

II. 潜在性 HBV 感染症例からの HBV 再活性化

母子感染を含めた新生児から小児期の HBV 感染による無症候性キャリアでは HBV 量が多いため、免疫抑制・化学療法を行う際に注意が必要となることは当然であるが、肝炎発症後に軽快し、肝機能が正常となった後もなお高リスクである。成人の初感染の場合、B型急性肝炎回復後は HBs 抗体陽性となり、いわゆる“治癒”という状態になるが、その場合でも肝細胞内には少量

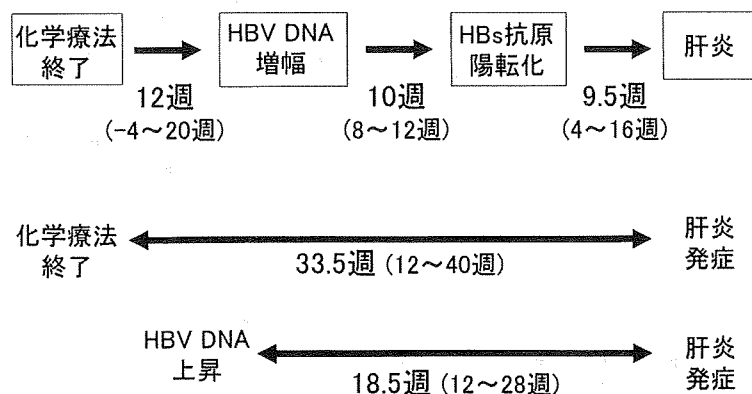
の HBV が残存していることがわかっている (図 3)。従って HBV は結核やヘルペス感染症などと同じく、一度感染すると体内から完全に消失することはないと考えられており、最近では“潜在性 HBV 感染”という表現で統一されている⁹⁾。

潜在性 HBV 感染は、血清 HBs 抗原陰性で HBe 抗体 and/or HBs 抗体陽性と定義される。この場合、特殊なウイルス遺伝子変異がない限り血中 HBV DNA は

表 1 HBV 再活性化に関係する免疫抑制・化学療法剤

| Class | Agents associated with HBV reactivation |
|-----------------------------------|--|
| Alkylators | Cyclophosphamide Ifosfamide Chlorambucil Carboplatin, Cisplatin |
| Antimetabolites | Cytarabine Fluorouracil Gemcitabine Mercaptopurine Methotrexate Thioguanine |
| Antitumor antibodies | Anthracyclines Bleomycin Mitomycin C Actinomycin D |
| Corticosteroides Immunotherapy | Prednisone/Dexamethasone etc. Rituximab (anti-CD20) Alemtuzumab (anti-CD52) Infliximab (anti-TNF) |
| Plant Alkaloids | Vincristine Vinblastine |
| Others | Asparaginase Procarbazine Docetaxel Etoposide Fludarabine Imatinib Mesylate Interferon alpha |

Lalazar G, et al : Br J Haematol 136 : 699-712, 2007 より引用



Hui CK, et al: Gastroenterology 131: 59-68, 2006より引用改変

図 2 HBV 再活性化のウイルスマーカーの動きと肝炎発症までの経過

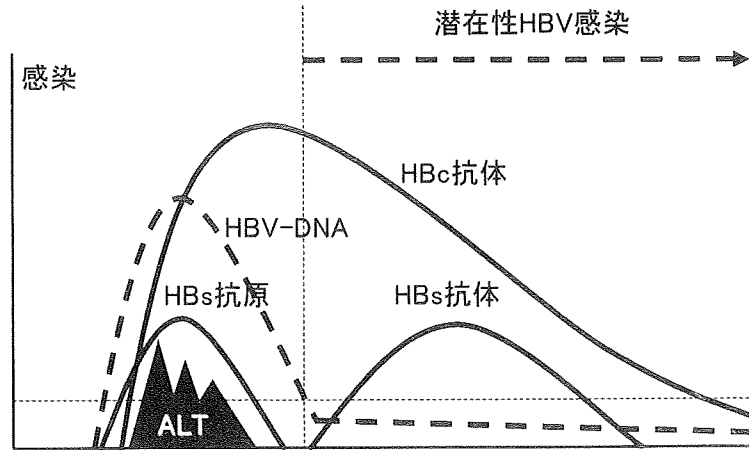
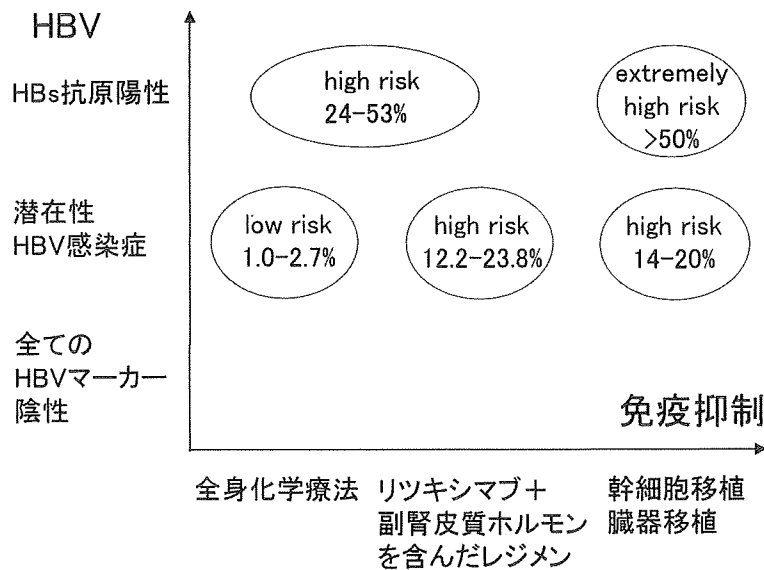


図3 HBV 初感染後のウイルスマーカーの動き



Kusumoto S, et al: Int J Hematol 90: 13-23, 2009より引用改変

図4 HBV 再活性化のリスク. 潜在性 HBV 感染症よりも HBs 抗原陽性キャリアが、また薬物による免疫抑制効果が強いほど、HBV 再活性化のリスクが高い。

検出されず、HBV による肝障害も起こらない。また、不顕性感染から潜在性 HBV 感染状態になったケースが多いため、患者本人も認識しておらず、病歴から絞り込むことはできない。従って、潜在性 HBV 感染のスクリーニングに最も有用なマーカーである HBe 抗体を測定する必要がある。福岡、北九州の赤十字血液センターからの報告によると、全献血者における HBe 抗体陽性の頻度は1.1%であるが、HBe 抗体陽性率は年齢とともに上昇するため、免疫抑制・化学療法を受ける患者の年齢層を考えると、潜在性 HBV 感染症例にはしばしば遭遇すると予想される¹⁰⁾¹¹⁾。

最近、この潜在性 HBV 感染症例からの HBV 再活性

化の報告が相次いでいる。固形癌に対して通常行われる化学療法は、潜在性 HBV 感染症例における HBV 再活性化のリスクはそれほど高くはないが、悪性リンパ腫に対するレジメンでリツキシマブと副腎皮質ホルモンを含んだ R-CHOP 療法は高リスクであることが知られている¹²⁾¹³⁾ (図4)。その他、クローン病や慢性関節リウマチに使用される抗 TNF 製剤のインフリキシマブやメソトレキセートでも HBV 再活性化の報告がある¹⁴⁾¹⁵⁾。

Ⅲ. B型急性肝炎と HBV 再活性化の違い

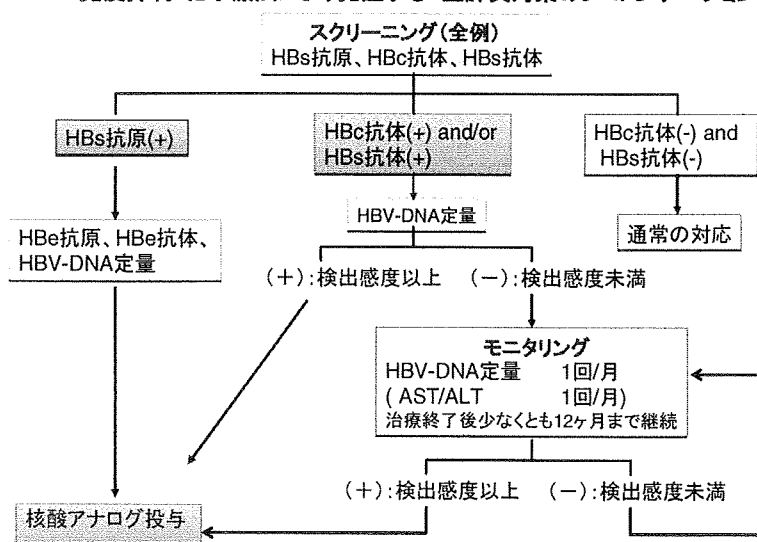
HBV による急性肝障害の代表である B 型急性肝炎は、成人ではほとんどが性行為感染によるものである

表2 HBV再活性化とB型肝炎の臨床像の比較

| | 再活性化 | B型肝炎 | P |
|----------------------------|------|-------|--------|
| 年齢 | 63 | 33 | <0.001 |
| 男性 | 59% | 71% | |
| ALT (IU/L) | 929 | 2,300 | <0.001 |
| T.Bil (mg/dL) | 10.3 | 6.4 | |
| Alb (g/dL) | 3.2 | 3.6 | <0.001 |
| PT (%) | 65.0 | 75.0 | |
| HBV DNA (log copies/mL) | 7.5 | 5.5 | <0.001 |
| 劇症化 | 22% | 9% | 0.048 |
| 肝関連死 | 26% | 4% | <0.001 |

Umemura T, et al : Clin Infect Dis 47 : e52-56, 2008 より引用改変

免疫抑制・化学療法により発症するB型肝炎対策の推奨



坪内博仁、他：肝臓 50: 38-42, 2009より引用

図5 厚生労働省の研究班からの推奨

表3 消化器内科・肝臓専門医へのコンサルトのタイミング

| |
|---|
| ①化学療法・免疫抑制療法予定患者の検査で、HBs抗原またはHBc抗体が陽性であった場合 |
| ②潜在性HBV感染において、治療中および治療後の経過中にHBV-DNAが検出されるようになった場合 |

が、肝細胞内に侵入し増殖したウイルスに対する正常な免疫応答によって肝障害は起こる。多くの場合、不顕性感染や軽度の急性肝炎で終わることが多い。ウイルス肝炎の中ではB型肝炎は劇症化率が高いことで知られているが、B型肝炎の中でみると、この劇症化率は1-2%である。これに比べてHBV再活性化は、HBV増殖や免疫状態などB型肝炎とはかなり異なる様相を呈する。

UmemuraらがB型肝炎とHBV再活性化を比

較した成績を報告している¹⁶⁾。HBVの感染経路、HBc抗体陽性率および基礎疾患を考えると当然であるが、HBV再活性化症例は有意に平均年齢が高い。ALTのピーク値はHBV再活性化症例のほうがむしろ有意に低い。肝予備能を表す検査値は悪化傾向にある。HBV量はHBV再活性化症例が100倍高値であり、特筆すべきは、劇症化と肝関連死がHBV再活性化症例において有意に高いことである。つまり、HBV再活性化は一旦起こると劇症化しやすく、劇症化すると内科的な救命率

は極めて低いと考えられている(表2)。従って、肝障害を発症させないような予防策が重要になる。

IV. HBV 再活性化の予防について

HBV 再活性化の予防のポイントは、①潜在性 HBV 感染を慎重に検討すること、②HBV DNA の陽性化を見逃さないことに尽きる。HBV DNA の増加は肝障害出現の数ヶ月も前に先行してみられるため、これを見逃さない限り、その後の対策を講じる時間的な余裕は十分ある。潜在性 HBV 感染であれば免疫抑制・化学療法を行いながら、月1回の HBV DNA の測定で経過観察するが、HBs 抗原陽性または HBV DNA が検出される場合は核酸アナログを投与する。厚生労働省の研究班からガイドラインが出されており(図5)、これに従えば HBV 再活性化は予防できる。

しかし、このガイドラインは多少煩雑であり、肝臓専門医がコンサルタントとして介入することで、最低限の項目のみに絞った福岡大学病院独自のマニュアルを提案し、これを免疫抑制・化学療法施行医に周知徹底して行う方が、安全管理という観点から見るとリーズナブルであると思われる。そこで、免疫抑制・化学療法前に HBs 抗原と HBc 抗体を測定し、いずれかが陽性であれば肝臓専門医にコンサルトを、さらに経過観察の指示があった場合は HBV DNA が検出された時点で再度肝臓専門医にコンサルトすることを提案する(表3)。核酸アナログの投与は必要に応じて行うが、その終了時期などコンセンサスが得られていない部分もあるため、核酸アナログ投与は肝臓専門医が行うことが望ましいと考える。

最近、HBc 抗体陰性症例からの HBV 再活性化の報告もある¹⁷⁾。長い経過によって HBc 抗体が陰性化した HBV キャリアであると考えられるが、このような例外的な症例もあるという認識をもっておく必要はある。また、最も強力な免疫抑制が必要となる造血肝細胞移植や臓器移植の場合は、HBs 抗体と HBV DNA の検査を必須にするなど、独自のプロトコルが必要である。

結 び

HBV 再活性化は化学療法などの治療後、思いもよらない時に突然起こる重篤な急性肝障害であり、免疫抑制・化学療法に関わる医師、看護師および薬剤師は、HBV 再活性化を常に認識していなければならない。さらに、マニュアルに従ってウイルスマーカーの検査を忘れないで行うことが重要であるが、それをチェックするシステム作りが、HBV 再活性化の確実な予防のために必要である。

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Ribavirin dose reduction raises relapse rate dose-dependently in genotype 1 patients with hepatitis C responding to pegylated interferon alpha-2b plus ribavirin

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SUMMARY. The impact of ribavirin exposure on virologic relapse remains controversial in combination therapy with pegylated interferon (Peg-IFN) and ribavirin for patients with chronic hepatitis C (CH-C) genotype 1. The present study was conducted to investigate this. Nine hundred and eighty-four patients with CH-C genotype 1 were enrolled. The drug exposure of each medication was calculated by averaging the dose actually taken. For the 472 patients who were HCV RNA negative at week 24 and week 48, multivariate logistic regression analysis showed that the degree of fibrosis ($P = 0.002$), the timing of HCV RNA negativation ($P < 0.001$) and the mean doses of ribavirin ($P < 0.001$) were significantly associated with relapse, but those of Peg-IFN were not. Stepwise reduction of the ribavirin dose was associated with a stepwise increase in relapse rate from 11%

to 60%. For patients with complete early virologic response (c-EVR) defined as HCV RNA negativity at week 12, only 4% relapse was found in patients given ≥ 12 mg/kg/day of ribavirin and ribavirin exposure affected the relapse even after treatment week 12, while Peg-IFN could be reduced to 0.6 μ g/kg/week after week 12 without the increase of relapse rate. Ribavirin showed dose-dependent correlation with the relapse. Maintaining as high a ribavirin dose as possible (≥ 12 mg/kg/day) during the full treatment period can lead to suppression of the relapse in HCV genotype 1 patients responding to Peg-IFN alpha-2b plus ribavirin, especially in c-EVR patients.

Keywords: chronic hepatitis C, drug exposure, pegylated interferon plus ribavirin, virologic relapse.

INTRODUCTION

Combination therapy of pegylated interferon (Peg-IFN) plus ribavirin is very effective for patients with chronic hepatitis C

Abbreviations: CH-C, chronic hepatitis C; c-EVR, complete early virologic response; ETR, end-of-treatment virologic response; Hb, haemoglobin; HCV, hepatitis C virus; IFN, interferon; LVR, late virologic response; Peg-IFN, pegylated interferon; PP, per protocol; Plt, platelet; RVR, rapid virologic response; SVR, sustained virologic response; VR, virologic response; WBC, white blood cell.

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(CH-C). However, sustained virologic response (SVR) in current therapy occurs in only 40–50% of patients with hepatitis C virus (HCV) genotype 1 [1–4]. Also, SVR is reduced in patients with genotype 1 who require reduction of either Peg-IFN or ribavirin, although dose reduction has little influence on SVR in those with genotype 2 or 3 [1–3,5,6]. Therefore, it is important to clarify the degree to which these medications can be reduced without adversely affecting SVR in patients with CH-C genotype 1.

In an early report on the relationship between drug exposure and antiviral effect in patients with CH-C genotype 1, patients who received $\geq 80\%$ of their total planned cumulative doses of Peg-IFN and ribavirin for $\geq 80\%$ of the scheduled duration of therapy had an SVR of 51% compared with only 34% for patients who received lesser amounts of one or both

medications [7]. On the other hand, Shiffman *et al.* [8] recently reported that reducing ribavirin did not affect SVR as long as the dose of Peg-IFN was maintained, while reducing the Peg-IFN dose significantly reduced SVR. The results of these observations are consistent with respect to the effect of Peg-IFN on SVR. However, what is controversial is whether or not reducing the ribavirin dose affects the antiviral effect.

Adding ribavirin to either interferon (IFN) or Peg-IFN monotherapy for patients with CH-C genotype 1 has been shown to reduce the relapse rate in large randomized trials [1,2,9–11]. In detail, adding ribavirin to the usual IFN monotherapy (3MIU, three-times-weekly) in 48-week treatment raised the end-of-treatment virologic response (ETR) rate from approximately 30% to 50% and also lowered the relapse rate from mid-40% to approximately 20% [9–11]. Lindsay *et al.* [12] reported that Peg-IFN alpha-2b (Peg-IFN α -2b) monotherapy (1.5 μ g/kg, once-weekly), as compared with IFN alpha-2b (IFN α -2b) monotherapy (3MIU, three-times-weekly), improved ETR (49% vs. 24%), but not the relapse rate (53% vs. 50%). In the trial of Peg-IFN alpha-2a (Peg-IFN α -2a) plus ribavirin vs IFN α -2b plus ribavirin or Peg-IFN α -2a alone, the ETR rates were 69%, 52% and 59%, and the relapse rates were 19%, 15% and 52%, respectively [2]. These findings from large-scale trials indicate that the main role of ribavirin is to reduce relapse in the combination therapy with Peg-IFN, although ribavirin affects both ETR and relapse in combination therapy with the usual IFN.

In the present study, we tried to determine whether or not dose reduction of ribavirin (or Peg-IFN) has an effect on virologic relapse in Peg-IFN plus ribavirin treatment for patients with CH-C genotype 1.

PATIENTS AND METHODS

Patients

This study was a multicentre trial conducted by Osaka University Hospital and other institutions participating in the Osaka Liver Forum. A total of 984 patients with CH-C were enrolled in this study between December 2004 and September 2006, and treated with a combination of Peg-IFN α -2b plus ribavirin. The baseline characteristics of the patients are shown in Table 1. All patients were Japanese infected with HCV genotype 1 and a viral load of more than 10^5 IU/mL. Patients were excluded from this study if they had decompensated cirrhosis or other forms of liver disease (alcohol liver disease, autoimmune hepatitis), coinfection with hepatitis B or anti-human immunodeficiency virus. This study was conducted according to the ethical guidelines of the 1975 Declaration of Helsinki and informed consent was obtained from each patient.

Treatment

All patients received Peg-IFN α -2b (PEGINTRON; Schering-Plough, Kenilworth, NJ, USA) plus ribavirin (REBETOL;

Table 1 Baseline characteristics of patients and drug doses at start of treatment

| Factor | Mean \pm SD or <i>n</i> |
|--|---------------------------|
| <i>n</i> | 984 |
| Age (years) | 56.3 \pm 10.1 |
| Sex (male/female) | 555/429 |
| Body weight (kg) | 61.8 \pm 11.5 |
| History of IFN treatment | 575/409 (160/182) |
| Naïve/experienced (relapser/nonresponder)* | |
| White blood cells (/mm ³) | 5052 \pm 1550 |
| Neutrophils (/mm ³) | 2577 \pm 1092 |
| Red blood cells ($\times 10^4$ /mm ³) | 442 \pm 47 |
| Haemoglobin (g/dL) | 14.1 \pm 1.4 |
| Platelets ($\times 10^4$ /mm ³) | 15.9 \pm 5.5 |
| AST (IU/L) | 66 \pm 45 |
| ALT (IU/L) | 79 \pm 61 |
| Serum HCV RNA (kIU/mL) [†] | 1600 |
| Histology (METAVIR) [‡] | |
| Fibrosis; 0/1/2/3/4 | 49/314/197/105/18 |
| Activity; 0/1/2/3 | 23/329/304/27 |
| Peg-IFN dose (μ g/kg/week) | 1.45 \pm 0.17 |
| Ribavirin dose (mg/kg/day) | 11.4 \pm 1.6 |

AST, aspartate aminotransferase; ALT, alanine aminotransferase; HCV, hepatitis C virus. *Viral response to previous treatment was unknown in 57 patients, and 10 patients had discontinued treatment. [†]Data shown are median values. [‡]301 missing.

Schering-Plough) for the duration of the study of 48 weeks. As a starting dose, Peg-IFN α -2b was given subcutaneously once weekly at a dosage of 60–150 μ g/kg based on body weight (body weight 35–45 kg, 60 μ g; 46–60 kg, 80 μ g; 61–75 kg, 100 μ g; 76–90 kg, 120 μ g; 91–120 kg, 150 μ g) and ribavirin was given orally twice a day at a total dose of 600–1000 mg/day based on body weight (body weight <60 kg, 600 mg; 60–80 kg, 800 mg; >80 kg, 1000 mg) according to the manufacturer's drug information available in Japan.

Dose reduction and discontinuance

Dose modification also followed, as a rule, the manufacturer's drug information according to the intensity of the haematologic adverse effects. The dose of Peg-IFN α -2b was reduced to 50% of the assigned dose when the white blood cell (WBC) count was below 1500/mm³, the neutrophil count below 750/mm³ or the platelet (Plt) count below 8×10^4 /mm³, and was discontinued when the WBC count was below 1000/mm³, the neutrophil count below 500/mm³ or the Plt count below 5×10^4 /mm³. Ribavirin was also reduced from 1000 mg to 600 mg, 800 mg to 600 mg, or 600 mg to 400 mg when the haemoglobin (Hb)

concentration decreased to less than 10 g/dL, and was discontinued when the Hb concentration decreased to less than 8.5 g/dL. Both Peg-IFN α -2b and ribavirin had to be discontinued if there was a need to discontinue one of the drugs. No ferric medicine or haematopoietic growth factors, such as epoetin alpha, or granulocyte-macrophage colony stimulating factor, were administered.

Virologic assessment and definition of virologic response

Serum HCV RNA level was quantified using the COBAS AMPLICOR HCV MONITOR test, version 2.0 (detection range 6–5000 kIU/mL; Roche Diagnostics, Branchburg, NJ, USA) and qualitatively analysed using the COBAS AMPLICOR HCV test, version 2.0 (lower limit of detection 50 IU/mL; Roche Diagnostics). Complete early virologic response (c-EVR) was defined as the absence of detectable serum HCV RNA at treatment week 12, the late virologic response (LVR) was defined as undetectable serum HCV RNA for the first time at 13–24 weeks of treatment, and the virologic response (VR) was defined as HCV RNA negativity at week 24 and week 48. SVR was defined as the absence of detectable serum HCV RNA at week 72. Patients with less than a 2-log decrease in HCV RNA level at treatment week 12 compared with the baseline had to stop treatment according to the protocol and were regarded as nonresponders. All patients with detectable serum HCV RNA at treatment week 24 were also considered to be nonresponders and were excluded from further treatment.

Assessment of drug exposure

The amounts of Peg-IFN α -2b and ribavirin actually taken by each patient during the full treatment period were evaluated by reviewing the medical records. The mean doses of Peg-IFN α -2b and ribavirin were calculated individually as averages on the basis of body weight at baseline: Peg-IFN α -2b expressed as μ g/kg/week, ribavirin expressed as mg/kg/day.

Evaluation of impact of drug exposure on virologic relapse

We evaluated the relationship between the drug exposure of both drugs and relapse by two different methods, univariate and multivariate analysis for relapse and independent evaluation of both drugs for relapse according to the degree of drug exposure. The former was performed with the factors of mean administration doses of both drugs, including the factors at baseline and the timing of HCV RNA negativiation. The latter was examined by classifying Peg-IFN α -2b exposure into five categories (up to 0.6 μ g/kg; from 0.6 to less than 0.9 μ g/kg; from 0.9 to less than 1.2 μ g/kg; from 1.2 to less than 1.5 μ g/kg; from 1.5 μ g/kg) and ribavirin exposure into five categories (up to 6 mg/kg; from 6 to less than 8 mg/kg; from 8 to less than 10 mg/kg; from 10 to less than 12 mg/kg; from 12 mg/kg).

Statistical analysis

Baseline data are expressed as means \pm SD or median values. Virologic response was evaluated using per protocol (PP) analysis. To analyse the difference between baseline data including drug exposure and virologic response, univariate analysis using the Mann–Whitney *U*-test or chi-square test and multivariate analysis using logistic regression analysis were performed. The significance of trends in values was determined with the Mantel–Haenszel chi-square test. A two-tailed *P* value <0.05 was considered significant. The analysis was conducted with SPSS version 15.0J (SPSS Inc., Chicago, IL, USA).

RESULTS

Progress of patients and dose reduction of Peg-IFN α -2b and ribavirin

The progress of patients in this study is shown in Fig. 1. Of the 984 patients, 903 completed 12 weeks of treatment and the c-EVR rate was 49% (445/903), based on PP study. To analyse for relapse, 472 patients with VR were assessed, with 178 (38%) showing Peg-IFN dose reduction without discontinuation and 246 (52%) with ribavirin dose reduction without discontinuation during the full (48 weeks) treatment period. The relapse rate was 26% (125/472) in the patients with undetectable HCV RNA level at the end of treatment. No difference was found in relapse rates between the IFN naïve patients and IFN experienced patients (IFN naïve; 25%, 72/287 vs IFN experienced; 29%, 53/185, *P* = 0.40). The SVR rate was 43% (347/812) in the PP study.

Impact of drug exposure during 0–48 weeks on relapse among patients with VR

The mean dose of Peg-IFN α -2b actually taken during the full treatment period by each patient was 1.32 μ g/kg/week (range, 0.49–2.16 μ g/kg/week; median, 1.38 μ g/kg/week) and that of ribavirin was 9.8 mg/kg/day (range, 3.3–16.2 mg/kg/day; median, 10.1 mg/kg/day) in patients with VR.

The result of univariate analysis for relapse among the patients with VR is shown in Table 2a. The degree of fibrosis, the timing of HCV RNA negativiation, Plt value and the mean doses of ribavirin were factors significantly associated with relapse, but those of Peg-IFN α -2b were not. The mean dose of ribavirin as well as the degree of fibrosis and the timing of HCV RNA negativiation was selected as a significant independent factor by multivariate logistic regression analysis (Table 2b).

Next, we analysed the relationship of the relapse rate and the mean ribavirin dose. The overall relapse rate among patients with VR was 26% (125/472). The

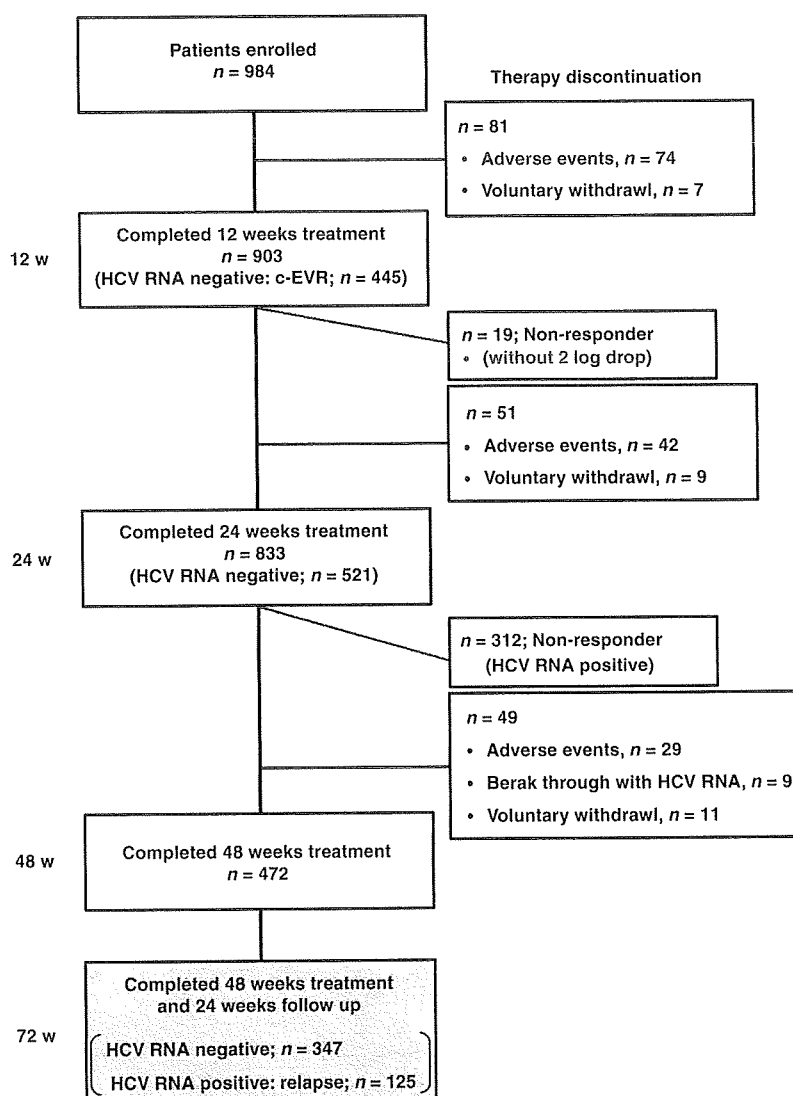


Fig. 1 Flow of patients throughout the study.

relapse rate was 60% (9/15) in patients receiving less than 6 mg/kg/day of ribavirin, and declined to 41% (32/79) at 6–8 mg/kg/day, 27% (34/124) at 8–10 mg/kg/day, 22% (43/193) at 10–12 mg/kg/day and 11% (7/61) in patients given ≥ 12 mg/kg/day ($P < 0.0001$). Figure 2 shows the relationship of the relapse rate and the mean ribavirin dose for two dosage groups of Peg-IFN α -2b: the group given ≥ 1.4 $\mu\text{g}/\text{kg}/\text{week}$ of Peg-IFN and that given < 1.4 $\mu\text{g}/\text{kg}/\text{week}$ (1.4 $\mu\text{g}/\text{kg}/\text{week}$ was the median value). In both groups, ribavirin was dose-dependently correlated with relapse. More than 12 mg/kg/day of the mean ribavirin exposure could suppress the relapse rate to 20% (4/20) in the group given < 1.4 $\mu\text{g}/\text{kg}/\text{week}$ and strongly suppress it to 7% (3/41) in the group given ≥ 1.4 $\mu\text{g}/\text{kg}/\text{week}$ of Peg-IFN.

Impact of drug exposure during 0–48 weeks on relapse according to the timing of HCV RNA negativation

Relapse rates among patients with c-EVR

The overall relapse rate among patients with c-EVR was 19% (75/391). We separately analysed the relapse rate among the patients with c-EVR according to the degree of exposure to both drugs. Table 3a shows the relapse rates among the patients with c-EVR according to the categories of Peg-IFN α -2b and ribavirin doses during the full treatment period. The relapse rate showed a decline according to the increase in the dose of ribavirin ($P = 0.0002$). The relapse rate was suppressed at an average of 15% (13–16%) in the patients who received 10–12 mg/kg/day of ribavirin, and the average was only 4% for those who received more than 12 mg/kg/day

Table 2 Factors associated with relapse among the patients with virologic response

| (a) Univariate analysis | | | | |
|--|---|-------------|--------------|---------|
| Factor | Nonrelapser | Relapser | P value | |
| <i>n</i> | 347 | 125 | | |
| Age (years) | 53.9 ± 10.7 | 56.2 ± 9.2 | 0.07 | |
| Sex (male/female) | 213/134 | 66/59 | 0.09 | |
| Serum HCV RNA (kIU/mL)* | 1600 | 1800 | 0.34 | |
| White blood cells (/mm ³) | 5335 ± 1517 | 5075 ± 1428 | 0.08 | |
| Neutrophils (/mm ³) | 2797 ± 1143 | 2625 ± 1021 | 0.17 | |
| Red blood cells (×10 ⁴ /mm ³) | 450 ± 45 | 446 ± 50 | 0.25 | |
| Haemoglobin (g/dL) | 14.3 ± 1.4 | 14.2 ± 1.5 | 0.45 | |
| Platelets (×10 ⁴ /mm ³) | 17.6 ± 5.3 | 16.4 ± 5.1 | 0.03 | |
| AST (IU/L) | 60 ± 42 | 58 ± 33 | 0.75 | |
| ALT (IU/L) | 75 ± 60 | 71 ± 50 | 0.98 | |
| Histology (METAVIR) [†] | | | | |
| Fibrosis: 0–2/3–4 | 222/20 | 74/19 | 0.002 | |
| Activity: 0–1/2–3 | 140/102 | 52/41 | 0.75 | |
| Peg-IFN dose (µg/kg/week) [‡] | 1.33 ± 0.26 | 1.27 ± 0.29 | 0.07 | |
| Ribavirin dose (mg/kg/day) [‡] | 10.1 ± 1.9 | 9.1 ± 2.1 | <0.001 | |
| Virologic response [§] : c-EVR/LVR | 316/31 | 75/50 | <0.001 | |
| (b) Multivariate analysis | | | | |
| Factor | Category | Odds ratio | 95% CI | P value |
| Platelets | By 1 × 10 ⁴ /mm ³ | – | – | NS |
| Fibrosis [¶] | 0–2/3–4 | 1/3.192 | 1.515–6.725 | 0.002 |
| Ribavirin dose [‡] | By 1 mg/kg/day | 0.790 | 0.696–0.896 | <0.001 |
| Virologic response [§] | c-EVR/LVR | 1/6.290 | 3.385–11.690 | <0.001 |

AST, aspartate aminotransferase; ALT, alanine aminotransferase; HCV, hepatitis C virus; c-EVR, complete early virologic response; LVR, late virologic response; NS, not significant difference Peg-IFN, pegylated interferon.

*Data shown are median values. †137 missing. ‡Mean doses during 0–48 weeks. §The timing of HCV RNA negativitation.

¶METAVIR fibrosis score.

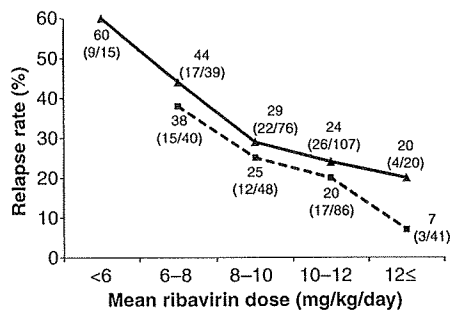


Fig. 2 Relapse rate according to Peg-IFN α -2b and ribavirin doses during treatment of patients who completed treatment, which was stratified with the mean ribavirin doses. (— \blacktriangle) Group with the mean Peg-IFN dose <1.4 μ g/kg/week; (--- \blacksquare) Group with the mean Peg-IFN dose \geq 1.4 μ g/kg/week. The ribavirin dose was dose-dependently correlated with the virologic relapse in both groups ($P < 0.0001$). There was no significant difference between the two Peg-IFN α -2b-dose groups ($P = 0.17$).

of ribavirin. In contrast, the relapse rate was not affected by the dose of Peg-IFN α -2b when the patients were given more than 0.9 μ g/kg/week of Peg-IFN α -2b. On the other hand, with respect to patients with rapid virologic response (RVR) defined as the absence of detectable serum HCV RNA at treatment week 4 ($n = 41$), none showed relapse and all attained SVR irrespective of the dose of Peg-IFN α -2b or ribavirin (prevalence of patients: the mean dose of Peg-IFN α -2b; <0.9 : 0.9–1.2 : 1.2–1.5 : 1.5 μ g/kg/week \leq 7 : 17 : 34 : 42%, the mean dose of ribavirin; <8 : 8–10 : 10–12 : 12 mg/kg/day \leq 15 : 24 : 41 : 20%).

Relapse rates among patients with LVR

Among the patients with LVR, the ribavirin exposure during treatment was also the factor correlated adversely with the relapse rate ($P = 0.03$). However, the overall relapse rate was 62% (50/81), which was much higher than that of the c-EVR patients ($P < 0.0001$) and 45% (5/11) of patients with LVR relapsed even in the group given more than 12 mg/kg/day of the average ribavirin dose (Table 3b).

Table 3 Relapse rate according to Peg-IFN and ribavirin doses during week 0–48 for patients with c-EVR and LVR who completed 48 weeks of treatment

| (a) C-EVR | | | | | | | | | | |
|--|-----------------------------|--------|-----|----------|-----|----------|-----|---------|-------|----------|
| Peg-IFN dose ($\mu\text{g}/\text{kg}/\text{week}$) [†] | Ribavirin dose (mg/kg/day)* | | | | | | | | Total | |
| | 12 \leq | 10–12 | | 8–10 | | <8 | | | | |
| ≥ 1.5 | 0% | (0/28) | 13% | (4/31) | 14% | (3/21) | 29% | (5/17) | 12% | (12/97) |
| 1.2–1.5 | 20% | (2/10) | 16% | (16/100) | 25% | (16/65) | 23% | (7/30) | 20% | (41/205) |
| 0.9–1.2 | 0% | (0/7) | 13% | (2/15) | 15% | (2/13) | 38% | (6/16) | 20% | (10/51) |
| <0.9 | 0% | (0/5) | 15% | (2/13) | 55% | (6/11) | 44% | (4/9) | 32% | (12/38) |
| Total | 4% | (2/50) | 15% | (24/159) | 25% | (27/110) | 31% | (22/72) | 19% | (75/391) |

| (b) LVR | | | | | | | | | | |
|--|---|--------|-----|---------|------|--------|------|---------|-------|---------|
| Peg-IFN dose ($\mu\text{g}/\text{kg}/\text{week}$) [§] | Ribavirin dose (mg/kg/day) [‡] | | | | | | | | Total | |
| | 12 \leq | 10–12 | | 8–10 | | <8 | | | | |
| ≥ 1.5 | 43% | (3/7) | 50% | (1/2) | 100% | (2/2) | 100% | (4/4) | 67% | (10/15) |
| 1.2–1.5 | | (1/1) | 60% | (12/20) | 29% | (2/7) | 82% | (9/11) | 62% | (24/39) |
| <1.2 | 33% | (1/3) | 50% | (6/12) | 60% | (3/5) | 86% | (6/7) | 59% | (16/27) |
| Total | 45% | (5/11) | 56% | (19/34) | 50% | (7/14) | 86% | (19/22) | 62% | (50/81) |

Peg-IFN, pegylated interferon; c-EVR, complete early virologic response; LVR, late virologic response.

* $P = 0.0002$ for comparison of the four ribavirin groups. [†] $P = 0.08$ for comparison of the four Peg-IFN groups. [‡] $P = 0.03$ for comparison of the four ribavirin groups. [§] $P = 0.57$ for comparison of the three Peg-IFN groups.

Impact of dose reduction after week 12 on relapse among patients with c-EVR

Among c-EVR patients with no or little reduction of Peg-IFN α -2b (the average dose ≥ 1.2 $\mu\text{g}/\text{kg}/\text{week}$) during the first 12 weeks, no significant difference was found in the relapse rate between those whose average dose of Peg-IFN α -2b was reduced to 0.6–1.2 $\mu\text{g}/\text{kg}/\text{week}$ during 12–48 weeks (17%, 7/41) and those without reduction of Peg-IFN α -2b (average dose ≥ 1.2 $\mu\text{g}/\text{kg}/\text{week}$) (18%, 53/295) ($P = 0.86$) (Table 4a). Reducing the dose of Peg-IFN α -2b after week 12 in patients in whom HCV RNA had already become undetectable before week 12 did not appear to adversely influence virologic relapse when the average dose of Peg-IFN α -2b was more than 0.6 $\mu\text{g}/\text{kg}/\text{week}$ during 12–48 weeks, irrespective of the mean dose of Peg-IFN α -2b during the first 12 weeks. On the other hand, the ribavirin dose reduction after week 12 tended to affect the relapse rate in patients given ≥ 10 mg/kg/day of the ribavirin dose during the first 12 weeks (Table 4b).

Impact of drug exposure during 0–48 weeks on relapse among VR patients with advanced fibrosis

In the evaluation of the 39 patients with VR with progression of fibrosis or cirrhosis (METAVIR fibrosis score 3 or 4) enrolled in this study, ribavirin exposure during treatment significantly correlated with relapse (nonrelapser, 10.5 ± 2.1 mg/kg/day vs relapser, 8.8 ± 2.3 mg/kg/day; $P = 0.007$). Among patients with advanced fibrosis (score 3–4),

the relapse rate in patients given ≥ 10 mg/kg/day of the average ribavirin dose was significantly low (36%, 9/25) in comparison with that in patients given < 10 mg/kg/day of ribavirin (71%, 10/14) ($P = 0.048$).

DISCUSSION

Previous studies have suggested that reducing the ribavirin dose within the first 12–20 weeks of treatment in patients with HCV genotype 1 was associated with a decline of SVR [7,13,14]. However, Shiffman *et al.* [8] recently reported that reducing the mean dose of ribavirin during the first 20 weeks of treatment had little impact on relapse for patients with CH-C genotype 1 and that SVR may not be adversely affected as long as the total cumulative ribavirin dose remains above 60%. As the reason for the inconsistency in the impact of reducing ribavirin on the antiviral effect, it was suggested that sample sizes of the previous studies were insufficient to assess the impact of reducing the dose of ribavirin independent of Peg-IFN. However, in Shiffman's study, while the impact of reducing the dose of Peg-IFN or ribavirin on SVR was indeed closely examined independently of each other with a large sample size, the subjects were limited to patients with advanced fibrosis or cirrhosis and prior nonresponse to Peg-IFN \pm ribavirin who were enrolled in the Hepatitis Antiviral Long-term Treatment Against Cirrhosis (HALT-C) trial. Reddy *et al.* [15] analysed the drug exposure retrospectively for 569 CH-C patients with genotype 1 enrolled in clinical trials of Peg-IFN α -2a plus

Table 4 Relapse rate according to drug doses during week 0–12 and 12–48 for patients with c-EVR who completed 48 weeks of treatment

| (a) Peg-IFN | | 12–48 weeks | | | |
|--|------------|--------------|--------------|--------------|-------------|
| Peg-IFN dose (mean, $\mu\text{g}/\text{kg}/\text{week}$) | | ≥ 1.2 | 0.9–1.2 | 0.6–0.9 | < 0.6 |
| 0–12 weeks | ≥ 1.2 | 18% (53/295) | 17% (5/30) | 18% (2/11) | (1/1) |
| | 0.9–1.2 | – | 22% (4/18) | 33% (4/12) | 60% (3/5) |
| | < 0.9 | (0/1) | (0/1) | 17% (2/12) | 20% (1/5) |
| Total* | | 18% (53/296) | 18% (9/49) | 23% (8/35) | 45% (5/11) |
| (b) Ribavirin | | 12–48 weeks | | | |
| Ribavirin dose (mean, $\text{mg}/\text{kg}/\text{day}$) | | ≥ 12 | 10–12 | 8–10 | < 8 |
| 0–12 weeks | ≥ 12 | 4% (2/47) | 13% (3/23) | 13% (1/8) | 33% (1/3) |
| | 10–12 | – | 15% (18/123) | 22% (12/54) | 20% (5/25) |
| | 8–10 | – | (1/1) | 26% (10/38) | 26% (10/39) |
| | < 8 | – | – | – | 40% (12/30) |
| Total† | | 4% (2/47) | 15% (22/147) | 23% (23/100) | 29% (28/97) |

c-EVR, complete early virologic response; Peg-IFN, pegylated interferon.

* $P = 0.18$ for comparison of the four Peg-IFN groups. † $P < 0.0001$ for comparison of the four ribavirin groups.

ribavirin, and concluded that SVR was not affected adversely by ribavirin reduction unless the cumulative ribavirin exposure was less than 60%. This supported Shiffman's data, but in Reddy's study, the stepwise reduction in ribavirin dose was shown to be associated with a stepwise increase in relapse rate from 19% to 54%. Thus, the impact of ribavirin drug exposure on the antiviral effect (relapse) in patients with CH-C genotype 1 remains unclear. Further examination is needed to determine whether or not ribavirin can be reduced to a certain degree without adversely affecting virologic relapse or SVR in Peg-IFN and ribavirin combination therapy for CH-C genotype 1.

In order to raise the SVR rate in patients with genotype 1, two strategies are possible: one is enhancing the virologic response of HCV RNA negativity and another is reducing relapse. In Peg-IFN plus ribavirin treatment, raising the doses of either or both drugs (dose-up strategy) is the only way to enhance the virologic response of HCV RNA negativity, but this is always accompanied by a high risk and the discontinuation rate can increase with the dose-up of drug, although the virologic response among patients completing the therapy can be improved [16,17]. Therefore, in this study, we tried to manage the drug dose to reduce relapse in virologic responders with HCV RNA negativity. Large-scale clinical trials [1,2,9–12] have revealed that adding ribavirin to IFN or Peg-IFN monotherapy for patients with CH-C reduced the relapse rate from approximately 50% to under 20%. Bronowicki *et al.* [18] examined the effect of ribavirin on CH-C genotype 1 in Peg-IFN α -2a plus ribavirin treatment

by randomizing patients with HCV RNA negativity by week 24 into two groups, one continuing with ribavirin and the other receiving Peg-IFN α -2a alone after week 24. As a result, the virologic responders who stopped ribavirin treatment at week 24 were found to have a significantly higher rate of breakthroughs during therapy and higher relapse rates after therapy in comparison with those who received Peg-IFN plus ribavirin for the full treatment period (relapse rate; 42% vs. 29%, $P = 0.02$). These findings indicate that ribavirin plays a very important role in reducing relapse. However, the relationship between ribavirin dose and relapse rate has not been examined in detail. Considering that ribavirin has little influence on HCV RNA negativation [1,2,9–12], its dose impact on the antiviral effect should be carefully examined, not for the SVR rate of all patients, but for the relapse rate of patients responding to Peg-IFN plus ribavirin, as evaluating of ribavirin by SVR including HCV RNA negativation cannot differentiate it from the strong influence of the Peg-IFN effect, which affects HCV RNA negativation dose-dependently [19]. Here, we examined the correlation between the average dose of drugs and the virologic relapse for patients responding to the treatment.

We performed univariate and multivariate analysis for relapse among the factors of mean administration doses of both drugs, including baseline factors and the timing of HCV RNA negativation. We found exposure to ribavirin dose, timing of HCV RNA negativation and the degree of liver fibrosis to be the independent factors affecting the virologic relapse in patients with VR. This indicates that management

of the ribavirin dose, which is the variable factor, unlike baseline factors, plays an important role in suppressing the virologic relapse in patients with CH-C genotype 1 treated by Peg-IFN plus ribavirin treatment. This suggests that maintaining the ribavirin dose should lower the relapse rate even in patients with advanced fibrosis who are liable to relapse. In fact, among patients with advanced fibrosis (METAVIR score 3–4), the relapse rate in those given ≥ 10 mg/kg/day of the average ribavirin dose was significantly lower than that in patients given < 10 mg/kg/day of ribavirin (36% vs. 71%). However, the sample size was too small for subsequent analysis with stratification. Further study is needed to clarify the impact of ribavirin dose on viral relapse in patients with progression of fibrosis.

The relapse rate among patients with c-EVR showed a decline according to the increase in ribavirin dose during treatment week 0–48 and was not affected by the Peg-IFN α -2b dose when the patients were given more than 0.9 μ g/kg/week of Peg-IFN α -2b. Among the patients with c-EVR, none with RVR had a relapse and all attained SVR irrespective of the dose of Peg-IFN α -2b or ribavirin. Examination of the impact of dose reduction after week 12 on relapse among patients with c-EVR showed that the ribavirin dose reduction after week 12 tended to affect the relapse rate in patients given ≥ 10 mg/kg/day of the ribavirin dose during the first 12 weeks, while the Peg-IFN α -2b dose after week 12 could be reduced without any increase in relapse rate in patients given more than 0.6 μ g/kg/week of the average dose of Peg-IFN α -2b. On the other hand, maintaining the ribavirin did not lead to reduce the relapse rate in patients with LVR. About half relapsed even when given ≥ 12 mg/kg/day of the average ribavirin dose. This suggested that the relapse rate could not be reduced by management of the ribavirin dose in patients with LVR. Extended therapy should be chosen in LVR patients as shown in the previous studies [20–23].

Shiffman *et al.* [24] recently reported that maintaining the Hb level with epoetin alpha did not enhance SVR if ribavirin was started at the standard dose (800–1400 mg/day, mean dose 13.3 mg/kg/day), although discontinuance and the reduction rates of ribavirin were decreased and a higher mean dose of ribavirin was administered in comparison with those treated with Peg-IFN plus ribavirin without epoetin. If these findings apply to patients with CH-C genotype 1, this would suggest that the ribavirin dose does not need to be maintained during treatment with Peg-IFN plus ribavirin, which would not agree with our findings. However, closer examination of the Shiffman *et al.* study shows that Peg-IFN plus a higher dose of ribavirin (1000–1600 mg/day, mean dose 15.2 mg/kg/day) with epoetin was found to suppress the relapse rate and enhance SVR. These data agree with ours with respect to the point that higher doses of ribavirin are associated with a lower relapse rate. What differs is the ribavirin dose needed to suppress the relapse. This is likely to be due to ethnic differences between the subjects. In Shiffman's study, approximately 40% were African-American

in whom the virologic response is well established as being significantly lower than those of other ethnic groups [25,26], while in our study, all subjects were Japanese. In the African-Americans treated with Peg-IFN plus standard-dose ribavirin, the relapse rate (calculated from 48% of ETR and 19% of SVR) was 60%, while 18% relapse (from 38% of ETR and 31% of SVR) occurred in those given Peg-IFN plus high-dose ribavirin. The relapse rate of patients with c-EVR in our study was 19%, which was very close to that for those with Peg-IFN plus high-dose ribavirin in Shiffman's study. Ribavirin does not have a direct antiviral action against HCV [27,28], and is considered to play an important role in accelerating HCV-infected cell clearance [29] and eradicating them completely when an immune response against infected cells is induced by IFN or Peg-IFN [30,31]. Therefore, the difference between patients who are easy or difficult to treat due to ethnic differences or differences in response to Peg-IFN can result in the need for different doses of ribavirin to suppress the relapse rate in patients with CH-C genotype 1.

In conclusion, our results have demonstrated that ribavirin is dose-dependently correlated with a relapse in patients with CH-C genotype 1 responding to Peg-IFN plus ribavirin. Maintaining a high dose (≥ 12 mg/kg/day) of ribavirin during the full treatment period could strongly suppress the relapse in such patients, while Peg-IFN α -2b could be reduced without affecting relapse in patients with c-EVR. This possibility should be explored in a prospective study.

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Pegylated interferon alpha-2b (Peg-IFN α -2b) affects early virologic response dose-dependently in patients with chronic hepatitis C genotype 1 during treatment with Peg-IFN α -2b plus ribavirin

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SUMMARY. Chronic hepatitis C (CH-C) genotype 1 patients who achieved early virologic response have a high probability of sustained virologic response (SVR) following pegylated interferon (Peg-IFN) plus ribavirin therapy. This study was conducted to evaluate how reducing drug doses affects complete early virologic response (c-EVR) defined as hepatitis C virus (HCV) RNA negativity at week 12. Nine hundred eighty-four patients with CH-C genotype 1 were enrolled. Drug doses were evaluated independently on a body weight base from doses actually taken. From multivariate analysis, the mean dose of Peg-IFN α -2b during the first 12 weeks was the independent factor for c-EVR ($P = 0.02$), not ribavirin. The c-EVR rate was 55% in patients receiving ≥ 1.2 $\mu\text{g}/\text{kg}/\text{week}$ of Peg-IFN, and declined to 38% at 0.9–1.2 $\mu\text{g}/\text{kg}/\text{week}$, and 22% in patients given < 0.9 $\mu\text{g}/\text{kg}/\text{week}$ ($P < 0.0001$). Even with stratified analysis according to

ribavirin dose, the dose-dependent effect of Peg-IFN on c-EVR was observed, and similar c-EVR rates were obtained if the dose categories of Peg-IFN were the same. Furthermore, the mean dose of Peg-IFN during the first 12 weeks affected HCV RNA negativity at week 24 ($P < 0.0001$) and SVR ($P < 0.0001$) in a dose-dependent manner. Our results suggest that Peg-IFN was dose-dependently correlated with c-EVR, independently of ribavirin dose. Thus, maintaining the Peg-IFN dose as high as possible during the first 12 weeks can yield HCV RNA negativity and higher c-EVR rates, leading to better SVR rates in patients with CH-C genotype 1.

Keywords: chronic hepatitis C, drug dose, early virologic response, HCV RNA negativity, pegylated interferon plus ribavirin, sustained virologic response.

Abbreviations: c-EVR, complete EVR; CH-C, chronic hepatitis C; EVR, early virologic response; G-CSF, granulocyte-macrophage colony stimulating factor; Hb, haemoglobin; HCV, hepatitis C virus; Peg-IFN, pegylated interferon; Plt, platelet; SVR, sustained virologic response; WBC, white blood cell.

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INTRODUCTION

Pegylated interferon (Peg-IFN) plus ribavirin therapy can improve anti-viral efficacy for patients with chronic hepatitis C [1–5], and the prognosis of patients in whom hepatitis C virus (HCV) is successfully eradicated improves markedly [6–10]. However, HCV still persists in approximately half of genotype 1 patients treated with Peg-IFN plus ribavirin [2–4]. Therefore, the treatment method needs to be well managed in order to maximize the virologic response in these patients with HCV genotype 1.

In order to achieve sustained virologic response (SVR), earlier virologic response is very important for patients with chronic hepatitis C (CH-C) genotype 1. A high SVR rate (65–72%) was found in patients who achieved early virologic response (EVR) defined as a 2-log decrease in HCV RNA level at week 12, but only 0–3% SVR was seen in patients without EVR [3,11]. Additionally, complete EVR (c-EVR), which means HCV RNA negativity at week 12, is more strongly related to SVR [3].

The relationship between drug exposure and anti-viral effect has been reported in several papers [2,11–15]. McHutchison *et al.* [12] demonstrated that the SVR rate in patients who received $\geq 80\%$ of their total planned doses of Peg-IFN and ribavirin for $\geq 80\%$ of the scheduled duration of therapy was significantly higher than that of patients who received $< 80\%$ of one or both drugs (51% vs 34%) and also suggested that the impact of dose reduction was greatest in patients for whom the dose had to be decreased within the first 12 weeks of treatment. In a subsequent analysis, reducing the dose of Peg-IFN and ribavirin to $< 80\%$ of the full planned dose within the first 12 weeks was reported to reduce EVR rate from 80 to 33% [11]. Thus, drug adherence during the first 12 weeks has been shown to be very important for attaining EVR and SVR, but it remains obscure whether either drug can be reduced to a certain degree without adversely affecting the treatment efficacy.

In the present study, we examined the correlation between c-EVR and drug doses which are evaluated on a body weight basis from drug doses actually taken, in order to clarify the necessary drug exposure of Peg-IFN and ribavirin for achieving a higher c-EVR rate in patients with CH-C genotype 1.

PATIENTS AND METHODS

Patients

The current study was a retrospective, multicenter trial conducted by Osaka University Hospital and other institutions participating in the Osaka Liver Forum. A total of 984 patients with CH-C treated with a combination of Peg-IFN α -2b plus ribavirin were enrolled in this study between December 2004 and September 2006. The baseline characteristics of the patients are summarized in Table 1. All patients were Japanese, their mean age was 56.3 ± 10.1 years, and 56% were males. The mean serum alanine aminotransferase level was 79 ± 61 IU/L.

Patients eligible for this study were those who were infected with HCV genotype 1 and had a viral load of more than 10^5 IU/mL, but were negative for hepatitis B surface antigen or anti-human immunodeficiency virus. Patients were excluded from this study if they had decompensated cirrhosis or other forms of liver disease (alcohol liver disease, autoimmune hepatitis). Informed consent was obtained from each patient included in this study. This study was conducted according to the ethical guidelines of the 1975 Dec-

Table 1 Baseline characteristics of patients

| Factor | Mean \pm SD or number |
|--|-------------------------|
| <i>n</i> | 984 |
| Age (year) | 56.3 ± 10.1 |
| Sex: male/female | 555/429 |
| Body weight (kg) | 61.8 ± 11.5 |
| History of interferon treatment | |
| Naïve/experienced | 575/409(160/182) |
| (relapser/nonresponder)* | |
| White blood cells (per mm ³) | 5052 ± 1550 |
| Neutrophils (per mm ³) | 2577 ± 1092 |
| Red blood cells ($\times 10^4$ /mm ³) | 442 ± 47 |
| Haemoglobin (g/dL) | 14.1 ± 1.4 |
| Platelets ($\times 10^4$ /mm ³) | 15.9 ± 5.5 |
| AST (IU/L) | 66 ± 45 |
| ALT (IU/L) | 79 ± 61 |
| Serum HCV RNA (kIU/mL) [†] | 1600 |
| Histology (METAVIR) [‡] | |
| Fibrosis; 0/1/2/3/4 | 49/314/197/105/18 |
| Activity; 0/1/2/3 | 23/329/304/27 |

AST, aspartate aminotransferase; ALT, alanine aminotransferase; HCV, hepatitis C virus.

*Viral response to previous treatment was unknown in 57 patients, and 10 patients had discontinued treatment. [†]Data shown are median values. [‡]301 missing.

laration of Helsinki and informed consent was obtained from each patient.

Treatment

All patients received Peg-IFN α -2b (PEGINTRON; Schering-Plough, Kenilworth, NJ, USA) plus ribavirin (REBETOL; Schering-Plough) for the duration of the study of 48 weeks. Peg-IFN α -2b was given subcutaneously once weekly at a dosage of 60–150 μ g/kg based on body weight (body weight 35–45 kg, 60 μ g; 46–60 kg, 80 μ g; 61–75 kg, 100 μ g; 76–90 kg, 120 μ g; 91–120 kg, 150 μ g) and ribavirin was given orally twice a day at a total dose of 600–1000 mg/day based on body weight (body weight ≤ 60 kg, 600 mg; 60–80 kg, 800 mg; > 80 kg, 1000 mg), according to a standard treatment protocol for Japanese patients.

Dose reduction

Dose modification followed, as a rule, the manufacturer's drug information according to the intensity of the haematological adverse effects. The dose of Peg-IFN α -2b was reduced to 50% of the assigned dose if the white blood cell (WBC) count declined to < 1500 /mm³, the neutrophil count to < 750 /mm³ or the platelet (Plt) count to $< 8 \times 10^4$ /mm³, and was discontinued if the WBC count declined to < 1000 /

mm³, the neutrophil count to <500/mm³ or the Plt count to <5 × 10⁴/mm³. Ribavirin was also reduced from 1000 to 600 mg, or 800 to 600 mg, or 600 to 400 mg if the haemoglobin (Hb) level decreased to <10 g/dL, and was discontinued if the Hb level decreased to <8.5 g/dL. Both Peg-IFN α -2b and ribavirin had to be discontinued if there was a need to discontinue one of the drugs. During this therapy, ferric medicine or haematopoietic growth factors, such as erythropoietin alpha, or granulocyte-macrophage colony stimulating factor (G-CSF), were not administered.

Virologic assessment and definition of virologic response

Serum HCV RNA level was quantified using the COBAS AMPLICOR HCV MONITOR test, version 2.0 (detection range 6–5000 IU/mL; Roche Diagnostics, Branchburg, NJ, USA) and qualitatively analysed using the COBAS AMPLICOR HCV test, version 2.0 (lower limit of detection 50 IU/mL). The c-EVR was defined as the absence of detectable serum HCV RNA at treatment week 12, and SVR was defined as the absence of detectable serum HCV RNA at week 72. Patients with less than a 2-log decrease in HCV RNA level at treatment week 12 compared with the baseline had to stop treatment and were regarded as nonresponders. All patients with detectable serum HCV RNA at treatment week 24 were also considered nonresponders and excluded from further treatment.

Assessment of drug exposure

The amounts of Peg-IFN α -2b and ribavirin actually taken by each patient during the first 12 weeks of the treatment were evaluated by reviewing the medical records. The mean doses of both drugs were calculated individually as averages on the basis of body weight at baseline: Peg-IFN α -2b expressed as μ g/kg/week, and ribavirin expressed as mg/kg/day.

Evaluation of impact of drug exposure on c-EVR

We evaluated the relationship between the drug exposure of both drugs and c-EVR by univariate and multivariate analysis for c-EVR, using the factors of mean administration doses of both drugs during the first 12 weeks and the factors at baseline. Furthermore, Peg-IFN α -2b dose (average dose per body weight and per week) was classified into five categories (up to 0.6 μ g/kg; from 0.6 to <0.9 μ g/kg; from 0.9 to <1.2 μ g/kg; from 1.2 to <1.5 μ g/kg; from 1.5 μ g/kg and above). Ribavirin exposure was classified into four categories (up to 8 mg/kg; from 8 to <10 mg/kg; from 10 to <12 mg/kg; from 12 mg/kg and above), in order to examine the impact of Peg-IFN dose exposure on c-EVR. This impact was also evaluated based on the percentage of the total prescribed dose and compared with that based on the mean dose per body weight.

Statistical analysis

Baseline data for various demographic, biochemical and virologic characteristics of the patients are expressed as mean \pm SD or median values. To analyse the relationship between baseline data including drug exposure and c-EVR, univariate analysis using the Mann–Whitney *U*-test or chi-squared test and multivariate analysis using logistic regression analysis were performed. The significance of trends in values was determined with the Mantel–Haenszel chi-square test. A two-tailed *P*-value < 0.05 was considered significant. Statistical analysis was conducted with SPSS version 15.0J (SPSS Inc., Chicago, IL, USA).

RESULTS

Progress of patients treated with Peg-IFN α -2b and ribavirin

Of the 984 patients, 81 discontinued treatment because of adverse events ($n = 74$) or voluntary withdrawal ($n = 7$) by treatment week 12. The 903 patients who completed 12 weeks of treatment were assessed for c-EVR. During 12–48 weeks of treatment, 331 of the nonresponders and nine of breakthrough discontinued treatment, as did 91 patients (adverse events, $n = 71$; voluntary withdrawal, $n = 20$). A total of 472 patients completed 48 weeks of treatment.

Drug reduction and virologic response

Peg-IFN α -2b was reduced without discontinuation in 29% ($n = 266$) and ribavirin was reduced without discontinuation in 40% ($n = 359$) of the 903 patients who completed 12 weeks of treatment. The c-EVR rate was 49% (445/903) and HCV RNA was negative at week 24 in 60% (542/903) of patients who completed 12 weeks of treatment. Of the 445 patients with c-EVR, 327 patients achieved SVR (73%). Only 7% of the 458 patients without c-EVR did so.

Impact of dose exposure of Peg-IFN α -2b and ribavirin on c-EVR

The mean dose of Peg-IFN α -2b actually taken during the first 12 weeks by each patient was 1.33 μ g/kg/week (range 0.41–2.16 μ g/kg/week; median 1.40 μ g/kg/week) and that of ribavirin was 10.4 mg/kg/day (range 2.9–16.2 mg/kg/day; median 10.6 mg/kg/day).

The mean doses of both drugs and the factors at baseline correlated with the c-EVR were assessed by univariate and multivariate logistic regression analyses. Univariate analysis showed that factors significantly associated with c-EVR were age, sex, WBC, neutrophils, red blood cells, Hb, Plt, aspartate aminotransferase, the degree of liver fibrosis and the mean doses of Peg-IFN α -2b and ribavirin during the first 12 weeks (Table 2). The factors selected as significant by the univari-

Table 2 Univariate analysis for c-EVR among patients who completed 12 weeks treatment

| Factor | c-EVR (+) | c-EVR (-) | P-value |
|--|-------------|-------------|---------|
| <i>n</i> | 445 | 458 | |
| Age (year) | 54.4 ± 10.4 | 57.5 ± 9.6 | <0.001 |
| Sex: male/female | 267/178 | 237/221 | 0.01 |
| Serum HCV RNA (kIU/mL)* | 1500 | 1600 | 0.28 |
| White blood cells (per mm ³) | 5336 ± 1536 | 4818 ± 1547 | <0.001 |
| Neutrophils (per mm ³) | 2789 ± 1133 | 2398 ± 1038 | <0.001 |
| Red blood cells (×10 ⁴ /mm ³) | 450 ± 46 | 435 ± 49 | <0.001 |
| Haemoglobin (g/dL) | 14.3 ± 1.4 | 13.9 ± 1.4 | <0.001 |
| Platelets (×10 ⁴ /mm ³) | 17.3 ± 5.2 | 15.0 ± 5.6 | <0.001 |
| AST (IU/L) | 62 ± 44 | 69 ± 44 | <0.001 |
| ALT (IU/L) | 77 ± 64 | 80 ± 57 | 0.07 |
| Histology (METAVIR) [†] | | | |
| Fibrosis: 0–2/3–4 | 273/37 | 247/74 | <0.001 |
| Activity: 0–1/2–3 | 171/139 | 159/162 | 0.16 |
| Peg-IFN dose (µg/kg/week) [‡] | 1.39 ± 0.22 | 1.28 ± 0.30 | <0.001 |
| Ribavirin dose (mg/kg/day) [‡] | 10.6 ± 1.7 | 10.1 ± 2.1 | 0.002 |

c-EVR, complete early virologic response; HCV, hepatitis C virus; AST, aspartate aminotransferase; ALT, alanine aminotransferase; Peg-IFN, pegylated interferon. *Data shown are median values. [†]272 missing. [‡]Mean doses during 0–12 weeks.

Table 3 Multivariate analysis for c-EVR among patients who completed 12 weeks treatment

| Factor | Category | Odds ratio | 95% CI | P-value |
|-----------------------------|---|------------|-------------|---------|
| Age | by 1 year | 0.982 | 0.966–0.999 | 0.04 |
| Sex | male/female | – | – | NS |
| Neutrophils | by 100/mm ³ | 1.017 | 1.002–1.033 | 0.03 |
| Red blood cells | by 1 × 10 ⁴ /mm ³ | – | – | NS |
| Haemoglobin | by 1 g/dL | – | – | NS |
| Platelets | by 1 × 10 ⁴ /mm ³ | 1.051 | 1.014–1.088 | <0.01 |
| AST | by 1 IU/L | – | – | NS |
| Fibrosis* | 0–2/3–4 | – | – | NS |
| Peg-IFN dose [†] | by 0.1 µg/kg/week | 1.079 | 1.011–1.151 | 0.02 |
| Ribavirin dose [†] | by 1 mg/kg/day | – | – | NS |

95% CI, 95% confidence interval; Peg-IFN, c-EVR, complete early virologic response; pegylated interferon; N.S., No Significant difference; AST, aspartate aminotransferase.

*METAVIR fibrosis score. [†]Mean doses during 0–12 weeks.

ate analysis were evaluated by multivariate logistic regression analysis. The mean dose of Peg-IFN α -2b during the first 12 weeks was the independent factor for c-EVR ($P = 0.02$), apart from the neutrophils ($P = 0.03$) and Plt value at baseline ($P < 0.01$) and age ($P = 0.04$) (Table 3). In contrast, the mean dose of ribavirin during the first 12 weeks showed no correlation with c-EVR.

The c-EVR rates were 54% (137/253) and 56% (246/443) for patients who received ≥ 1.5 and 1.2–1.5 µg/kg/week of Peg-IFN α -2b on average during the first 12 weeks, and declined to an average rate of 38% (40/105) in patients given 0.9–1.2 µg/kg/week of Peg-IFN α -2b, and an average rate of 22% (22/102) in patients given < 0.9 µg/kg/week ($P < 0.0001$) (Table 4). The c-EVR rate among the patients

with ≥ 1.2 µg/kg/week of Peg-IFN α -2b was significantly higher than that of the patients with < 1.2 µg/kg/week [≥ 1.2 µg/kg/week, 55% (383/696) vs < 1.2 µg/kg/week, 30% (62/207), $P < 0.0001$].

Next, we analysed the impact of Peg-IFN α -2b on c-EVR in stratified analysis according to ribavirin dose. Figure 1 shows the relationship of c-EVR and the degree of Peg-IFN α -2b exposure for two groups of ribavirin doses: the group with ≥ 10.6 mg/kg/day of ribavirin and that with < 10.6 mg/kg/day (10.6 mg/kg/day was the median value). In either group, the mean dose of Peg-IFN α -2b was dose-dependently correlated with c-EVR ($P < 0.0001$), and c-EVR rates were very similar in both groups if the dose categories of Peg-IFN α -2b were the same.

Table 4 The c-EVR rate according to Peg-IFN and ribavirin doses during weeks 0–12 for patients who completed 12 weeks treatment

| Ribavirin dose (mg/kg/day)** | Peg-IFN α -2b dose (μ g/kg/week),* | | | | Total |
|------------------------------|--|---------------|--------------|--------------|---------------|
| | ≥ 1.5 | 1.2–1.5 | 0.9–1.2 | <0.9 | |
| ≥ 12 | 57% (60/105) | 61% (22/36) | 38% (6/16) | 22% (2/9) | 54% (90/166) |
| 10–12 | 54% (46/85) | 58% (154/267) | 36% (14/39) | 23% (11/47) | 51% (225/438) |
| 8–10 | 50% (25/50) | 53% (52/99) | 52% (15/29) | 18% (4/22) | 48% (96/200) |
| <8 | 46% (6/13) | 44% (18/41) | 24% (5/21) | 21% (5/24) | 34% (34/99) |
| Total | 54% (137/253) | 56% (246/443) | 38% (40/105) | 22% (22/102) | 49% (445/903) |

c-EVR, complete early virologic response; Peg-IFN, pegylated interferon.

* $P < 0.0001$ for comparison of the four Peg-IFN groups. ** $P = 0.05$ for comparison of the four ribavirin groups.

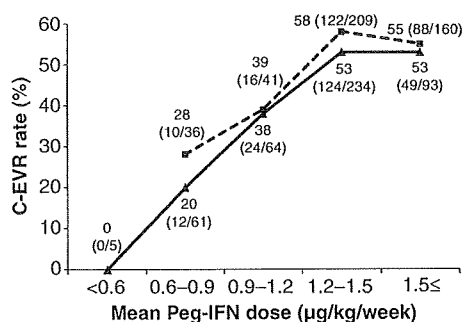


Fig. 1 Complete-EVR rate according to pegylated interferon alpha-2b (Peg-IFN α -2b) and ribavirin doses during weeks 0–12 for patients who completed 12 weeks of treatment. (— \blacktriangle) Group with the mean ribavirin dose <10.6 mg/kg/day. (--- \blacksquare) Group with the mean ribavirin dose ≥ 10.6 mg/kg/day. The Peg-IFN α -2b dose was dose-dependently correlated with c-EVR in both groups ($P < 0.0001$). There was no significant difference between the two ribavirin-dose groups ($P = 0.19$).

c-EVR rates according to Peg-IFN α -2b drug exposure using a percentage cut off and mean dose cut off

Table 5 shows the c-EVR rates according to the category of Peg-IFN α -2b doses during the first 12 weeks based on the

Table 5 The c-EVR rate according to Peg-IFN dose during weeks 0–12 based on the percentage of the planned dose and the mean doses

| Peg-IFN α -2b dose (μ g/kg/week) | $\geq 80\%$ | 60–80% | <60% | Total |
|--|----------------|---------------|-------------|---------------|
| ≥ 1.2 | 55%* (371/679) | 71%** (12/17) | – | 55% (383/696) |
| <1.2 | 32% (6/19) | 38% (35/92) | 22% (21/96) | 30% (62/207) |
| Total | 54% (377/698) | 43% (47/109) | 21% (21/96) | 49% (445/903) |

c-EVR, complete early virologic response; Peg-IFN, pegylated interferon.

* $P < 0.05$; patients with ≥ 1.2 μ g/kg/week vs <1.2 μ g/kg/week among the patients with more than 80% of the total prescribed dose of Peg-IFN α -2b. ** $P = 0.01$; patients with ≥ 1.2 μ g/kg/week vs <1.2 μ g/kg/week among the patients with more than 60–80% of the total prescribed dose of Peg-IFN α -2b.

percentage of the total prescribed dose and the mean doses. The whole c-EVR rate was 54% (377/698) for patients who received more than 80% of the prescribed dose, and 43% (47/109) in patients given 60–80% of the prescribed dose, and 21% (21/96) in patients given <60% of the prescribed dose of Peg-IFN α -2b. Among patients given $\geq 80\%$ of the prescribed dose of Peg-IFN α -2b, the c-EVR rate was significantly lower in patients given <1.2 μ g/kg/week of Peg-IFN α -2b than those given ≥ 1.2 μ g/kg/week (32% vs 55%, $P < 0.05$). On the other hand, even in patients given 60–80% of the prescribed dose of Peg-IFN α -2b, if they were given ≥ 1.2 μ g/kg/week of Peg-IFN α -2b, a higher c-EVR rate was attained in comparison with those given <1.2 μ g/kg/week (71% vs 38%, $P = 0.01$); the c-EVR rate in patients given 60–80% of the prescribed dose and ≥ 1.2 μ g/kg/week of Peg-IFN α -2b was not inferior to that in patients given $\geq 80\%$ of the prescribed dose and ≥ 1.2 μ g/kg/week of Peg-IFN α -2b.

Impact of dose exposure of Peg-IFN α -2b during the first 12 weeks of the treatment on HCV RNA negativity at week 24 and SVR

Patients positive for HCV RNA at week 24 week during Peg-IFN α -2b and ribavirin treatment were regarded as non-responders and stopped treatment [11]. We analysed the

relationship between the dose exposure to Peg-IFN α -2b during the first 12 weeks and HCV RNA negative rates at week 24 or SVR in 903 patients completing 12 weeks of treatment. As a result, HCV RNA negative rates at week 24 and SVR rates declined according to the decrease in the dose of Peg-IFN α -2b during the 12 weeks of treatment; patients given ≥ 1.5 , 1.2–1.5, 0.9–1.2 and < 0.9 $\mu\text{g}/\text{kg}/\text{week}$ of Peg-IFN α -2b during the first 12 weeks of the treatment showed HCV RNA negativity of 63%, 66%, 48% and 39%, respectively ($P < 0.0001$), and SVR of 46%, 43%, 30% and 20%, respectively ($P < 0.0001$).

DISCUSSION

Adherence to ribavirin was reported to be the important factor for EVR as well as that to Peg-IFN in most previous studies [2,11,12]. However, the drug exposure of Peg-IFN α -2b and ribavirin had not been analysed independently with respect to their individual influence on the anti-viral effect in these studies. Adherence to both drugs may be related factors, i.e. most patients who can tolerate a high dose of Peg-IFN are in good condition and thus can also receive a high dose of ribavirin. In the present study, the impact of the dose of Peg-IFN α -2b and ribavirin on the anti-viral effect was evaluated by multivariate logistic regression analysis, using the mean administration doses of both drugs during the first 12 weeks and baseline factors. As a result, the dose exposure of Peg-IFN α -2b was found to be the significant factor affecting c-EVR as well as baseline factors such as age, neutrophils and Plt values, but not ribavirin. This suggests that the c-EVR rate can be raised by maintaining the dose of Peg-IFN α -2b during the first 12 weeks in patients with disadvantageous factors at baseline. In fact, the c-EVR rate was higher in those who received ≥ 1.2 $\mu\text{g}/\text{kg}$ of Peg-IFN α -2b than in those given < 1.2 $\mu\text{g}/\text{kg}$ of Peg-IFN α -2b for aged patients over 60 years of age (≥ 1.2 $\mu\text{g}/\text{kg}$; 46% vs < 1.2 $\mu\text{g}/\text{kg}$; 28%, $P < 0.01$) or for patients with a low Plt value ($< 12 \times 10^4/\text{mm}^3$) (≥ 1.2 $\mu\text{g}/\text{kg}$; 45% vs < 1.2 $\mu\text{g}/\text{kg}$; 22%, $P < 0.001$). Therefore, a marked dose reduction of Peg-IFN α -2b should not be risked at the start even for aged patients or patients with lower Plt value, which is indicative of advanced fibrosis. The administration of ≥ 1.2 $\mu\text{g}/\text{kg}/\text{week}$ of Peg-IFN α -2b is desirable as a starting dose for achieving c-EVR even in these patients: that of < 1.2 $\mu\text{g}/\text{kg}/\text{week}$ can lead to a non-viral response or a late viral response. Independent evaluation of the c-EVR rate according to the degree of the ribavirin dose showed a stepwise decline as the total cumulative dose of Peg-IFN α -2b decreased. Therefore, the dose of Peg-IFN α -2b should be maintained as high as possible even in patients who have to reduce Peg-IFN α -2b to < 1.2 $\mu\text{g}/\text{kg}/\text{week}$. Using G-CSF for patients who develop severe neutropenia and are forced to decrease Peg-IFN can be beneficial, especially in the first 12 weeks.

The goal of 80% of the planned drug dosage for 80% of the assigned duration was derived from an adherence criterion

that had been adopted previously for assessment of the efficacy of other pharmaceutical agents, such as drugs to treat cancer and human immunodeficiency virus [16]. However, in Peg-IFN plus ribavirin therapy for patients with CH-C, the planned administration dose [17,18] differs on a body weight basis by 27% for Peg-IFN α -2b and 40% for ribavirin among patients of 50–100 kg of body weight, which would be equivalent to the same rate differences for 80% of the planned drug dosage. In detail, the target dose of Peg-IFN α -2b scheduled to be administered is 1.5 $\mu\text{g}/\text{kg}$, but the usual dose for the individual patient is from 1.28 to 1.76 $\mu\text{g}/\text{kg}/\text{week}$ based on body weight among patients weighing 50–100 kg according to the practice guidelines of the American Association for the Study of Liver Diseases and the manufacturer's drug information in the USA and Europe [17,18]. The range of ribavirin dose per kg of body weight is from 12 to 20 mg/kg/day. Therefore, in this study, the drug exposure was assessed from the average dose per kg of body weight.

In the evaluation of c-EVR rates according to Peg-IFN α -2b drug exposure using a percentage cut off and mean dose cut off in this study, the c-EVR rate of patients given < 1.2 $\mu\text{g}/\text{kg}/\text{week}$ of Peg-IFN α -2b was low (32%) even in those who received $\geq 80\%$ of the total planned doses of Peg-IFN α -2b. If given ≥ 1.2 $\mu\text{g}/\text{kg}/\text{week}$ of Peg-IFN α -2b, the c-EVR rate (71%) in patients who received 60–80% of the total doses was not inferior to that in patients given $\geq 80\%$ of the total dose of Peg-IFN α -2b (54%). This means that patients whose starting dose of Peg-IFN α -2b is < 1.5 $\mu\text{g}/\text{kg}/\text{week}$ should not have their dosage reduced to 80% of the planned dose (< 1.2 $\mu\text{g}/\text{kg}/\text{week}$) in order to have a higher probability of c-EVR, while those given ≥ 1.5 $\mu\text{g}/\text{kg}/\text{week}$ of Peg-IFN α -2b at the start can have their dosage reduced to 80% (≥ 1.2 $\mu\text{g}/\text{kg}/\text{week}$) without lowering the c-EVR rate. Thus, the drug dose on a body weight basis itself should be examined as an index of the drug exposure in order to evaluate the anti-viral effect of both drugs accurately for patients with CH-C.

As for the impact of the drug exposure to ribavirin on c-EVR, the drug dose of ribavirin during the first 12 weeks was shown to have no relationship with the c-EVR rate, although it was precisely evaluated in this study, using doses actually taken on body weight. However, ribavirin can be more effective for decreasing the viral relapse after interferon or Peg-IFN α -2b and ribavirin combination therapy in patients with CH-C genotype 1 [2,3,19–24]. Recently, Shiffman *et al.* [15] have reported that a higher starting dose of ribavirin (1000–1600 mg/day) plus a regular dose of Peg-IFN α -2b with epoetin was associated with a lower relapse rate in treatment with CH-C genotype 1. Considering the viral relapse after treatment, it is thought that the ribavirin dose should not be reduced quickly in patients with mild side effects, even though it does not affect c-EVR. In fact, among the patients who attained c-EVR, a higher rate of viral relapse was found in the patients given < 10 mg/kg/day of the mean ribavirin dose during 48 weeks in comparison