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Boceprevir and telaprevir for the treatment of chronic hepatitis C: Safety management in clinical practice

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Abbreviations:

AEs, adverse events

DAAs, direct-acting antivirals

PEG-IFN, pegylatedinterferon

RBV, ribavirin

SVR, sustained virologic response

EPOerythropoietin alfa

SCAR, severe cutaneous adverse reaction

BSA, body surface area

DRESS, drug rash with eosinophilia and systemic symptoms

SJS,Stevens-Johnson Syndrome

TEN, toxic epidermal necrolysis

AGEP, acute generalized exanthematouspustulosis

Keywords: hepatitis C, direct acting antiviral, protease inhibitor, anemia, rash, safety

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Abstract

Effective management of adverse events (AEs) is important to prevent treatment discontinuation and optimize hepatitis C virus (HCV) infection eradication rates. The addition of direct-acting antiviral agents (DAA), telaprevir (TVR) or boceprevir (BOC) to pegylated interferon (PEG-IFN) and ribavirin (RBV) represents a new era of therapy associated with an improvement in treatment response rates and an impairment of the safety profile compared to PEG-IFN/RBV. An increase in the frequency and severity of anemia was reported in clinical trials for both drugs, and skin disorders including rash and pruritus occurred more frequently with the TVR-based regimen. These AEs are generally manageable and do not lead to early discontinuation. The management of anemia has not been clearly established and the impact of RBV dose reductions and erythropoietin alpha (EPO) use on treatment efficacy and safety must be clarified. The management of rashes, which were mild and moderate in more than 90% of the cases, is well planned, does not require TVR discontinuation and can be treated using emollients and topical corticosteroids. However, approximately 5% of rashes were severe and a few cases were classified as severe cutaneous adverse reactions (SCAR) leading to treatment discontinuation.

Introduction:

Because of the consequences of treatment failure in patients with chronic hepatitis C virus (HCV) infection, optimizing treatment efficacy and safety is essential to prevent the development of morbidities and increase survival rates (1-3). Managing adverse events (AEs) during antiviral treatment plays an important role in improving adherence and reducing premature treatment discontinuation. In addition, interactions among different drugs in the regimen may affect the efficacy and/or safety of therapy and must be managed appropriately in each patient. While AEs and drug-drug interactions are generally well established for standard peginterferon (PEG-IFN)/ribavirin (RBV) therapy (4-6), the addition of direct-acting antiviral agents (DAAs) to PEG-IFN/RBV as part of a triple therapy regimen will change the factors to be taken into account in patient management. The oral DAA, telaprevir (TVR) and boceprevir (BOC), in combination with PEG-IFN/RBV have led to a significant improvement in sustained virologic response (SVR) in HCV genotype 1 patients, but have been associated with an increase in several AEs compared to PEG-IFN/RBV alone. In clinical practice, the effective management of safety and drug-drug interactions will be essential to optimize the benefits provided by these agents for patients infected with HCV genotype 1.

Managing safety and tolerance to triple combination therapy

In placebo-controlled phase II/III studies, the most common AEs that occurred more frequently with TVR than with placebo (>5% difference) included pruritus, rash, anemia and gastrointestinal disorders (anorectal symptoms, nausea and diarrhea), which were generally manageable and did not lead to premature discontinuation (7-9). In both BOC phase III studies, the main AEs were an increase in the occurrence of fatigue, anemia, nausea, diarrhea, dysgeusia (taste alteration), and neutropenia (10-12).

Anemia

Anemia is a well known RBV-related event which is exacerbated by the addition of TVR and BOC. The mechanism of anemia with both DAA was not due to hemolysis but was thought to be the result of a bone-marrow suppressive effect. In clinical trials, triple therapy with TVR or BOC was associated with an increase in the incidence (approximately 20%) and severity of anemia compared to PEG-IFN/RBV alone (Table 1). The frequency of anemia, defined as hemoglobin levels below 10 g/dL, was about 50% for triple therapy with BOC and 40% with TVR. Hemoglobin values gradually improved after the end of TVR dosing at Week 12, and were similar to those with PEG-IFN/RBV alone by week 20. The impact of anemia on the SVR rate was different for the two drugs. Anemia had no effect on efficacy outcomes in treatment-naïve patients (13) with TVR (Figure 1). In contrast, for BOC the SVR rate was more frequently achieved in patients with anemia than in those without in treatment-naïve and -experienced patients (14) (Figure 2). Premature discontinuation of antiviral treatment because of anemia remained rare (Table 1). The management of anemia in TVR and BOC clinical trials is summarized in Table 1. In BOC trials, 43% of patients received erythropoietin alpha (EPO) for the management of anemia, *de facto* using a quadruple combination therapy (10-12). This may be problematic in clinical practice, as BOC will have to be administered for

24 or 44 weeks. In phase II/III TVR trials, the use of EPO was generally prohibited but EPO was used in 1% of patients only (9). Blood transfusion was required in less than 5% of the patients for both drugs. A similar proportion of patients underwent RBV dose reductions to manage anemia, approximately 22 to 26% (Table 1) (7-12). The retrospective analysis of phase III clinical trials with BOC and TVR showed that RBV dose reduction did not seem to have a negative impact on SVR (Figure 1, 2) (13,14). In addition, the use of EPO did not seem to have a positive impact on SVR rate in BOC studies. These initial analyses must be discussed in relation to the results of large retrospective studies of patients treated with PEG-IFN/RBV. These studies have shown that a dose reduction of RBV only has a negative effect on the efficacy of outcomes when the cumulative dose is less than 60% of the initially planned dose. If the dose reduction of RBV occurs when HCV RNA is undetectable, the impact on SVR seems to be reduced (15). When EPO is used, the full dose of RBV can often be maintained and the quality of life is improved (16). In a *post hoc* analysis of a controlled study involving more than 3,000 patients, it has been shown that patients who developed anemia during a course of PEG-IFN/RBV had higher SVR rates than those who did not develop anemia (17). In this study, EPO increased the chance of eradicating HCV when it was administered in the first 8 weeks, probably when HCV RNA was still detectable. After the eighth week of treatment, EPO had no beneficial effect on the SVR rate. If these results are extrapolated to triple therapy, it may be necessary to maintain the full dose of RBV until HCV RNA becomes undetectable. Thus, EPO could be used according to the local regulations of each country. If anemia occurs when HCV RNA is undetectable, the RBV dose could be reduced by stages of 200 mg daily. The value of administering EPO or reducing the dose of RBV in patients with hemoglobin levels below 10 g/dL is under investigation in a prospective clinical trial with BOC. The results of this important study will be available in 2012. The initial dose of protease inhibitors must be maintained in all cases. Finally, a few patients with cirrhosis were included in phase III clinical trials with BOC and TVR (7,8,10,11). The first

safety report of the CUPIC cohort, related to the French early access program and including a large number of treatment experienced patients with cirrhosis treated with triple therapy showed a poor safety profile. BOC or TVR in combination with PEG-IFN/RBV was associated with high rates of serious adverse events (40% to 57%) with a median treatment period of 84 to 89 days. EPO was used in 41% to 45% of patients and blood transfusions were required in 4% to 17% of patients, suggesting that triple therapy must be administered cautiously with intensive safety monitoring, including anemia, in patients with cirrhosis (18).

Dermatological adverse events

Dermatological reactions with PEG-IFN/RBV are well established and tend to be a uniform entity of dermatitis: generalized pruritus and skin xerosis, with eczematiform lesions accentuated by erythematous papules and microvesicles that are often excoriated, predominantly located on the extremities and on truncal skin sites exposed to friction (19). These eruptions can be managed using the same approach as for chronic eczema (topical corticosteroids, gradually replaced by emollients), and there is usually no need to discontinue antiviral treatment (20). The new treatment era with DAA is accompanied by additional patient management considerations for HCV-treating physicians. In particular skin disorders are expected to be more frequent and more severe with triple combination regimens than with PEG-IFN/RBV alone.

In clinical trials, dermatological AEs have been reported at higher frequencies with TVR-based and sometimes with BOC-based therapy, compared to PEG-IFN/RBV alone (7-12). In placebo-controlled TVR Phase II/III studies, in which 2012 patients received at least one dose of TVR and 764 patients received at least one dose of placebo, 55% of TVR-treated patients developed a rash compared to 33% of patients treated with PEG-IFN/RBV alone (9). Although it was more extensive and severe, the typical rash in people who received a TVR-based regimen was virtually indistinguishable from the PEG-IFN/RBV rash visually and on

histopathology. Rashes were primarily pruritic and eczematous, although some had an additional maculopapular component, which is not consistent with a typical hypersensitivity. Histologically, the rash appeared to be a spongiform dermatitis, with predominantly lymphatic or eosinophilic perivascular infiltration. Most (>90%) rashes were mild or moderate (grade 1 and 2), involving less than 30% of the body surface area (BSA), and progression to more severe rash was infrequent (<10%) (9,21). Approximately 50% of rashes developed within the first 4 weeks of treatment, with the remaining 50% starting between 5–12 weeks and the median time to onset of rash (any grade) was 25 days (range 1–350) (21). Therefore, skin eruptions can occur at any time during TVR treatment. Following the end of TVR treatment at week 12, all patients continued to receive PEG-IFN/RBV, and the incidence of rash was similar between TVR and placebo-treated patients.

Overall, the incidence of severe or grade 3 rash (primarily eczematous, pruritic and involving more than 50% of BSA) was 4.8% *versus* 0.4% with PEG-IFN/RBV alone (9). Rash led to premature discontinuation of TVR alone in 5.8% of patients and of TVR combination therapy in 2.6% of patients compared to none of those receiving PEG-IFN/RBV. Following the end of TVR treatment or discontinuation, symptoms improved and usually resolved, although rashes may take several weeks to resolve.

A few cases of rash were classified as severe cutaneous adverse reactions (SCAR), which can be life-threatening if unrecognized or unmanaged, and require immediate discontinuation of antiviral treatment. In placebo-controlled Phase II and III trials, 11 patients (0.4%) were recorded as having drug reactions with eosinophilia and systemic symptoms (DRESS) and three patients (<0.1%) had suspected Stevens-Johnson Syndrome (SJS) (9,21). Among the 11 reported cases of DRESS, three were confirmed by a systematic retrospective assessment by expert dermatologists. One of these cases has been reported separately (22). Among the three SJS cases, one occurred 11 weeks after TVR was discontinued and was not considered to be related to TVR. Of the two cases of suspected SJS that occurred during the TVR treatment

phase, one was considered to be possible SJS by expert dermatologists, and the other probable SJS. All of these severe reactions resolved when treatment was discontinued (9,21). Finally, the mechanism of TVR-related rash remains unknown and no predictors have been identified. The second dermatological AEs that was frequently reported with TVR was pruritus. This event was generally reported when rash was present, but could also be seen without it. Pruritus may be invalidating and cause rare treatment discontinuations.

Guidance for managing rashes

The goal of physicians should be to give patients the best chance of eradicating HCV, i.e. to continue antiviral therapy when possible in accordance with treatment and rash management protocols. However, to avoid exposing patients to the risk of severe drug-induced cutaneous reactions, physicians treating HCV should be able to easily distinguish between dermatitis and SCAR. The recommendations for grading and monitoring of dermatological reactions, and for discontinuation of TVR, PEG-IFN, and RBV because of such events are summarized in Table 2 (9). Figure 3 provides a guide to estimate body surface area as an indicator of the severity of a dermatological reaction (23). A number of clinical and biological criteria should help physicians to distinguish between TVR-related dermatitis and potential SCAR. If drug rash with eosinophilia and systemic symptoms (DRESS) alert criteria are present, including an onset between 5 - 10 weeks after the first dose of TVR that rapidly progresses to exanthema with a prolonged fever ($> 38.5^{\circ}\text{C}$) that is not related to the PEG-IFN injection and facial oedema, the following confirmation criteria should be assessed: enlarged lymph nodes (at least 2 sites), eosinophilia ($\geq 700/\mu\text{L}$ or $\geq 10\%$), atypical lymphocytes, internal organ involvement (liver and kidney) (21). Patients presenting with rapidly progressing exanthema, skin pain, atypical or typical target lesions, mucosal involvement in at least 2 sites, or with blisters or epidermal detachment should be suspected of having Stevens-Johnson Syndrome (SJS) and toxic epidermal necrolysis (TEN) (21).

In the case of Grade 1 or 2 rash, patients can benefit from guidance on optimal skin care techniques that may limit symptoms and allow optimal antiviral therapy to be continued for as long as possible. Emollient cream, rather than lotions or ointments may be effective in relieving eczematous reactions. Cream should be applied for at least 15 minutes, beginning with areas around the joints, and progressing with broad strokes across the rest of the skin. This should be begun 15 minutes after showering or bathing, and should be applied daily. Rashes can be primarily treated with topical corticosteroids. Permitted systemic antihistaminic drugs may also be used for the treatment of pruritus. Regular follow up is important, and the patient should be advised to limit exposure to sun/heat. Baking soda or oatmeal baths and loose-fitting clothes can be suggested. If grade 3 rash is present TVR must be discontinued immediately and can be managed with topical corticosteroids without antiviral treatment (PEG-IFN/RBV) discontinuation. In case of SCAR, all antiviral treatment must be discontinued immediately and the patient must be hospitalized in an appropriate department.

Anorectal disorders

In placebo-controlled TVR Phase II/III trials, anorectal adverse events occurred more frequently in TVR arms than in control arms: 26.2% *versus* 5.4%, respectively (7-9). Events usually developed within the first 2 weeks of treatment. Reported events included hemorrhoids, anal pruritus, anal discomfort or rectal burning. Most of these events were mild to moderate, very few led to treatment discontinuation, and they resolved after completion of TVR dosing. The mechanism is unknown and no evident association was found with either generalized pruritus or skin rash. An anal examination should be performed to exclude lesions that could explain the symptoms, especially hemorrhoids, fissure nor fistula. Generally, an anal examination shows non specific erythema secondary to itching. Standard symptomatic care may be considered for managing anorectal disorders, including short-term use of non-specific

topical ± including local anesthetic in case of rectal burning. Topical corticosteroids and allowed systemic antihistaminic drugs may also be used for the treatment of pruritus.

Managing drug-drug interactions

Drug-drug interactions that lower antiviral or concomitant medication drug levels to below therapeutic ranges can result in a loss of efficacy, with suboptimal drug pressure potentially leading to drug resistance (24). In contrast, drug-drug interactions that elevate drug levels and exposure can increase the risk of AEs (24). Both of these effects may reduce the chances of treatment success, but with effective management they can be lessened.

The mechanisms of drug-drug interactions include absorption, gastrointestinal metabolism or transport, and hepatic metabolism or transport. For example, liver enzymes such as cytochrome P450 (CYP) 3A and transporters such as P-glycoprotein (P-gp) may affect plasma drug concentrations. Ultimately, whether an interaction takes place is dependent upon characteristics of both the drug and the patient.

Table 3 summarizes the key pharmacological characteristics of TVR and BOC that need to be accounted for when considering the risk of drug-drug interactions (9,12).

As a result of these characteristics, TVR and BOC are contraindicated with a number of drugs, in particular those that are highly dependent on CYP 3A (TVR) or CYP 3A4/5 (BOC) for clearance and for which elevated plasma concentrations are associated with serious or life-threatening events. Concomitant administration of TVR with active substances that strongly induce CYP 3A, and thus may lead to lower exposure and loss of efficacy of TVR, is also contraindicated (Table 4) (9,12)

In summary, it is important to review all medications prior to initiation of triple combination therapy. Once this information has been collected, a key source of information and recommendations regarding co-administration with different compounds is the drug product

label (9,12). As with HIV, online tools are now also becoming available to help healthcare professionals predict, avoid and manage drug interactions in HCV.

Conclusion

The addition of DAA (TVR or BOC) to PEG-IFN/RBV therapy will change the spectrum of elements to be taken into consideration for patient management compared to PEG-IFN/RBV alone (27). The main adverse events reported in clinical trials evaluating the efficacy and safety of TVR or BOC in combination with PEG-IFN/RBV were generally manageable and did not lead to premature discontinuation, although an increase in the frequency and severity of anemia and skin disorders relative to PEG-IFN/RBV alone were noted. In clinical practice, monitoring and effective management of AEs and drug-drug interactions will be essential to optimize treatment with TVR and BOC, and improve cure rates across different patient populations.

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Figureslegends

Figure 1:Impact of anemia and RBV reduction dose on SVR rate in treatment-naïve patients receiving triple therapy with TVR or PEG-IFN/RBV alone (13).

Figure 2: Impact of anemia and management of anemia RBV on SVR rate in treatment-naïve and -experienced patients receiving triple therapy with BOC (14).

Figure 3: Estimating body surface area (23).

Table 1

Incidence and management of anemia with TVR and BOC in triple combination compared to PEG-IFN/RBV alone in controlled clinical trials (7-12).

	TVR phase II/III placebo-controlled trials (7-9)	BOC phase III clinical trials (10-12)
Incidence of anemia	32% (TVR) vs 15% (control)	49% (BOC) vs 29% (control)
RBV dose reductions due to anemia	22% (TVR) vs 9% (control)	26% (BOC) vs 13% (control)
EPO use	Not allowed (1% use in TVR arms)	43% (BOC) vs 24% (control)
Blood transfusions	4.6% (TVR) vs 1.6% (control)	3% (BOC) vs <1% (control)
Premature discontinuation	TVR/placebo alone: 2% vs 0.5% All treatment at the same time: 1% (TVR) vs 0.5% (control)	0-3% (BOC) vs 0-1% (control)

Table 2

Grading and recommendations for managing dermatological reactions with TVR-based triple combination therapy (9,21).

Extent and features of dermatological reactions	Recommendations for monitoring of dermatological reactions and discontinuation of TVR, PEG-IFN, and RBV
Mild rash (grade 1) Localized skin eruption and/or a skin eruption with a limited distribution (up to several isolated sites on the body)	-Monitor for progression or systemic symptoms until the rash is resolved.
Moderate rash (grade 2) Diffuse rash <50% of BSA	<ul style="list-style-type: none"> -Monitor for progression or systemic symptoms until the rash is resolved. -Consider consultation with a dermatologist. -For moderate rash that progresses, permanent discontinuation of TVR should be considered. If the rash does not improve within 7 days following TVR discontinuation, RBV should be interrupted. Interruption of RBV may be required sooner if the rash worsens despite discontinuation of TVR. PEG-IFN may be continued unless interruption is medically indicated. -For moderate rash that progresses to severe, permanently discontinue TVR (see below).
Severe rash (grade 3) Extent of rash > 50% of BSA or associated with significant systemic symptoms, mucous membrane ulceration, target lesions, epidermal detachment.	<ul style="list-style-type: none"> -Permanently discontinue TVR immediately. -Consultation with a dermatologist. -Monitor for progression or systemic symptoms until the rash is resolved. -PEG-IFN and RBV may be continued. If improvement is not observed within 7 days of TVR discontinuation, sequential or simultaneous interruption or discontinuation of RBV and/or PEG-IFN should be considered. If medically indicated, earlier interruption of discontinuation of PEG-IFN and RBV might be needed.
SCAR: Generalized bullous eruption, DRESS, SJS/TEN, AGEP, erythema multiform	<ul style="list-style-type: none"> -Permanent and immediate discontinuation of TVR, PEG-IFN, and RBV. -Consider consultation with dermatologist.

BSA: Body surface area; DRESS: drug rash with eosinophilia and systemic symptoms; SJS: Stevens-Johnson Syndrome; TEN: toxic epidermal necrolysis; AGEP: acute generalized exanthematous pustulosis

Table 3

Summary of key pharmacological characteristics of TVR and BOC (9,12, 25,26).

Drug	Dosing regimen	CYP	P-glycoprotein	Non-CYP metabolism
TVR	q8h No significant boosting by ritonavir	CYP3A4: ▪ Substrate ▪ Inhibitor	▪ Substrate ▪ Inhibitor	—
BOC	tid No significant boosting by ritonavir	CYP 3A4/5: ▪ Substrate ▪ Inhibitor	▪ Substrate	▪ Substrate (aldo-ketoreductase 1C2/1C3)

q8h: every 8 hours; tid: three times daily

Table 4

Contraindications to TVR and BOC (9,12).

Class	Agent	TVR (9)	BOC (12)
Alpha-1receptor antagonists	Alfuzosin	Contraindicated	No recommendation
Antiarrhythmics	Amiodarone, bepridil, quinidine	Contraindicated (contraindicated with class Ia/III, except IVlidocaine)	Bepridil contraindicated. Caution with amiodarone/quinidine
Anticonvulsants	Carbamazepine, Phenobarbital, phenytoin	Contraindicated	No data available not recommended
Antihistamines	Astemizole, terfenadine	Contraindicated	No recommendation
Antimalarials	Lumefantrine, halofantrine	No recommendation	Contraindicated
Antimycobacterials	Rifampicin	Contraindicated	No data available not recommended
Antipsychotics	Pimozide	Contraindicated	No data available not recommended
Benzodiazepines	Oral midazolam, oral triazolam	contraindicated	contraindicated
Digestive motility stimulants	Cisapride	Contraindicated	No recommendation
Ergot rye derivatives	Dihydroergotamine, ergonovine, ergotamine, methylergonovine	Contraindicated	Contraindicated
Herbal products	St John's wort (Hypericumperforatum)	Contraindicated	No recommendation
HMG-CoA reductase inhibitors	Atorvastatin, simvastatin, lovastatin	Contraindicated	No data available. Therapeutic monitoring recommended (atorvastatin, simvastatin)
PDES inhibitors	Sildenafil, tadalafil	Contraindicated	No recommendation
Tyrosine Kinase inhibitors	Not specified	No recommendation	Contraindicated