratamento e realizaram avaliação da resposta virológica sustentada em semana 24 pós-tratamento. Resultados – Foram incluídos 24 pacientes monoinfectados por hepatite C crônica genótipo 1. Dezenove (79%) pacientes eram previamente experimentados. Treze (54,2%) pacientes apresentavam cirrose. Dezenove (79,2%) pacientes completaram o tratamento planejado. Ao final do tratamento, 14 (58,3%) dos 24 pacientes apresentaram carga viral indetectável. Conclusão – Embora o presente estudo tenha avaliado um pequeno número de casos, possivelmente reflete a população submetida à terapia tripla na vida real, despida das restrições dos estudos de registro.