Safety and on-treatment efficacy of telaprevir: the early access programme for patients with advanced hepatitis C

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ABSTRACT

Background and aim  Severe adverse events (AEs) compromise the outcome of direct antiviral agent-based treatment in patients with advanced liver fibrosis due to HCV infection. HEP3002 is an ongoing multinational programme to evaluate safety and efficacy of telaprevir (TVR) plus pegylated-interferon-α (PEG-IFNα) and ribavirin (RBV) in patients with advanced liver fibrosis caused by HCV genotype 1 (HCV-1).

Methods  1782 patients with HCV-1 and bridging fibrosis or compensated cirrhosis were prospectively recruited from 16 countries worldwide, and treated with 12 weeks of TVR plus PEG-IFN/RBV, followed by 12 or 36 weeks of PEG-IFN and RBV (PR) alone dependent on virological response to treatment and previous response type.

Results  1587 patients completed 12 weeks of triple therapy and 4 weeks of PR tail (53% cirrhosis, 22% HCV-1a). By week 12, HCV RNA was undetectable in the vast majority of patients (85%) of naïve, 88% of relapsers, 80% of partial responders and 72% of null responders. Overall, 931 patients (59%) developed grade 1 anaemia (grade 3/4 in 31%), 630 (40%) dose reduced RBV, 332 (21%) received erythropoietin and 157 (10%) were transfused. Age and female gender were the strongest predictors of anaemia. 64 patients (4%) of naïve, 31% of relapsers, 7% of partial responders and 4% of null responders. Overall, 931 patients (59%) developed grade 1 anaemia (grade 3/4 in 31%), 630 (40%) dose reduced RBV, 332 (21%) received erythropoietin and 157 (10%) were transfused. Anemia was the main side effect as 59% of patients receiving RBV had anaemia. Discontinuation of TVR due to AEs was necessary in 193 patients (12%). Seven patients died (0.4%, six with cirrhosis).

Conclusions  In compensated patients with advanced liver fibrosis due to HCV-1, triple therapy with TVR led to satisfactory rates of safety, tolerability and on-treatment virological response with adequate managements of AEs.

INTRODUCTION

Chronic infection with HCV is a leading cause of liver-related morbidity and mortality worldwide, for which an effective antiviral treatment with pegylated-interferon (PEG-IFNα) and ribavirin (PR) has been available since early 2000.1 2 The treatment outcome for difficult-to-cure patients, such as those chronically infected with HCV genotype 1 (HCV-1), has remarkably improved following the addition of the oral HCV protease inhibitors (PI) boceprevir (BOC) or telaprevir (TVR) to PR therapy.3 4 While eradication of HCV by PI+PR therapy has been available since early 2000.12 The treatment outcome for patients with chronic hepatitis C due to genotype 1 of HCV has remarkably improved following the addition of the oral HCV protease inhibitors boceprevir (BOC) or telaprevir (TVR) to pegylated interferon+ribavirin (RBV) (PR) therapy.5

What is already known on this subject?

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▸ However, advanced hepatic fibrosis may limit both the access as well as the response rates of patients to triple therapy with significant issues of tolerability and safety.

▸ A French observational study with TVR- or BOC-based triple therapy in patients with cirrhosis reported high frequency of anaemia and severe adverse events (AEs) particularly in patients with low platelet counts and low serum values of albumin (beyond the selection criteria of registration trials).

What are the new findings?

▸ We gained further insights into the safety and efficacy of TVR+PR treatment in 1587 patients with advanced liver fibrosis or cirrhosis due to HCV-1 who were enrolled in an open label expanded access programme to TVR-based regimen and were selected following registration trials criteria for platelets, neutrophils and albumin levels.

▸ During 16 weeks of treatment (12 weeks of triple therapy followed by 4 weeks of PR) anaemia was the main side effect as 59% of patients developed grade 1–4 anaemia (grade 3/4 in 31%), 630 (40%) dose reduced RBV, 332 (21%) received erythropoietin and 157 (10%) were transfused.

▸ Treatment was safe as 12% of patients had to discontinue TVR due to AEs and only seven died (0.4%, six with cirrhosis).

▸ At the end of triple therapy serum HCV RNA was undetectable in the vast majority of previous untreated patients and importantly in 72% of 436 patients who had a prior null response to PR. The latter is the most difficult to cure population that was under-represented in a previous field practice study in France.

Significance of this study

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How might it impact on clinical practice in the foreseeable future?

- In compensated patients with advanced liver fibrosis due to HCV-1 who fulfilled the selection criteria of registration trials, 16 weeks of TVR triple therapy proved to be safe, tolerated and effective.

related mortality,6–8 factors that may limit both the access as well as the response rates of patients to triple therapy have been identified. Perhaps the most relevant is the presence of advanced hepatic fibrosis,9–11 which affects tolerability and safety of triple therapy. The impact of this factor on treatment access and response rates is driven by an increased rate of myelosuppression-related adverse events (AEs) such as anaemia and infections in patients treated with triple therapy.10–13 In addition, suboptimal dosing due to poor treatment tolerability may impact the outcome of PI+PR therapy in patients with advanced fibrosis, leading to reduced rates of virus eradication and also to unaffordable cost to utility ratios.14 A French observational study with TVR- and BOC-based triple therapy in patients with cirrhosis revealed that 33%–46% of the patients had been ineligible for registration studies with one of these PIs due to their severe level of liver disease. Up to 11.7% of patients had to discontinue therapy until week 16 due to the onset of AEs including anaemia, neutropenia, rash, clinical decompensation and bacterial infections. In addition, more than half of the patients on TVR required treatment with bone marrow stimulating factors and 16.1% received blood transfusions to manage treatment-related anaemia.15 Another real-world study in Germany showed a high frequency of anaemia and SAEs in patients with cirrhosis who were treated with triple therapy, particularly in patients with low platelet counts.16 Therefore, the safety of triple therapy is now being perceived by hepatologists as a potential barrier to its use in patients with the most need, such as those with advanced fibrosis, who are more often vulnerable to myelosuppression-related complications.

To gain further insights into the safety and efficacy of TVR +PR treatment in patients with advanced liver fibrosis or cirrhosis due to HCV-1 infection, an open label expanded access programme (EAP) was launched in late 2011 involving 16 nations worldwide. Cumulatively, the programme enrolled more than 2000 treatment naive or experienced patients with bridging fibrosis or cirrhosis. We report here the analysis of the on-treatment virological response and safety in the first 1587 patients who reached week 16 of therapy.

PATIENTS AND METHODS

Patients

From August 2011 to March 2013, compensated (Child–Pugh A) patients with bridging fibrosis or cirrhosis due to HCV-1 infection were enrolled in the TVR EAP at study centres located in 16 countries across Europe, South America and Australasia. Key inclusion criteria were male and female gender; ages 18–70; infection with HCV-1; quantifiable serum HCV RNA; and documentation of liver fibrosis assessed by liver biopsy or a non-invasive test like Fibrotest or Fibroscan showing severe fibrosis (Metavir F3 or Ishak 3–4 (S3,4)) or cirrhosis (Metavir F4 or Ishak 5–6 (S5,6)). Eligible patients had to have an absolute neutrophil count >1500/mm3, a platelet count >90 000/mm3 and haemoglobin (Hb) >12 g/dL for women and 13 g/dL for men. Patients were excluded if they had history or other evidence of decompensated liver disease, hepatocellular carcinoma, or if they were infected or co-infected with HCV other than genotype 1, HBV, or HIV or had a history of alcohol abuse. Additionally, patients were excluded if they had a history of receiving HCV protease or polymerase inhibitors. The enrolment criteria for the TVR EAP were similar to the REALIZE study.12

The protocol was signed by the principal site investigators and approved by the independent ethics committee at each participating study centre. All patients provided written informed consent prior to the conduct of any procedures for the EAP.

Treatment

During the first 12 weeks of the programme, all patients received oral administration of TVR at a dose of 750 mg every 8 h in combination with PR. The type (α2a vs α2b PEG-IFN; copegus vs rebetol) and doses of PR were selected according to local guidelines. After week 12, PR was administered for another 12 or 36 weeks depending on the virological response to treatment and/or previous response type. The concomitant administration of PR followed label recommendations.17 In patients with bridging fibrosis who were treatment naive or prior treatment relapers, PR was administered for another 12 or 36 weeks based on virological response to treatment as measured by week 4 and 12 plasma HCV RNA levels. Patients with undetectable HCV RNA at weeks 4 and 12 received an additional 12 weeks of PR alone (total treatment duration of 24 weeks); patients with detectable HCV RNA (but not meeting stopping rules) at either week 4 or 12 received an additional 36 weeks of PR alone (total treatment duration of 48 weeks). Patients with bridging fibrosis with prior partial or null response to previous treatment with PR including a viral breakthrough and all patients with cirrhosis were treated for a subsequent 36 weeks. At weeks 4 and 12, patients were assessed as to whether they met the predefined stopping rules based on virological response. If at either time point the HCV RNA levels were greater than 1000 IU/mL, all treatment was permanently discontinued.

Measures of disease severity and treatment efficacy

Liver fibrosis was staged on histological specimens obtained by percutaneous liver biopsy using either the Metavir or Ishak score or by Fibrotest or Fibroscan. Fibroscan cut-off to diagnose bridging fibrosis and cirrhosis was ≥9.5 kPa and ≥12.5 kPa, respectively.18 The Child–Pugh score system was used to define the clinical status of patients, Child–Pugh A stage indicating compensated liver disease.

A range of assays were used to measure HCV RNA levels at local investigational sites. The majority of sites used the Roche COBAS TaqMan or Abbott RealTime assays (96%). Roche COBAS TaqMan (versions 1 and 2) has a lower limit of quantification (LLOQ) of 15–25 IU/mL depending on serum volume and the method of RNA extraction with a lower limit of detection (LLOD) of 10 IU/mL. Abbott RealTime has an LLOQ of 12 IU/mL and an LLOD of 10–12 IU/mL.

In the intent to treat analysis, the number of patients who had HCV RNA levels below LLOD was calculated for the overall population (n=1587) and in subgroups by prior treatment history.
Safety assessments
Throughout the treatment period and follow-up, safety assessments were carried out including laboratory assessments, physical examinations, evaluation of vital signs and the reporting of AEs. Any clinically significant abnormalities persisting at the end of the EAP/early withdrawal were followed by the investigator until a resolution or clinical stable endpoint was reached.

The DAIDS criteria were used to grade AEs except for rash which had protocol-specific definitions of severity grades: grade 1 mild, localised to one or several isolated sites; grade 2 moderate, diffuse skin eruption involving up to 50% of the body surface area; grade 3 severe, involving more than 50% of the body surface area or with significant systemic signs or symptoms; and grade 4 life-threatening, diagnosis of generalised bullous eruption, Steven Johnson syndrome or TEN. For patients who experienced grade 1 or 2 rash, medical management was left to the discretion of the investigator. For patients who had grade 2 rash progressed to grade 3, and for those experiencing grade 3 rash, TVR was permanently discontinued. If the rash did not improve, symptomatically or objectively within 7 days following TVR discontinuation, R was also discontinued. Immediate and permanent discontinuation of all study drugs was mandatory for all patients diagnosed with grade 4 rash.

Hb levels were assessed before treatment and at weeks 2, 4, 8 and 12, and as clinically appropriate thereafter; additional visits could be performed at the discretion of the investigator. Anaemia was defined as an AE by the investigator in the case report form, according to the following guidelines: grade 1 Hb values between 10.0 and 10.9 g/dL, or any decrease from baseline between 2.5 and 3.4 g/dL, grade 2 Hb values between 9.0 and 9.9 g/dL, or any decrease of Hb between 3.5 and 4.4 g/dL; grade 3 Hb values between 7.0 and 8.9 g/dL or any decrease of Hb ≥ 4.5 g/dL; and grade 4 an Hb value less than 7.0 g/dL. If anaemia developed during treatment, R dose was modified according to label recommendations. TVR was discontinued only if reductions of R dose or discontinuation did not result in an improvement of anaemia. TVR dose reductions were prohibited and TVR could not be reinitiated if treatment was discontinued. Use of blood transfusions, erythropoietin (EPO) or iron-based products were allowed during the trial.

Statistical methods
Fisher’s exact tests were used to detect differences in the prevalence of AEs between patients who were F3 versus F4 at baseline. Multivariate linear regression was used to identify baseline factors associated with a higher probability of extended rapid virological response (eRVR) (defined as HCV RNA undetectable at both weeks 4 and 12) and development of anaemia (defined as Hb below 10 g/dL at any time on treatment).

RESULTS
Baseline characteristics
Patient demographics and disease severity are shown in table 1A,B. A total of 746 patients (47%) had bridging fibrosis and 835 patients (53%) a cirrhosis (F4 or S5.6).

Disease stage was assessed by Fibroscan in 1149 patients (72%), biopsy in 308 patients (19%) and fibrosis markers in 130 patients (8%). Fibrosis stage was classified by Metavir in 1412 patients (89%) and Ishak in 175 patients (11%). The mean and SD of the baseline score of Fibroscan was 11.6 kPa (2.8) for patients classified as F3 at baseline, and 25.0 kPa (12.6) for patients classified as F4 at baseline.

Overall, the mean age of the patients was 53; 1012 patients (64%) were male and 1557 (98%) were white. In all, 357 patients (22%) were infected with HCV-1a and 1055 (66%) had HCV RNA greater than or equal to 800 000 IU/mL. At baseline, 374 patients (24%) had grade 1–3 thrombocytopenia (platelets <125 000/mm3), while 134 patients (8%) had a grade 2–4 thrombocytopenia (platelets <100 000/mm3). A total of 23 patients (1%) had grade 1–3 reductions in serum albumin (<3.5 g/dL).

Among the 532 out of 835 cirrhotic patients with currently available information, 78 patients (14.7%) had either grade or presence of oesophageal varices reported.

Overall, 321 patients (20%) were treatment naive, 531 (33%) were prior treatment relapsers, 203 (13%) were prior partial responders, 436 (27%) were prior null responders and 49 (3%) had had a viral breakthrough. In all, 47 patients (3%) had had an unspecified previous non-response. The demographic and clinical characteristics were similar between the bridging fibrosis and cirrhosis study groups.

Efficacy
Overall, 82% of patients at week 4 had a serum HCV RNA level less than 25 IU/mL, a rate that increased to 86% at week 12. Furthermore, 60% and 82% of patients had undetectable HCV RNA at weeks 4 and 12, respectively. The percentage of treatment naive patients who had undetectable HCV RNA levels at week 12 was 85% whereas among treatment experienced patients, 88% of treatment relapsers, 80% of partial responders, 72% of null responders and 84% of viral breakthrough patients had undetectable HCV RNA levels at week 12.

Figure 1 shows the outcome of treatment at weeks 4 and 12 for each subgroup of patients with respect to previous PR treatment. At week 12, null responders had a significantly higher rate of virological failure (14%) than the other groups (treatment naive (4%), prior relapsers (2%) and partial responders (5%)). In all, 37 patients (2%) stopped TVR at week 4 having met the futility rules criteria. Three patients (0.2%) continued triple therapy despite having met the week 4 virological criteria for anticipated TVR interruption. The number of patients discontinuing triple therapy for AEs was similar in all groups (7% for treatment naive, 4% for prior relapsers, 7% for partial responders and 7% for null responders).

In multivariate analysis, four baseline factors were associated with a higher chance of eRVR: baseline viral load <800 000 IU/mL (OR=1.47, 95% CI 1.18 to 1.85), genotype 1b (OR=1.52, 95% CI 1.16 to 1.96), α-fetoprotein <10 pg/mL (OR=2.36, 95% CI 1.82 to 3.23) and naive, relaper or prior partial response versus prior null response (OR=2.0, 95% CI 1.56 to 2.5). The rates of eRVR were 20/50 (40%), 77/235 (33%), 262/500 (52%), 349/584 (60%) and 173/200 (79%) for patients with 0, 1, 2, 3 or 4 of these predictive factors, respectively. There was no association between eRVR and the type of PEG-IFN used.

Safety and tolerability
Through week 16, 1014 (64%) patients experienced grade 2–4 AEs that were considered related to TVR treatment. The most common AEs were anaemia (n=698, 44%), rash (n=201, 13%), thrombocytopenia (n=120, 8%), pruritus (n=95, 6%) and asthenia (n=91, 6%). Patients with cirrhosis developed more AEs than those with bridging fibrosis (67% vs 60%, p=0.01). Overall, 12% of patients experienced AEs that ultimately led to TVR discontinuation (table 2). Of the 1587 patients, seven patients died during the PR tail as a consequence of

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Table 1  Baseline characteristics of the study patients according to disease severity

| Characteristic                      | Bridging fibrosis (N=752) | Cirrhosis (N=835) | Overall (N=1587) |
|-------------------------------------|---------------------------|-------------------|------------------|
| **Panel A**                         |                           |                   |                  |
| Age year—mean (range)               | 52 (22–73)                | 54 (19–75)        | 53 (19–75)       |
| Body mass index (BMI)†             | 26±3.7                    | 27±4.2            | 27±4.0           |
| BMI—range                          | 18–42                     | 19–47             | 18–47            |
| Males sex—no. (%)                  | 463 (62)                  | 549 (66)          | 1012 (64)        |
| Race or ethnic group—no. (%)§      |                           |                   |                  |
| White                               | 740 (98)                  | 817 (98)          | 1557 (98)        |
| Black, Asian or other              | 12 (2)                    | 18 (2)            | 30 (2)           |
| HCV-1 subtype—no. (%)             |                           |                   |                  |
| 1a                                  | 168 (22)                  | 189 (23)          | 357 (22)         |
| 1b                                  | 562 (75)                  | 609 (73)          | 1171 (74)        |
| Missing or unknown                 | 22 (3)                    | 37 (4)            | 59 (4)           |
| HCV RNA log_{10} IU/mL, mean (SD)¶ | 6.2±0.66                  | 6.1±0.74          | 6.1±0.71         |
| HCV RNA > 800 000 IU/mL—no. (%)§   | 507 (67)                  | 548 (66)          | 1055 (66)        |
| Model for End Stage Liver Disease score | 7 (6–8)               | 7 (7–8)          | 7 (6–8)         |
| α-Fetoprotein—μg/L                 | 5.6 (3.5–10.0)            | 9.0 (5.3–16.9)    | 7.1 (4.2–13.1)   |
| Albumin—g/L                        | 44.0 (41.1–46.1)          | 42.2 (40.0–45.0)  | 43.0 (40.4–46.0) |
| Bilirubin—μmol/L                   | 11.8 (8.2–15.4)           | 13.5 (10.0–17.5)  | 12.3 (9.0–16.7)  |
| Creatinine—μmol/L                  | 69.8 (59.2–79.6)          | 69.0 (60.0–79.0)  | 69.0 (59.2–79.6) |
| Glucose—mmol/L                     | 5.2 (4.7–5.8)             | 5.4 (4.8–6.3)     | 5.3 (4.8–6.0)    |
| Haemoglobin—g/L                    | 151 (141–160)             | 149 (140–159)     | 150 (140–159)    |
| Neutrophils ×10^9/L                | 3.2 (2.5–4.0)             | 3.0 (2.3–3.8)     | 3.0 (2.4–3.9)    |
| Platelets ×10^9/L                   | 182 (149–226)             | 144 (114–184)     | 161 (126–205)    |
| Prothrombin intl. normalised ratio | 1.0 (1.0–1.1)             | 1.1 (1.0–1.1)     | 1.04 (1.00–1.11) |
| **Panel B**                         |                           |                   |                  |
| IFN-α—3 no. (%)§                   | 577 (77)                  | 620 (74)          | 1197 (75)        |
| CC                                  | 23 (3)                    | 50 (6)            | 73 (5)           |
| CT                                  | 117 (16)                  | 121 (14)          | 238 (15)         |
| TT                                  | 35 (5)                    | 44 (5)            | 79 (5)           |
| Previous type of response—no. (%)  |                           |                   |                  |
| Prior null responder                | 180 (24)                  | 256 (31)          | 436 (27)         |
| Prior partial responder             | 91 (12)                   | 112 (13)          | 203 (13)         |
| Total non-responders†               | 291 (39)                  | 394 (47)          | 685 (43)         |
| Relapsers                           | 265 (35)                  | 266 (32)          | 531 (33)         |
| Treatment naive                     | 169 (22)                  | 152 (18)          | 321 (20)         |
| Viral breakthrough                  | 26 (3)                    | 23 (3)            | 49 (3)           |
| Unknown                             | 1 (0)                     | 0 (0)             | 1 (0)            |

*Values are medians and IQR unless otherwise indicated. Percentages may not total 100 because of rounding.
†Includes three F1 patients and three F2 patients.
‡BMI is the weight in kilograms divided by the square of the height in metres.
§Race or ethnic group was self-reported. Patients of any race could also identify themselves as Hispanic.
¶Log10 values for HCV RNA are means±SE.
**Values are means±SD unless otherwise indicated. Percentages may not total 100 because of rounding.
††Includes prior null responders, prior partial responders and non-responders-unspecified.

hepatic failure, pneumonia, haemorrhage, septic shock or ischaemic colitis leading to subsequent multi-organ failure. Detailed results are shown in table 3.

By week 16, 931 patients (59%) developed any grade anaemia (table 4). Among patients with bridging fibrosis and cirrhosis, 55% and 62%, respectively, had any grade anaemia, with 25% and 29% of the patients developing grade 3 anaemia and 3% and 4% developing grade 4 anaemia, respectively. As expected, grade 2–4 anaemia judged to be related to TVR treatment occurred more frequently in patients with cirrhosis than in those with bridging fibrosis (41% vs 47%, p<0.01, table 2). For treatment of anaemia, 630 patients (40%) underwent R dose reduction, 332 (21%) received EPO and 157 (10%) received a blood transfusion. There was combined use of EPO and blood transfusion in 74 patients (5%), EPO and R dose reduction in 234 patients (15%) and combined use of transfusion and R dose reduction in 141 patients (9%).

Figure 2 shows the incidence and prevalence rate of any grade TVR-related anaemia over the 16-week period of study. In all, 68% of cases of anaemia occurred within the first 8 weeks of treatment, and 83% of patients who developed anaemia throughout treatment were still considered anaemic at week 16. Table 4 shows the different levels of anaemia recorded during the first 16 weeks of treatment, by baseline fibrosis stage. Overall, 45 patients (3%; 31 cirrhosis and 14 bridging fibrosis) discontinued TVR for anaemia, while 27 patients (2%) discontinued R for anaemia.

There were reductions in Hb below 10 g/dL in 48% of patients. In a multivariate analysis, the four strongest predictors of Hb <10 g/dL at any time on treatment were female sex...
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Figure 1  Outcome of treatment at weeks 4 and 12, by prior treatment. Shown are the week 4 and 12 outcomes of treatment by subgroup: treatment naive (n=321), prior treatment relapers (n=531), previous partial responders (n=436) and previous null (n=203). Data for patients who had previously experienced viral breakthrough (n=49) and whose prior response to treatment was unspecified (n=47) have not been represented.

Table 2 Reasons for discontinuation of telaprevir and the incidence of the most common grade 2–4 drug-related AEs and serious AEs

| Variable                      | Bridging fibrosis (F3)* (N=752) | Cirrhosis (F4) (N=835) | Overall (N=1587) | p Value |
|-------------------------------|---------------------------------|------------------------|------------------|---------|
| Grade 2–4 drug-related AE†—no. (%) |                                 |                        |                  |         |
| Patients with one or more AE  |                                 |                        |                  |         |
| Anaemia‡                       | 307 (41)                        | 391 (47)               | 698 (44)         | 0.02    |
| Rash‡                         | 90 (12)                         | 111 (13)               | 201 (13)         | NS      |
| Thrombocytopenia‡              | 37 (5)                          | 83 (10)                | 120 (8)          | <0.01   |
| Pruritus‡                     | 37 (5)                          | 58 (7)                 | 95 (6)           | NS      |
| Anaemia                        | 44 (6)                          | 47 (6)                 | 91 (6)           | NS      |
| Rash                           | 24 (3)                          | 36 (4)                 | 60 (4)           | NS      |
| Abnormal liver test†           | 26 (3)                          | 35 (4)                 | 61 (4)           | NS      |
| Serious AEs§—no. (%)           |                                 |                        |                  |         |
| Patients with one or more serious AEs |               |                        |                  |         |
| Anaemia                        | 32 (4)                          | 43 (5)                 | 75 (5)           | NS      |
| Rash                           | 12 (2)                          | 16 (2)                 | 28 (2)           | NS      |
| Infection                      | 6 (1)                           | 20 (2)                 | 26 (2)           | NS      |
| Pyrexia                        | 4 (1)                           | 8 (1)                  | 12 (1)           | NS      |
| Reason for discontinuation¶—no. (%) |                               |                        |                  |         |
| Any AE                         | 80 (11)                         | 113 (14)               | 193 (12)         | NS      |
| Rash                           | 36 (5)                          | 36 (4)                 | 72 (5)           | NS      |
| Anaemia                        | 14 (2)                          | 31 (4)                 | 45 (3)           | 0.01    |
| Nausea                         | 8 (1)                           | 9 (1)                  | 17 (1)           | NS      |
| Pyrexia                        | 6 (1)                           | 10 (1)                 | 16 (1)           | NS      |
| Pruritus                        | 3 (0)                           | 10 (1)                 | 13 (1)           | NS      |
| Abdominal pain                 | 1 (0)                           | 8 (1)                  | 9 (1)            | NS      |

*Includes three F1 patients and three F2 patients.
†Listed are grade 2–4 drug-related AEs that occurred in at least 4% of the overall population.
‡Included in this category are all related events that were described with a variety of descriptive terms.
§Listed are serious AEs that occurred in at least 0.5% of the overall population.
¶Listed are discontinuations that occurred in at least 1% of the overall population.
*Includes three F1 patients and three F2 patients.
Table 3  Adverse events with fatal outcome (n=1587)

| Patient 1 | Patient 2 | Patient 3 | Patient 4 | Patient 5 | Patient 6 | Patient 7 |
|-----------|-----------|-----------|-----------|-----------|-----------|-----------|
| Age       | 52        | 51        | 50        | 55        | 60        | 66        | 56        |
| Gender    | Male      | Female    | Female    | Male      | Male      | Male      | Male      |
| Fibrosis stage | F4        | F4        | Undetectable | F3        | F4        | F4        | F4        |
| BSL VL    | 1 200 000 | 2 387 203 | 389 340   | 3 180 000 | 1 430 000 | 977 100   | 2 502 935 |
| Last obs. VL | Undetectable | Undetectable | 20        | 570       |          |          |
| Dates of TVR treatment | July 2011 to September 2011 | December 2011 to February 2012 | January 2012 to April 2012 | September 2011 to December 2011 | November 2011 to February 2012 | May 2012 to July 2012 | April 2012 to July 2012 |
| Date of death | 4 weeks after TVR d/c | 2 weeks after TVR d/c | 4 weeks after TVR d/c | 30 weeks after TVR d/c | 4 weeks after TVR d/c | 24 weeks after TVR d/c | 8 weeks after TVR d/c |
| Adverse event | Anaemia, dehydration, hepatic failure, hepatorenal syndrome, hyper-catabolism, keto-acidosis, multi-organ failure | Ischaemic colitis, septic shock, multi-organ failure | Hepatic failure, bone marrow failure, multi-organ failure | Anaemia, hepatic neoplasm malignant, intra-abdominal haemorrhage, pneumonia | Anaemia, oesophageal variceal haemorrhage, pneumonia | Diarrhoea, vomit, hypotension, septic shock, coma |
| Causality | Possibly related | Related | Unlikely related | Unlikely/not related | Unlikely related | Not related | Unlikely related |
| Medical history | Diabetes | Low platelets (74 000) | | | | Neutropenia (830 cells/mm³) |

BSL, baseline; d/c, discontinued; TVR, telaprevir; VR, viral load.

Owing to the multifactorial origin of anaemia in patients with advanced liver fibrosis including age, myelosuppression and impaired renal function, higher rates of grade 3 or 4 anaemia were observed in EAP patients than those observed in phase II/III studies (31% in the EAP compared with 4%–18% previously reported). These discrepancies can be accounted for by the fewer number of patients enrolled in clinical development studies who had either bridging fibrosis or cirrhosis (ie, an advanced liver disease entailing a significant risk of developing myelosuppression-related AEs). Additionally, our choice of defining anaemia as either Hb below given threshold levels or any decrease of Hb following triple therapy could have increased the estimate of anaemia in some patients. In all, 40% of the overall cohort had anaemia managed with R dose reduction and blood transfusion was used in 10%. In contrast to the registration trial protocols where the use of EPO to correct anaemia was not permitted,11 12 332 patients (21%) received EPO. In addition, a significant proportion of patients had their anaemia treated through a combination of R dose reduction/interruption and either EPO or blood transfusion. It is possible that R dose reduction to manage anaemia was avoided because of concerns that treatment effectiveness may be compromised; however, it is now appreciated from both a retrospective and prospective study with TVR and BOC, respectively, that R dose reduction may not affect sustained virological response (SVR) rates. Overall, the strategies of anaemia management in this study resulted in only 3% of patients discontinuing TVR because of anaemia. This compares with 2% of patients in the REALIZE study who discontinued TVR for anaemia.12

Similar rates of anaemia were reported in patients with compensated cirrhosis who were enrolled in the field practice study CUPIC in France, where 29% had grade 3 anaemia (defined as Hb <9 g/dL);13 this was despite a higher number of CUPIC patients (33%) with severe liver impairment exceeding the enrolment criteria adopted in REALIZE than in the EAP (9%).12 The clinical burden of anaemia in CUPIC was higher than in the EAP, with EPO use in more than half of the patients (57%) or the need for blood transfusion in 15% possibly reflecting slightly more advanced liver disease in CUPIC patients.

Not unexpectedly, anaemia defined as a Hb drop below 10 g/dL following PR+TVR therapy more often occurred in >65-year-old patients, women and patients with low pretreatment Hb values or those with higher weight-based dosing of R. While age is a well-recognised risk factor for anaemia in patients with advanced liver disease, likely reflecting an increased susceptibility to treatment-related bone marrow and renal toxicity, female gender and low pretreatment Hb values

Table 4  Week 16: Prevalence and management of anaemia by fibrosis stage

| Characteristic | Bridging fibrosis (F3)* (N=752) | Cirrhosis (F4) (N=835) | Overall (N=1587) |
|---------------|--------------------------------|-----------------------|------------------|
| Grade 1–4 anaemia | 413 (55) | 518 (62) | 931 (59) |
| Grade 3 anaemia | 189 (25) | 238 (29) | 427 (27) |
| Grade 4 anaemia | 26 (3) | 35 (4) | 61 (4) |
| d/c TVR due to anaemia | 14 (2) | 31 (4) | 45 (3) |
| Initial RBV dose (mg/day)—mean | 1106 | 1120 | 1114 |
| Initial RBV dose (mg/kg/day)—mean | 14.6 | 14.3 | 14.4 |
| RBV dose reductions—no. (%) | 270 (36) | 356 (43) | 630 (40) |
| EPO use—no. (%) | 138 (19) | 194 (23) | 332 (21) |
| Blood transfusion—no. (%) | 60 (8) | 96 (12) | 157 (10) |
| RBV dose reduction+other intervention (EPO or blood transfusion)—no. (%) | 126 (17) | 182 (22) | 309 (20) |

*Includes three F1 patients and three F2 patients.
†Included in this category are all related events that were described with a variety of descriptive terms.
‡d/c, discontinued; EPO, erythropoietin; RBV, ribavirin; TVR, telaprevir.

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have long been recognised as risk factors for anaemia also in patients exposed to dual therapy with PR.24 25

One reassuring finding of the TVR EAP was the low rate of cutaneous rash (13% grade 2–4 treatment-related), and in particular of grade 3–4 rash affecting 4% of the population only. This may reflect the improved standard of care based on interventions and counselling to prevent cutaneous toxicity of TVR. This is not unprecedented since the rates of any grade rash were already reduced in phase III registration trials compared with the phase II trials where rash was a leading AE causing a shortened period of TVR administration.15 26 Similarly, rather low rates (4.8%) of grade 3/4 rash were also observed in the CUPIC study.15

In the EAP, a number of patients developed AEs other than anaemia or rash, including infections, thrombocytopenia, pruritus, weakness, nausea and anorectal discomfort. Similar to the cutaneous rash, anorectal discomfort was reported for a lower proportion of patients than in registration trials,11 12 20 21 26 again possibly reflecting improved standard of care based on the use of prophylactic and therapeutic remedies. Ultimately, only 12% of patients experienced SAEs including anaemia (5%), rash (2%), infection (2%) and pyrexia (1%). Thus, enrolment of patients fulfilling the selection criteria to the registration trials and well compensated liver status resulted in a satisfactory safety record of TVR-based regimens in patients with advanced liver fibrosis. This explains also the low mortality rates (n=7, 0.4%) in the EAP cohort up to week 16. Deaths were the consequence of multi-organ failure caused by infection in two patients, pneumonia in two patients, and septic shock, ischaemic colitis, and haemorrhage in one patient each. Notably, one of these patients suffered from diabetes, a disease known to increase the risk of infection, and six of the seven patients had cirrhosis. PR-related deaths were reported also in patients with cirrhosis due to HCV (2%) who were enrolled in a trial aiming to evaluate the safety and efficacy of elthrombopag,27 a synthetic compound able to increase the level of circulating platelets. Slightly higher rates (2.8%) of TVR treatment-related deaths were reported in patients with cirrhosis who were enrolled in the CUPIC study15 where mortality was almost invariably associated with liver failure and predicted by signs of impaired liver function including a baseline platelet count less than 100 000/mm² and serum albumin lower than 3.5 g/dL. In this study, TVR-based regimens were also associated with a significant rate of infection (6.5%). With respect to the TVR EAP, the greater burden of AEs and serious complications (29 among 429 patients) observed in the CUPIC study during the first 16 weeks of triple therapy underscore the enrolment of a significant number of patients with deteriorated liver function. In the French study, 20% of patients...
had <100,000 platelets and 12% <3.5 g albumin compared with TVR EAP where these figures were 8% and 1%, respectively. Poor safety signals in patients with more profound liver derangement were reported in two studies in Germany and Austria where the prevalence of patients with severe liver impairment was intermediate between CUPIC and TVR EAP.

The finding that 85% of previously untreated patients had undetectable HCV RNA at week 12 with minor differences between patients with bridging fibrosis and those with cirrhosis (87% vs 83%) is similar to the antiviral efficacy of TVR regimens seen in registration trials. Based on previous phase III trial results, one may therefore predict that a significant proportion of previously untreated patients with advanced fibrosis will ultimately achieve an SVR upon completion of the treatment schedule. Still, the SVR results need to be confirmed in long-term follow-up. In most countries, treatment naive patients with advanced fibrosis have been prioritised to receive triple therapy with TVR. The EAP study also provides encouraging though preliminary data of the efficacy of TVR triple therapy in treatment-experienced patients with bridging fibrosis and cirrhosis. These findings were particularly rewarding for those individuals with a previous relapse or partial response to PR, reaching very high HCV RNA undetectability rates of 88% and 80% at week 12. A preliminary report of the CUPIC study is in line with our observations of SVR in experienced patients with advanced fibrosis/cirrhosis, with 46% of 107 patients enrolled in the study achieving an SVR at week 12 post-treatment. One important limit of the CUPIC study, however, is the under-representation of null responders, who are most difficult to cure, as a consequence of their poor IFN sensitivity, leading to a high risk of virological breakthroughs and post-treatment relapse of hepatitis. By enrolling 436 patients who had had a prior null response (180 bridging fibrotic and 256 cirrhotic patients), the EAP is the largest study of prior null responders.

In conclusion, the 16-week interim analysis of the TVR EAP in 1587 patients provided encouraging insights on the safety, tolerability and preliminary efficacy of TVR triple therapy in difficult-to-cure categories of hepatitis C patients with advanced fibrosis.

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Local ethic committee.

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Data sharing

The principal investigator had full access to the results of the trial.

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