Improved Sustained Virological Response Following Treatment with Pegylated-Interferon Alpha-2b Compared with Alpha-2a, Both with Ribavirin, for Chronic Hepatitis C Infection with Genotypes 2 and 3

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ABSTRACT

Purpose: The optimal formulation of pegylated interferon α (PEG-IFα) as a part of combination therapy with ribavirin (RBV) is uncertain for patients infected with hepatitis C Genotypes 2 and 3. Methods: A multivariate analysis of prospectively collected treatment data from two tertiary centres on 351 treatment naïve HCV Genotype 2 or 3 patients who received PEG-IFα-2a or b plus ribavirin. Results: Univariate analyses demonstrate that PEG-IFα-2b based on regimens achieved a higher sustained virological response (SVR) than PEG-IFα-2a (77.9% versus 62.0%, P = 0.0012). On multivariate analyses, PEG-IFα-2b appeared superior to PEG-IFα-2a with an odds ratio (OR) and 95% confidence interval (CI 95) for SVR of 2.19 (CI 95 1.35 - 3.52, P = 0.0005). Genotype was a significant predictor of outcome in the multivariate model with 80% of Genotype 2 but only 67.7% of Genotype 3 subjects achieving SVR (OR 2.66 [CI 95 1.35 - 5.92]). Increasing age was negatively associated with SVR (OR 0.97 [CI 95 0.94 - 0.99]). Some of the differences in SVR are explained by higher relapse rates with PEG-IFα-2a (P = 0.009). Conclusions: PEG-IFα-2b and RBV achieve higher SVR rates than PEG-IFα-2a and RBV in Genotypes 2 and 3 chronic HCV infections. There is less relapse with PEG-IFα-2b. Genotype 2 infections are considerably easier to cure. SVR is higher in younger patients. These findings should influence a choice of PEG-IFα in the era of direct acting anti-viral drugs in therapy of Genotypes 2 and 3.

KEYWORDS

Hepatitis C; Pegylated Interferon; Ribavirin; SVR; Multivariate Analysis; Genotype 2; Genotype 3

1. Introduction

Chronic hepatitis C virus (HCV) infection is a significant public health issue. Worldwide, approximately 170 million people are infected [1]. The risk of developing chronic infection after acute exposure to HCV approaches 85% [2]. It is estimated that approximately 1% of the Australian community have chronic HCV infection [2]. Approximately 5% - 20% of chronically infected patients progress to cirrhosis in the long term with 3% - 5% of these patients developing hepatocellular carcinoma annually (HCC) [3].

Successful treatment of chronic HCV infection is defined by a negative PCR for HCV-RNA 6 months post therapy—a sustained virological response (SVR). Interferon (IF) alpha was the first established treatment [4], with further enhancement of outcomes seen after the addition of the antiviral nucleoside analogue, ribavirin (RBV) [5,6]. The advent of pegylated (PEG) forms of IF alpha led to further improvements in response rates [7,8]. The combination of PEG-IF with ribavirin has remained the standard of care throughout the 2000s [9-11].
With broadening of choices of therapy to include genotype specific direct acting antivirals (DAAs) [12,13] and the understanding that numerous host and viral factors affect outcomes, choices affecting therapy are likely to become highly individualised [14]. As such, any additional information that might affect the choice of IF backbone to that therapy is especially valuable. International guidelines [15] suggest that there is no practical difference between the two commercially available formulations: PEG-IFα-2a (Pegasys, Roche Pharmaceuticals, Geneva, Switzerland) and PEG-IFα-2b (Pegintron, Schering Plough Corporation, NJ, USA) both combined with RBV. A recent Cochrane review and meta-analysis [16] have reported improved SVR with PEG-IFα-2a, particularly for HCV Genotype 1. However, the evidence for the optimal choice of PEG-IFα in the treatment of other HCV genotypes is less clear.

The aim of this study was to compare treatment outcomes between PEG-IFα-2a and PEG-IFα-2b (plus RBV) in patients with Genotype 2 and 3 infections, in a “real-world” clinical environment, through a retrospective review of our experience in two large, tertiary-referral, hospital-based hepatitis treatment services in Australia.

2. Materials and Methods

Data on all patients treated for HCV between 2002 and 2008 were prospectively collected through the clinical hepatitis services of two tertiary metropolitan hospitals in Perth, Western Australia [Fremantle Hospital (FH) and Royal Perth Hospital (RPH)]. These two hospitals serve a population of approximately one million. Patient demographic factors, viral genotype, liver biopsy result, and treatment allocation were prospectively documented. HCV viral load at baseline was not routinely available at our centre during this time. The degree of histologic liver fibrosis was reported using the METAVIR score [17].

The choice of PEG-IF formulation was at the treating physician’s discretion and was prescribed with RBV according to licensed indications. Haematological side effects were managed by dose reduction of RBV or PEG-IF in a standard fashion. Sustained virological response was the primary outcome of interest and was defined by the presence of a negative HCV PCR result 6 months after cessation of treatment. Any patient who received at least one dose of PEG-IF therapy, but did not attain this outcome was considered to have not achieved SVR. An end of treatment (EOT) HCV PCR was considered a secondary outcome as was relapse during the 6 month period after cessation of treatment.

We used R for all basic and multivariate analyses [18]. Data are presented as median (interquartile range [IQR]) or proportions for continuous and categorical variables, respectively. Mann-Whitney U and Chi-squared tests were applied for univariate analyses for continuous and categorical variables, respectively. The METAVIR score was considered to be an ordinal variable. Multivariate analysis was performed using logistic regression. Variables other than age were included based on biological plausibility and P < 0.10 on univariate analysis. Backward stepwise logistic regression was applied and the most parsimonious model was chosen using Aikake’s Information Criterion (AIC). To account for the large amount of missing data for weight and fibrosis scores, we imputed missing data using the R package “AMELIA” [19]. Briefly, each variable of interest was defined as nominal, ordinal or continuous. Non-parametric, continuous variables were log-transformed and intuitive constraints placed on the possible output data. Following multivariate imputation, AMELIA provides visual and statistical diagnostics that ensure that the imputed data are representative of measured data and consistent with clinical expectations. The output from AMELIA consists of 5 complete datasets. Logistic regression modeling was then performed on each of the 5 completed imputed datasets and the final adjusted odds ratios determined by calculating the mean from each model [19].

3. Results

During the study period, 730 patients with HCV received at least one dose of PEG-IF and RBV and were included in the analysis. Of these, 351 were infected with either HCV Genotypes 2 or 3. The baseline demographic and laboratory characteristics of the study population are presented in Table 1. The median (IQR) age at treatment was 42 (33 - 48) years. There was a predominance of male subjects (64.7%). A liver biopsy was performed on 54% of participants with the majority of these patients found to have early stage fibrosis. There was an even spread of prescription of the two interferon types across each of the study centres (48% and 54% for PEG-IFα-2a at FH and RPH, respectively [P = 0.30]). Details regarding weight were available for 48% of patients.

Comparisons according to PEG-IFα formulation by univariate analysis are shown in Table 2. The overall SVR rate was superior for PEG-IFα-2b compared to PEG-IFα-2a (77.9% versus 62.0%; odds ratio (OR) for SVR of 2.16 [CI95 1.35 - 3.46; P = 0.0012]) (Figure 1). The only other factor associated with SVR was lower age with the median age of those attaining an SVR being 41 years versus 44 years for non-SVR (OR 0.98 CI95 0.95 - 1.00, P = 0.04). Multivariate analysis confirmed a superior SVR with PEG-IFα-2b (OR 2.19; CI95 1.35 - 3.52). Patient age remained independently associated with SVR (OR 0.97; CI95 0.94 - 0.99). HCV Genotype 2 had a significantly higher SVR rate than Genotype 3 (OR 2.66; CI95 1.28 - 5.97).
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Table 1. Baseline demographics and univariate comparisons between patients receiving different formulations of pegylated interferon alpha. *METAIVIR fibrosis score [17].

<table>
<thead>
<tr>
<th></th>
<th>Number of patients with data available (%)</th>
<th>Median (IQR)</th>
<th>Peg-interferon α2a (N = 179)</th>
<th>Peg-interferon α2b (N = 172)</th>
<th>P-value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Age, years (IQR)</td>
<td>351 (100)</td>
<td>42 (33-48)</td>
<td>41 (34-49)</td>
<td>42.0 (33-47)</td>
<td>0.36</td>
</tr>
<tr>
<td>Sex – Total</td>
<td>351 (100)</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Male (%)</td>
<td>227 (100)</td>
<td>112 (63)</td>
<td>115 (67)</td>
<td></td>
<td>0.42</td>
</tr>
<tr>
<td>Female (%)</td>
<td>124 (100)</td>
<td>67 (54)</td>
<td>57 (46)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Weight (kg), median (IQR)</td>
<td>171 (48)</td>
<td>77 (68 - 87)</td>
<td>74 (67 - 85)</td>
<td>79 (68 - 95)</td>
<td>0.09</td>
</tr>
<tr>
<td>Centre, Fremantle</td>
<td>157 (100)</td>
<td>75 (48)</td>
<td>82 (52)</td>
<td></td>
<td>0.30</td>
</tr>
<tr>
<td>Centre, Royal Perth</td>
<td>194 (100)</td>
<td>104 (54)</td>
<td>90 (46)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Hepatitis C genotype</td>
<td>351 (100)</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Genotype 3</td>
<td>296 (100)</td>
<td>149 (83)</td>
<td>147 (86)</td>
<td></td>
<td>0.40</td>
</tr>
<tr>
<td>Genotype 2</td>
<td>55 (100)</td>
<td>30 (56)</td>
<td>25 (44)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Fibrosis score*, median (IQR)</td>
<td>188 (54)</td>
<td>2 (1 - 3)</td>
<td>2 (1 - 2)</td>
<td></td>
<td>0.71</td>
</tr>
<tr>
<td>Ribavirin dose, mg/kg, median (IQR)</td>
<td>171 (47)</td>
<td>10.8 (9.4 - 11.9)</td>
<td>11.6 (10.0 - 12.7)</td>
<td>0.41</td>
<td></td>
</tr>
</tbody>
</table>

Table 2. Predictors of sustained virological response (SVR) in patients treated with pegylated-interferon and ribavirin for chronic hepatitis C infection with Genotypes 2 and 3. Data are shown as percentages (%) or medians with interquartile ranges (IQR). Odds ratios are given with 95% confidence intervals. *METAIVIR fibrosis score [17].

<table>
<thead>
<tr>
<th></th>
<th>SVR N (%)</th>
<th>No SVR N (%)</th>
<th>P-value</th>
<th>Unadjusted OR (95% CI)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Overall</td>
<td>245 (69.8)</td>
<td>106 (30.2)</td>
<td>0.04</td>
<td>0.98 (0.95 - 1.00)</td>
</tr>
<tr>
<td>Age, years – median (IQR)</td>
<td>41 (32-47)</td>
<td>44 (35-49)</td>
<td>0.12</td>
<td>0.67 (0.41 - 1.11)</td>
</tr>
<tr>
<td>Sex, Male (Yes)</td>
<td>154 (63)</td>
<td>73 (69)</td>
<td>0.36</td>
<td></td>
</tr>
<tr>
<td>Weight, kg (median, IQR)</td>
<td>78 (69-89)</td>
<td>72.5 (65-86)</td>
<td>0.53</td>
<td>0.53 (0.26 - 1.07)</td>
</tr>
<tr>
<td>Centre (FH vs. RPH):</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>FH</td>
<td>114 (72)</td>
<td>43 (28)</td>
<td>0.35</td>
<td>1.27 (0.80 - 2.02)</td>
</tr>
<tr>
<td>RPH</td>
<td>131 (68)</td>
<td>63 (32)</td>
<td></td>
<td></td>
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<tr>
<td>Hepatitis C genotype:</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Genotype 3</td>
<td>201 (67.9)</td>
<td>95 (32.1)</td>
<td>0.08</td>
<td>0.53 (0.26 - 1.07)</td>
</tr>
<tr>
<td>Genotype 2</td>
<td>44 (80.0)</td>
<td>11 (20.0)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Fibrosis score*, median (IQR)</td>
<td>2 (1 - 2)</td>
<td>2 (1 - 3)</td>
<td>0.21</td>
<td></td>
</tr>
<tr>
<td>Pegylated interferon formulation:</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>PEG-IFα-2a</td>
<td>111 (62.0)</td>
<td>68 (38.0)</td>
<td>0.0012</td>
<td>2.16 (1.35 - 3.46)</td>
</tr>
<tr>
<td>PEG-IFα-2b</td>
<td>134 (77.9)</td>
<td>38 (22.1)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Ribavirin dose, mg/kg median (IQR)</td>
<td>10.7 (9.6 - 11.8)</td>
<td>11.1 (9.5 - 12.5)</td>
<td>0.70</td>
<td></td>
</tr>
</tbody>
</table>

Logistic regression on the 5 imputed datasets from AMELIA did not identify any additional independent variables associated with SVR (data not shown).

In our study, patients with HCV Genotype 2 were significantly older (median 48 years versus 40, $P < 0.001$) and fewer were male (53% versus 67%, $P = 0.04$) than patients with Genotype 3. Genotype-specific logistic regression did not identify any variable to be independently associated with SVR for Genotype 2, whilst only PEG-IFα-2b (OR 2.64; CI 1.59 - 4.44) and lower age (OR 3.34; CI 1.59 - 1.42 - 10.1, $P = 0.01$) were independently associated with SVR for Genotype 3.

In analyses for end of treatment HCV RNA PCR negativity (EOT response) PEG-IFα-2b formulation (OR
Two meta-analyses, published simultaneously, have reported superiority of PEG-IFα-2a when compared with PEG-IFα-2b for HCV Genotype 1 infection [16,24]. Because of inconsistencies in study selection, inclusion criteria and randomisation stratification, critics have urged caution in applying these results to HCV treatment in all clinical situations [25].

Such a situation is treatment of HCV Genotypes 2 and 3. In the Cochrane analysis, only 5 studies were selected for inclusion in the Genotype 2/3 sub-group analyses [16]. These studies included those only published in abstract form, [26] studies enrolling HIV co-infected patients [27] and studies of previous non-responders, [28] often with small patient numbers [27,28]. Furthermore, because the reported SVR was particularly high in at least two studies, it is likely that they lacked statistical power to detect differences of limited magnitude [26,29,30].

At least one other study seems not to have been included at all [31]. In this study, no statistically significant difference in SVR was observed between different formulations, but the authors noted lower relapse rates with PEG-IFα-2b and RBV suggesting that cost-effectiveness issues might favour that choice, irrespective of SVR, a proposition also favoured by other commentators [32].

Given the limitations of other studies and their associated meta-analyses, the present study, involving over 350 patients in a “real-world” clinical environment extends the literature on this topic. As in the Genotype 1 IDEAL [23], study we found a similar EOT response with PEG-IFα-2a and 2b, but higher relapse rates within the PEG-IFα-2a group. In HCV Genotype 1 infection, relapse has been postulated to be related to levels of RBV or IF drug exposure during treatment [33,34]. In Australia, weight based on RBV is not licensed for HCV Genotype 2 and 3 infections treated with PEG-IFα-2a. Despite this in our population, there were no significant differences in weight or RBV dosage between the two groups, so RBV dose is unlikely to explain the observed differences in SVR. Therefore, the formulation of PEG-IFα may be relevant. Commentaries on different pharmacokinetics, binding characteristics, and interferon exposures between the two brands broadly suggest that PEG-IFα-2b exhibits greater virological activity [35-37].

The other important finding in the present study is that SVR for HCV Genotype 2 is better than that for Genotype 3. This phenomenon has been recognised elsewhere and questions the validity of continuing to group these HCV genotypes together [30].

Similar to findings in previous studies, we also observed improved SVR in patients treated at younger ages [38]. The discovery of the role of IL28B polymorphisms in response in Genotype 1 infection has occurred since our study was completed [20-22] and IL28B data were...
not available for our analysis. Subsequently, other authors using both brands of either PEG-IF or non PEG-IF together with RBV have found similar IL28B polymorphisms may be important in SVR in infections with other HCV genotypes [38]. However, there is no reason to suspect that there would be any differences in IL28B polymorphisms between our groups treated with either PEG-IF formulation.

This study is limited in that it did not include information about adherence to therapy, a factor important to our outcome [39]. Meta-analyses suggest that PEG-IF is associated with fewer adverse events leading to treatment discontinuation [16]. Although speculative, this could underestimated the magnitude of the beneficial SVR response observed in our patients with HCV Genotypes 2 and 3 treated with PEG-IF.

With the addition of direct acting antivirals (DAAs) to the therapy of HCV [12,13], the importance of a difference between the two forms of PEG-IF might appear irrelevant. But with the high cost of DAAs, increasingly personalised choices based on DAA, viral and host characteristics, and excellent results with the present standard of care in patients with HCV Genotypes 2 and 3, the choice of the optimal PEG-IF might remain important for those who do not need or who cannot afford new DAAs.

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REFERENCES


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http://dx.doi.org/10.1016/j.cgh.2006.10.008

http://dx.doi.org/10.1080/00365520802647400

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http://dx.doi.org/10.1016/j.jhep.2006.03.008

http://dx.doi.org/10.1016/j.jhep.2008.05.009

http://dx.doi.org/10.1016/j.jhep.2010.07.041

http://dx.doi.org/10.1053/gast.2002.35950