Hepatitis C virus related cirrhosis: time to occurrence of hepatocellular carcinoma and death

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Abstract

Background—In patients with hepatitis C virus (HCV) infection and cirrhosis, long term outcome and the incidence of hepatocellular carcinoma (HCC) are still debated.

Design—From January 1987 to January 1997, 416 patients (240 male, median age 57 years) with uncomplicated Child-Pugh A HCV related cirrhosis were followed in two Paris area centres from diagnosis of cirrhosis until death or reference date (1 June 1998). The analysis used a three state disability model generalising the Cox model.

Results—Of the 416 patients, 60 developed HCC with a five year rate of 13.4% (95% confidence interval (CI) 9.0–17.8%) and 83 died (including 34 with HCC), with a five year death rate of 15.3% (95% CI 12.6–18.0%). By multivariable analysis, time to HCC relied on age (hazard ratio (HR) 1.05 per year; p=0.0005), male sex (HR 2.13; p=0.01), oesophageal varices (HR 2.36; p=0.008), decreased platelet count (HR 0.99; p=0.03), and bilirubin level (HR 1.01; p=0.003), while death after HCC was mainly related to tobacco consumption (HR 1.04; p=0.0006). In contrast, death free of HCC was dependent on age (HR 1.04; p=0.01), oesophageal varices (HR 2.75; p=0.001), low platelet count (HR 0.99; p=0.006), and albumin level (HR 0.90; p=0.0001).

Conclusion—The incidence of HCC and mortality should be higher in these patients than previously stated, and prognostic factors for HCC and death are closely related age and symptoms of portal hypertension.

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Keywords: hepatitis C; cirrhosis; hepatocellular carcinoma; survival

Hepatitis C virus (HCV) infection has become a major worldwide health problem because of the potential natural course of the disease to cirrhosis and then hepatocellular carcinoma (HCC).1 14. At the present time, HCV related cirrhosis is the most common indication for orthotopic liver transplantation in Western countries1 although limited data have been reported concerning long term mortality and incidence of HCC in these patients.1 15 16 Most previous retrospective cohort studies had several limitations caused by either the absence of HCV serology with focus on non-A non-B hepatitis1 6 10 or a short and incomplete follow-up. Prospective cohort studies used small samples with no consistent report of causes of death,16 except for a recent multicentre European study that enrolled, over a 10 year period, 384 patients with Child-Pugh A HCV related cirrhosis.17 Fair rates of survival of 91% and 79% at five and 10 years, respectively, were found. However, these estimates may have suffered from selection bias as some centres only included five patients.

Finally, the current understanding of prognostic factors for HCV related cirrhosis progression is based on consideration of only two states: the initial stage (cirrhosis) and an outcome stage (either HCC or death). Several authors were only interested in the occurrence of the outcome, ignoring the time scale and thus censoring.18 Others used time to outcome as the main end point, defining survival models.1 17 20 Almost all studies on prognostic factors have used this two state model as the basis for identifying prognostic factors, with no details of interactions among the prognostic factors and outcome.

The aim of our study was to assess and predict the outcome of HCV related cirrhosis patients based on a cohort of 416 patients from two centres in the Paris area. Two main end points were considered, HCC and death, analysed jointly through the use of a three state disability model (or illness-death model).21

Patients and methods

PATIENTS

Inclusion in the study was considered at the time of diagnosis of HCV related cirrhosis. All consecutive patients with HCV related cirrhosis referred to our two hepatology departments between 1 January 1987 and 31 December 1996 for liver biopsy were included in the study if they met all of the following criteria: (1) positive serum anti-HCV antibodies (second or third generation ELISA test); (2) cirrhosis compatible with HCV origin proved on biopsy either by the transparietal or transvenous route; (3) Child Pugh class A; (4) absence of HCC defined by the absence of focal liver mass on ultrasonography or CT scan, and serum a fetoprotein (AFP) <50 ng/ml; (5) absence of coinfection with other viruses (HIV, HBV) or associated liver diseases; (6) precise evaluation of alcohol consumption; and (6) residence in

Abbreviations used in this paper: HCV, hepatitis C virus; HCC, hepatocellular carcinoma; AFP, a fetoprotein; SMR, standardised mortality ratio; HR, hazard ratio.
France allowing regular follow up. The date of the liver biopsy establishing the diagnosis of cirrhosis defined the starting time in the study.

BASELINE CLINICAL AND BIOLOGICAL DATA
Routine clinical and biological data were collected: age, sex, tobacco consumption, serum albumin (g/l), bilirubin (µmol/l), platelet count (10⁹/l), prothrombin time (%), and AFP level (ng/ml). Daily alcohol consumption was evaluated as < 40 g, 40–80 g, and > 80 g. Oesophageal varices were looked for at endoscopy in 325 patients (table 1).

LIVER HISTOLOGY
All patients underwent transapertural or transvenous liver biopsy depending mainly on platelet count (with a cut off of 100×10⁹/l) and bleeding time. Transvenous biopsies were also performed for protracted prothrombin times or for the purpose of measuring wedged hepatic pressure. Liver biopsy samples were fixed and embedded in paraffin. Knodell fibrosis and activity scores were assessed. Of the 416 biopsies, 278 were independently re-examined by two liver pathologists (CD and CG), without knowledge of the clinical data, to detect the presence of large cell dysplasia. Small liver samples obtained from transvenous liver biopsies were not analysed for the presence of large cell dysplasia. Non-concordant results involved reviewing the slides until consensus was established.

PROSPECTIVE FOLLOW UP AND HCC SCREENING
In accordance with the screening programme, ultrasonographic study of the liver was performed every six months. A diagnosis of HCC was based on histology obtained by ultrasound guided liver biopsy. In patients in whom liver biopsy was not possible, a diagnosis was based on: (a) liver cirrhosis; (b) focal lesion compatible with HCC, demonstrated by two imaging techniques (ultrasound, CT scan, or MRI); and (c) AFP level above 400 ng/ml, portal thrombosis, or growing arterial hypervascular tumour at CT scan and/or angiography and/or MRI. Doubtful cases were not considered as HCC until a precise histological diagnosis was obtained. All patients considered as HCC without histological proof were followed up and had an increase in the size of their tumour.

INTERFERON TREATMENT
Interferon treatment was recorded. Sustained response was defined on biological criteria (normalisation of alanine aminotransferase activity six months after discontinuation of treatment). Because of the length of the study, we did not evaluate the virological response to treatment.

STATISTICAL ANALYSIS
Analysis was based on the reference date of 1 June 1998. The disability model was specified by three transition intensities: the intensity (or incidence) of developing HCC, the incidence of death without HCC, and the incidence of death intensity with HCC (fig 1). Analysis of the disability model is performed one transition at a time, using time since diagnosis of HCV related cirrhosis as the time scale. Patients were at risk of death unrelated to HCC from a diagnosis of cirrhosis until the time to death without HCC, censoring, or development of HCC. They were at risk of death related to HCC from the time to development of HCC until the time to death or censoring. We first estimated time to events using the Kaplan Meier method, and the hazard rate over one year intervals by the actuarial life table method. Standardised mortality ratios (SMR) were computed to compare survival distributions with those of the general French population, matched for age and sex.

Prognostic analyses considered for each transition intensity a full multivariable model with proportional hazards that allowed estimation of the hazard ratio (HR) with 95% confidence intervals (95% CI). The fit of the final model was checked by testing for interaction between the prognostic factors in the model and the underlying time (that is, for proportional intensities) and by testing the log linearity of the continuous predictors. The latter were also introduced as binary covariates after dichotomisation according to median values.
Two sided tests were computed with p values <0.05 defining statistical significance. SAS (SAS Inc, Cary, North Carolina, USA) and S-plus software packages were used for statistical analysis.

**Results**

**BASELINE CHARACTERISTICS**

From 1 January 1989 to 31 December 1996, 651 consecutive patients with Child A HCV cirrhosis were referred to our two departments for liver biopsy. Of these, 215 were excluded because of viral coinfection, the presence of an ultrasound detectable nodule at inclusion, and/or non-virus C related liver disease. Accordingly, 416 patients fulfilled the inclusion criteria. A liver biopsy was performed by the transvenous transjugular route in 58% of patients. Haemodynamic measurements were not available for all patients.

The sample included 240 males and 176 females, median age 57 years. Table 1 summarises the distribution of the baseline patient characteristics at the time of diagnosis of HCV related cirrhosis. The majority of patients (80%) were asymptomatic, the diagnosis of cirrhosis being the result of screening liver biopsy, while 20% were symptomatic: 45 patients (11.7%) had minimal US detectable ascitis and bilirubin was >30 <50 µmol/l in 38 patients (9.7%).

According to the Knodell score, all 416 patients had a fibrosis index of 4 while the activity index was minimal (<4) in 206 (50%) patients, moderate (4–8) in 169 (41%), and severe (8) in 41 (9%). Large cell dysplasia was observed in 44/287 patients (15%).

![Graphs showing estimated time to outcome](image-url)

*Figure 2 Estimated time to outcome (hepatocellular carcinoma (HCC), death after HCC, or death free of HCC) with associated estimated hazard of developing the event over one year time intervals. This refers to the conditional failure rate (that is, the probability of an individual free of an event at the end of the time interval experiencing the event in the next interval). An interesting feature of these data is that the hazard rate for HCC increases over time while the hazard rates for death remain roughly constant over time.*
HR, hazard ratio; 95 CI, 95% confidence interval.

†Grade 2 or >2.

The size of the confirmed HCC tumour was >10 mm in all but two patients. AFP level was >400 ng/ml in 17% of patients (fig 3). At five years the HCC rate was 13.4% (95% confidence interval (CI) 9.0–17.8%) and the death rate was 15.25% (95% CI 12.6–17.9%). Death was not due to liver disease in 19 patients (including suicide in one untreated patient): cancer (ovarian, digestive, brain tumours (n=6); cardiac failure (n=4); severe infection (n=3); brain haemorrhage (n=2); other (n=3). Compared with age and sex matched life tables, mortality for HCV related cirrhosis whatever the cause was increased by about threefold (SMR 2.85; p<0.0001). Analysing only mortality of HCV related cirrhosis (that is, excluding HCC related mortality) yielded an estimated SMR of 1.76 (p= 0.001). Interestingly, the hazard rate of HCC tended to increase with time, when the hazard rates for death remained fairly constant (fig 2).

PROGNOSTIC ANALYSES

The results of the prognostic multivariate analyses are summarised in table 2. The occurrence of HCC was related to increased age (hazard ratio (HR) 3.78 above 57 years; 95% CI 1.77–8.05), male sex (HR 2.13; 95% CI 1.18–3.85), presence of oesophageal varices (HR 2.36; 95% CI 1.26–4.45), decreased platelet count (HR 1.69 below 100×10⁹/l; 95% CI 0.99–3.17), and bilirubin level (HR 1.14 above 15 mg/dl; 95% CI 0.61–2.13), all assessed at the time of diagnosis of cirrhosis. Alcohol consumption was not significantly associated with the occurrence of HCC.

To explain the heterogeneity in survival times without HCC, the most relevant variables were close to those related to the occurrence of HCC, suggesting a homogeneous baseline high risk group based on age (HR 1.31; 95% CI 0.63–2.72), presence of oesophageal varices (HR 2.75; 95% CI 1.48–5.10), decreased albumin level (HR 3.70 below 41 g/l; 95% CI 0.99–3.17), and bilirubin level (HR 1.14 above 15 mg/dl; 95% CI 0.61–2.13), all assessed at the time of diagnosis of cirrhosis. Alcohol consumption was not significantly associated with the occurrence of HCC.

Table 2 Prognostic analyses according to outcome: hepatocellular carcinoma (HCC), death after hepatocellular carcinoma, and death unrelated to hepatocellular carcinoma

<table>
<thead>
<tr>
<th>Event</th>
<th>HCC (60 events)</th>
<th>Death after HCC (34 events)</th>
<th>Death unrelated to HCC (48 events)</th>
</tr>
</thead>
<tbody>
<tr>
<td>HR (95 CI)</td>
<td>p Value</td>
<td>HR (95 CI)</td>
<td>p Value</td>
</tr>
<tr>
<td>Age</td>
<td>1.05 (1.02–1.08)</td>
<td>0.0005</td>
<td>1.03 (0.98–1.08)</td>
</tr>
<tr>
<td>Female</td>
<td>0.47 (0.26–0.85)</td>
<td>0.013</td>
<td>0.96 (0.38–2.40)</td>
</tr>
<tr>
<td>Oesophageal varices*</td>
<td>2.36 (1.25–4.45)</td>
<td>0.008</td>
<td>0.63 (0.26–1.53)</td>
</tr>
<tr>
<td>Platelets</td>
<td>0.99 (0.98–0.99)</td>
<td>0.029</td>
<td>1.002 (0.99-1.01)</td>
</tr>
<tr>
<td>Bilirubin</td>
<td>1.01 (1.005–1.02)</td>
<td>0.003</td>
<td>1.000 (0.99-1.01)</td>
</tr>
<tr>
<td>Alcohol &gt;40 g</td>
<td>0.56 (0.26–1.23)</td>
<td>0.15</td>
<td>0.45 (0.14–1.46)</td>
</tr>
<tr>
<td>Tobacco</td>
<td>1.01 (0.99–1.02)</td>
<td>0.42</td>
<td>1.04 (1.02-1.06)</td>
</tr>
<tr>
<td>Albumin</td>
<td>1.03 (0.97–1.09)</td>
<td>0.36</td>
<td>0.93 (0.86–1.01)</td>
</tr>
</tbody>
</table>

*Grade 2 or >2.

HR, hazard ratio; 95 CI, 95% confidence interval.
for example a five year death rate of 15% compared with 9%) although all patients had a similar inclusion criterion (that is, Child-Pugh class A). This could be explained by the extensive use of transcutaneous liver biopsy in our study, allowing us to include patients with more severe liver disease. Moreover, inclusion in our study, contrary to the Eurohep study,17 of 10% of patients with a large (>80 g/day) alcohol consumption (provided that histological examination showed a pattern of HCV related disease) may have contributed to our overall high death rate. However, most of these patients stopped or reduced their drinking habits during follow up (evaluation based on questioning by the physician) and alcohol consumption at the initiation of follow up was not found to be a prognostic factor in our study.

With regard to the occurrence of HCC, we first showed that the instantaneous risk of HCC increased with each passing year, as shown by Mazella and colleagues.20 This could be explained in part by exclusion of cirrhotic patients with a non-identified, even small liver nodule at entry, as well as patients with AFP >50 ng/ml. Moreover, it could be related to increasing risk factors such as more advanced age and poorer liver function. Overall, the incidence of HCC was high, with an estimate of 13.4% at five years compared with previous reports of 7%17 or 5%.20 Such a discrepancy could be due to improvements in the accuracy of diagnosis and closer follow up for early detection of HCC. Nevertheless, our estimate is in the range of previous reports using a similar mean time and periodicity of follow up, and showing an incidence of 11% at four years16 and 13.5% at five years.21

In understanding the variability of the occurrence of HCC in HCV related cirrhosis patients, we confirmed the importance of sex, age, and signs of portal hypertension, such as oesophageal varices and low platelet count, as previously published.15 20 The prognostic value of haemodynamic measures and the correlation with platelet count require further study. We did not find any relationship between the occurrence of HCC and previously reported prognostic factors such as alcohol consumption and cigarette smoking,14 20 but, as discussed below, surprisingly cigarette smoking was a predictive factor of death in patients with HCC.

Levels of AFP at inclusion did not add any prognostic information on HCC occurrence in contrast with a previous report.14 This relates to AFP levels obtained at the start point of the study and below 50 ng/ml. It does not rule out a prognostic value of serum levels of AFP above 50 ng/ml or of a transient elevation during the follow up, as only the initial value was taken into account. More interestingly, in our study, two other controversial predictive factors were not found to be associated with the occurrence of HCC, namely the presence of large cell dysplasia and treatment with interferon. The liver biopsies at inclusion of 278/416 patients were reviewed independently by two pathologists (CD and CG), blinded to the outcome, so that there would be no bias in the results. The predictive value of large cell dysplasia may depend on the cause of cirrhosis as previous findings were based on patients with heterogeneous causes (alcohol, hepatitis B virus infection, hepatitis C).21 22 and not specifically on HCV. In fact, large cell dysplasia is probably more informative of outcome in HBV infection22 or even alcoholic cirrhosis which largely predominates in some studies.27

The role of interferon in the prevention of HCC has been widely discussed16 18 20 21 28 29 and our results only underline the need for a large prospective randomised trial. No conclusion can be drawn from the absence of significant results in our study. Finally, virological factors such as the presence of HBV markers18 of HCV replication detected by PCR and infection with genotype 119 20 that have been described as predictive of HCC could not be assessed in our patients. HBV markers have been previously studied in patients from one centre (Bondy) and did not seem to add any prognostic value to the occurrence of HCC.

In the small group of 34 patients with HCC, the occurrence of death was independently related to smoking habits. The importance of tobacco in hepatocarcinogenesis has been stressed by previous studies. Tobacco smoking, when associated with alcohol consumption, could induce metabolites that have a carcinogetic effect, and presumably for explaining our data could lead to more malignant forms of HCC.

In our series, the predictive factors of survival in patients free of HCC were similar to those found in the Eurohep study,17 namely age, low platelet count, and high bilirubin level at initial examination. We also found that decreased albumin level and the presence of oesophageal varices were predictive of poor survival. The various treatments of portal hypertension were not recorded and their influence on mortality was therefore not studied. In fact, these factors together reflect the importance of age on the severity of liver disease in predicting outcome unrelated to HCC.

It should be noted that the data requirements for a multi-state model are quite strict. Not only are the dates of entry and death or censoring needed, but also, in principle, the exact transition times between the transient states must be known for each individual. In our data set, patients with HCV related cirrhosis underwent a screening programme that allowed for similar detection of HCC. Nevertheless, it should be kept in mind that similar problems may arise with any clinical data when the detection of non-fatal complications depends strongly on the frequency with which patients are followed. Finally, the value of any set of prognostic factors depends on reproducibility in a new patient sample. Our results agree with previously published data when the detection of non-fatal complications depends strongly on the frequency with which patients are followed. Finally, the value of any set of prognostic factors depends on reproducibility in a new patient sample. Our results agree with previously published data when the detection of non-fatal complications depends strongly on the frequency with which patients are followed. Finally, the value of any set of prognostic factors depends on reproducibility in a new patient sample. Our results agree with previously published data when the detection of non-fatal complications depends strongly on the frequency with which patients are followed. Finally, the value of any set of prognostic factors depends on reproducibility in a new patient sample.
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