Carvedilol for portal hypertension in cirrhosis: systematic review with meta-analysis

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ABSTRACT

Objective: To assess the clinical and haemodynamic effects of carvedilol for patients with cirrhosis and portal hypertension.

Design: A systematic review and meta-analysis.

Data sources: We searched PubMed, Cochrane library databases, EMBASE and the Science Citation Index Expanded through December 2015. Only randomised controlled trials (RCTs) were included.

Outcome measure: We calculated clinical outcomes (all-cause mortality, bleeding-related mortality, upper gastrointestinal bleeding) as well as haemodynamic outcomes (hepatic venous pressure gradient (HVPG) reduction, haemodynamic response rate, post-treatment arterial blood pressure (mean arterial pressure; MAP) and adverse events).

Results: 12 RCTs were included. In 7 trials that looked at haemodynamic outcomes comparison carvedilol versus propranolol, showing that carvedilol was associated with a greater reduction (%) of HVPG within 6 months (mean difference −8.49, 95% CI −12.36 to −4.63) without a greater reduction in MAP than propranolol. In 3 trials investigating differences in clinical outcomes between carvedilol versus endoscopic variceal band ligation (EV), no significant differences in mortality or variceal bleeding were demonstrated. 1 trial compared clinical outcomes between carvedilol versus naldol plus isosorbide-5-mononitrate (ISMN), and showed that no significant difference in mortality or bleeding had been found. 1 trial comparing carvedilol versus nebivolol showed a greater reduction in HVPG after 14 days follow-up in the carvedilol group.

Conclusions: Carvedilol may be more effective in decreasing HVPG than propranolol or nebivolol and it may be as effective as EV or naldol plus ISMN in preventing variceal bleeding. However, the overall quality of evidence is low. Further large-scale randomised studies are required before we can make firm conclusions.

Trial registration number: CRD42015020542.

INTRODUCTION

Acute variceal bleeding is the most feared and devastating complication of cirrhosis. Six-week mortality is ~10–20%, ranging from 0% for patients with Child-Pugh class A to ~30% for patients with Child-Pugh class C. The 1-year rate of recurrent variceal bleeding is ~60%. Although some risk factors are known, such as red wale marks on the varices, large variceal size and advanced pre-existing liver disease, measurement of portal pressure with hepatic venous pressure gradient (HVPG) is the best method for estimating the risk of bleeding varices. HVPG >10 mm Hg is the strongest predictor of the development of varices; HVPG >12 mm Hg is associated with a high risk of variceal bleeding. For patients with acute variceal bleeding, HVPG >20 mm Hg has been associated with high mortality. Bleeding is less likely if HVPG decreases to <12 mm Hg or decreases by 20% from the baseline figure.

Various types of drugs have been used to decrease HVPG. The European Association for the Study of the Liver (EASL) guidelines recommended using non-selective β-blockers (NSBBs), such as propranolol or nadolol, with or without isosorbide-5-mononitrate (ISMN) to prevent variceal bleeding. Reduction of HVPG is achieved by decreasing the cardiac output (β1-blockade) and constricting the splanchic vessels (β2-blockade). Only ~40% of treated patients reach therapeutic levels, and the risk of variceal bleeding remains high for haemodynamic non-responders. Carvedilol, which blocks both α and β receptors, was reported to have better results than NSBBs by additionally decreasing intrahepatic
rate was de-

Methods

The protocol has been registered on the PROSPERO registry under registration number CRD42015020542.

Types of studies

We included only randomised controlled trials (RCTs) comparing carvedilol versus propranolol or other interventions for participants with cirrhosis and portal hypertension, irrespective of the publication status, language or blinding. Abstracts were included.

Types of participants

The participants were patients with cirrhosis and portal hypertension, regardless of the aetiology of cirrhosis or the severity of cirrhosis, who were >18 years old.

Types of interventions

We included trials comparing carvedilol versus propranolol or any other intervention. Any co-interventions were allowed if they were used in both arms of the trial.

Treatment outcomes

The primary outcomes were: (1) all-cause mortality, (2) bleeding-related mortality and (3) upper gastrointestinal bleeding. The secondary outcomes were: (1) HVPG reduction, assessed as a percentage; (2) haemodynamic response rate; (3) post-treatment MAP and (4) adverse events.

For all-cause mortality, upper gastrointestinal bleeding and bleeding-related mortality, trials with follow-ups longer than 7 days were included. Percentage of HVPG reduction, haemodynamic response rate and post-treatment MAP were assessed separately within 24 h (acute term), at 24 h to 6 months (short term) and at >6 months (long term). The haemodynamic response rate was defined as a rate of HVPG reduction ≥20% of the baseline value or ≤12 mm Hg.

Search methods for identification of studies

We searched the PubMed, Cochrane Central Register of Controlled Trials, EMBASE and Science Citation Index Expanded databases for published articles. The search strategies with the expected time spans are displayed in onlinesupplementary appendix1.

We also searched clinical trials databases (ClinicalTrials.gov; WHO International Clinical Trial Registry Platform; International Standard Randomized Controlled Trial Number (ISRTN) registry) for planned and ongoing trials. We searched the Conference Proceedings Citation Index (CPCI-S/CPCI-SSH), BIOSIS previews and Derwent Innovation Index (DII) databases for conference proceedings and innovations.

We reviewed the reference lists of the retrieved articles for potentially relevant studies; we attempted to contact the corresponding authors of relevant studies to request information on unpublished articles. We also sent letters to the authors of abstracts and articles with incomplete data to obtain additional information.

Data collection and analysis

Three authors (TL, XC and JL) selected studies for inclusion following the PRISMA process. Disagreements were resolved by discussion with other authors (PS and AB). The reasons for exclusion were recorded. Two authors (WK and WX) extracted data from the included studies; disagreements were discussed with another author (YH).

Assessment of risk of bias in included studies

The risk of bias in the included studies was assessed according to the recommendations of the Cochrane Handbook for Systematic Reviews of Interventions.14 Two authors (WK and YH) conducted the assessment of the risk of bias. Disagreements were discussed with other authors (QZ and AB). We classified trials with a low risk of bias if none of the domains were associated with an unclear or high risk of bias; otherwise, an unclear (at least one domain was assessed as having unclear risk without any high-risk domains) or high risk of bias was classified.

Statistical methods

We used relative ratios (RRs) with 95% CIs to calculate dichotomous data and the mean differences (MDs) with 95% CIs to calculate continuous data. We used HRs with 95% CIs as relevant effect measures for mortality and upper gastrointestinal bleeding. We estimated HRs from the log-rank $\chi^2$ statistic, log-rank p values, given numbers of events or Kaplan-Meier curves, using methods presented by Tierney et al.15 In one three-arm study, only two arms were used.16

We performed statistical analysis following the guidelines of the Cochrane Handbook for Systematic Reviews of Interventions and using Review Manager software (RevMan, V.5.3, Copenhagen, Denmark: The Nordic Cochrane Centre, the Cochrane Collaboration, 2014).14 A random-effects model was chosen a priori for all of the analyses, and then the fixed-effects model was performed as a sensitivity test. If the two models yielded the same results, no significant heterogeneity was considered; if the 95% CI for the average intervention effect was wider in the random-effects model, the heterogeneity between studies was considered. We also calculated the $\chi^2$ and $I^2$ statistics. The p<0.10 or $I^2\geq 50%$ was
considered to indicate substantial heterogeneity. If significant heterogeneity was found, potential reasons for heterogeneity were explored.

For missing data, analyses were performed using the intention-to-treat (ITT) principle (best-case/worst-case scenario for dichotomous outcomes: mortality, upper gastrointestinal bleeding, haemodynamic response rate), as well as per protocol principle (for all of the outcomes).

**Subgroup analysis**

We performed subgroup analysis to identify the impact of type of controlled group (trials using ISMN in addition to NSBBs, compared with trials using NSBBs alone).

**Sensitivity analysis**

We excluded the trials published as abstracts and trials that used ISMN in addition to NSBBs to perform sensitive analysis. We did not exclude trials with a high risk of bias because only a few studies with a low risk of bias were included in the meta-analysis.

**Summary of findings tables**

We used ‘summary of findings’ tables to present our assessment of the body of evidence associated with some outcomes, using GRADEPro software (ims.cochrane.org/revman/other-resources/gradepro). The quality of a body of evidence considers five factors regarding the limitations of the studies: risk of bias, inconsistency, indirectness, imprecision and publication bias.

**RESULTS**

**Results of the search**

From 172 identified publications, 67 duplicates were removed. From the remaining 105 publications, 67 were removed due to non-randomised designs or irrelevance regarding our topic. We assessed the full-text versions of the remaining 38 publications. Five publications were excluded: two were published as Master’s theses referring to one trial, which reported largely different results, and the methods of randomisation were questionable;17,18 one was a non-randomised trial;19 one was irrelevant to our topic;20 and one included non-cirrhotic participants.21 Finally, of the remaining 33 publications, 25 referring to 12 trials were included in the quantitative synthesis;10,16,22-31 5 referring to 2 trials were awaiting classification,32,33 and 5 referring to 4 trials were ongoing projects (figure 1).34-37

**Description of the individual comparisons in the trials**

The characteristics of the included studies are shown in table 1 and online supplementary appendices 2 and 3. The summary of findings table is shown in online supplementary appendix 4.

Twelve trials were associated with a low (one), unclear (seven) or high (four) risk of bias. The details can be seen in the ‘risk of bias graph’ (figure 2A) and ‘risk of bias summary’ (figure 2B).

**Carvedilol versus propranolol**

**Description of included studies**

Seven trials (one abstract) with 379 participants compared carvedilol versus propranolol;10,16,22,23,27,30,31 six trials were two-arm (carvedilol vs propranolol) studies, and one trial was a three-arm study (carvedilol vs propranolol vs placebo).16 One of the studies used ISMN in addition to propranolol in the controlled group.22 Haemodynamic outcomes had been reported in each trial, and treatment effects were assessed within 24 h (acute term)10,16,22,23 and within 6 months (24 h to 6 months; short term).10,23,27,30,31 No trials reported long-term outcomes.

Six trials reported the sex of the participants: 199 were male, and 79 were female.10,16,22,23,27,31 The mean age of the participants ranged from 42 to 61 years. Most...
trials mainly included participants with Child-Pugh class A cirrhosis; the most common causes of cirrhosis were alcohol abuse (three trials), hepatitis B virus infection (one trial) and hepatitis C virus infection (one trial). Two trials did not report the aetiologies of cirrhosis.

The characteristics of included studies and included participants as well as the administration of each drug could be seen in Table 1 and online supplementary appendices 2 and 3.

### Effects of interventions

One trial recruited 47 participants with a follow-up about 90 days reported mortality (RR 2.88, 95% CI 0.12 to 67.29), bleeding-related mortality (RR 2.88, 95% CI 0.12 to 67.29) and upper gastrointestinal bleeding (RR 0.96, 95% CI 0.06 to 14.43), but no significant difference had been found for these outcomes. The risk of bias was high because of the incomplete outcome data.

### Percentage (%) of HVPG reduction

Percentage (%) of HVPG reduction within 24 h: Four trials with an unclear (2) or a high (2) risk of bias reported acute-term HVPG reductions after drug administration. No significant heterogeneity was found; the fixed-effects model showed that carvedilol was associated with a greater reduction (%) in HVPG (MD −8.66, 95% CI −12.66 to −4.65) (figure 3A). The result

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**Table 1 Characteristics of included studies**

<table>
<thead>
<tr>
<th>Author year</th>
<th>Group</th>
<th>N of Pati</th>
<th>Administration of intervention</th>
<th>Time of outcome assessment</th>
<th>Drop Carv/Cont ITT</th>
</tr>
</thead>
<tbody>
<tr>
<td>Banares 1999</td>
<td>Carv/Prop</td>
<td>14/14</td>
<td>Carv: 25 mg four times a day orally Prop: 0.15 mg/kg intravenously followed by a continuous infusion of 0.2 mg/kg</td>
<td>60 min</td>
<td>0/0 No</td>
</tr>
<tr>
<td>Banares 2002</td>
<td>Carv/Prop</td>
<td>26/25</td>
<td>Carv: 31±4 mg orally Prop: 73±10 mg orally</td>
<td>11.1±4.1 weeks</td>
<td>2/3 No</td>
</tr>
<tr>
<td>De 2002</td>
<td>Carv/Prop</td>
<td>18/18</td>
<td>Carv: start at 25 mg four times a day orally followed by 6.25 mg twice daily Prop: start at 80 mg four times a day orally followed by 40 mg twice daily</td>
<td>90 min; 7 days</td>
<td>I/2 No</td>
</tr>
<tr>
<td>Hobolth 2012</td>
<td>Carv/Prop</td>
<td>24/23</td>
<td>Carv: start at 3.125 mg twice daily, followed by 14±7 mg/day orally Prop: start at 40 mg twice daily, orally followed by 122±64 mg/day</td>
<td>90 min; 92.7±13.6 days</td>
<td>3/6 No</td>
</tr>
<tr>
<td>Lin 2004</td>
<td>Carv/Prop + ISMN</td>
<td>11/11</td>
<td>Carv: 25 mg, four times a day, orally Prop: 40 mg plus ISMN (20 mg), four times a day, orally</td>
<td>90 min</td>
<td>0/0 No</td>
</tr>
<tr>
<td>Lo 2012</td>
<td>Carv/Nado + ISMN</td>
<td>61/60</td>
<td>Carv: 10.4±2.2 mg/day orally Nado: 45±13 mg±various mg ISMNs/day orally</td>
<td>30 months</td>
<td>5/6 Yes</td>
</tr>
<tr>
<td>Mo 2014</td>
<td>Carv/Prop</td>
<td>48/48</td>
<td>Carv: started at 12.5 mg, four times a day orally then adjusted according to the BP, HR Prop: started at 10 mg, three times a day, orally then adjusted according to the BP, HR</td>
<td>7 days</td>
<td>0/0 No</td>
</tr>
<tr>
<td>Shah 2014</td>
<td>Carv/EVL</td>
<td>82/86</td>
<td>Carv: 6.25 mg four times a day for 1 week, 6.25 mg twice daily thereafter orally E VL: underwent EVL within 48 h of randomisation, repeated every 3 weeks</td>
<td>13 months</td>
<td>2/0 Yes</td>
</tr>
<tr>
<td>Silkauskaite 2013</td>
<td>Carv/Nebi</td>
<td>10/10</td>
<td>Carv: 25 mg four times a day orally Nebi: 5 mg four times aday orally</td>
<td>60 min; 14 days</td>
<td>1/2 No</td>
</tr>
<tr>
<td>Sohn 2013</td>
<td>Carv/Prop</td>
<td>50/49</td>
<td>Carv: 11.6±2.2 mg/day orally Prop: 153.5±100.2 mg/day orally</td>
<td>6 weeks</td>
<td>8/11 Yes</td>
</tr>
<tr>
<td>Stanley 2014</td>
<td>Carv/EVL</td>
<td>33/31</td>
<td>Carv: 6.25 mg four times a day for the first week, 12.5 mg/day thereafter orally E VL: underwent EVL within 1 week, then repeated every 2 weeks</td>
<td>26.4 months</td>
<td>14/11 Yes</td>
</tr>
<tr>
<td>Tripathi 2009</td>
<td>Carv/EVL</td>
<td>77/75</td>
<td>Carv: started at 6.25 mg four times a day for the first week; 12.5 mg thereafter orally E VL: underwent EVL every 2 weeks</td>
<td>26 months</td>
<td>25/23 Yes</td>
</tr>
</tbody>
</table>

BP, blood pressure; Carv, carvedilol; Cont, control; EVL, endoscopic variceal band ligation; ISMN, isosorbide-5-mononitrate; ITT, intention-to-treat analysis; Nado, nadolol; Nebi, nebivolol; Pati, patients; Prop, propranolol.
of a sensitive analysis that excluded the trial using propranolol plus ISMN as the control remained consistent with the pooled analysis (see online supplementary appendix 5).

Percentage (%) of HVPG reduction at 24 h to 6 months: Five trials with an unclear (3) or a high (2) risk of bias reported the short-term percentage (%) of HVPG reduction after drug administration.\textsuperscript{10} \textsuperscript{23} \textsuperscript{27} \textsuperscript{30} \textsuperscript{31} Four were reported as full texts, while the other was presented as an abstract.\textsuperscript{30} No significant heterogeneity was found. The fixed-effects model showed that carvedilol was associated with a greater reduction (\%) in HVPG (MD $-8.49$, 95\% CI $-12.36$ to $-4.63$; \textit{figure} 3A). The results of sensitivity analysis that excluded the trial reported as an abstract remained consistent with the pooled analysis (see online supplementary appendix 5).

**Haemodynamic response rate**

\textit{Haemodynamic response rate within 24 h:} Three trials with an unclear (2) and a high (1) risk of bias reported the acute-term haemodynamic response rate after drug administration.\textsuperscript{10} \textsuperscript{16} \textsuperscript{23} It was 32 of 51 in the carvedilol group versus 18 of 48 in the propranolol group. The $\chi^2$ result was 3.35, $I^2$ was 40\%. The fixed-effects model found a higher haemodynamic response rate in the carvedilol group (RR 1.67, 95\% CI 1.09 to 2.54; \textit{figure} 3B), while the random-effects model found no significant difference between the studies (RR 1.59, 95\% CI 0.89 to 2.84). Best-case and worst-case scenario analyses were consistent with the per protocol analysis (see online supplementary appendix 5).

Haemodynamic response rate at 24 h to 6 months: Five trials with an unclear (5) or a high risk of bias (2) reported a short-term haemodynamic response rate.\textsuperscript{10} \textsuperscript{23} \textsuperscript{27} \textsuperscript{30} \textsuperscript{31} It was 87 of 160 in the carvedilol group versus 58 of 152 in the propranolol group. No significant heterogeneity was found. The fixed-effects model found that the carvedilol group had better results (RR 1.42, 95\% CI 1.11 to 1.82; \textit{figure} 3B). The results of the ITT analysis, as well as the sensitivity analysis that excluded the trial reported as an abstract, remained consistent with the pooled analysis (see online supplementary appendix 5).

**Post-treatment MAP**

\textit{Post-treatment MAP within 24 h:} Three trials with an unclear (2) or a high (1) risk of bias reported acute-term post-treatment MAP.\textsuperscript{16} \textsuperscript{22} \textsuperscript{23} The $\chi^2$ result was 2.93, and $I^2$ was 32\%. Both the random-effects and fixed-effects models found no statistically significant difference between groups (fixed effects: MD $-4.96$, 95\%
CI −10.76 to 0.84; figure 3C). The results of the sensitive analysis, which excluded the trial using propranolol plus ISMN as a control, remained consistent with the pooled analysis (see online supplementary appendix 5).

Figure 3 Carvedilol versus propranolol. Percentage of hepatic venous pressure (HVPG) reduction (1.1.1 outcome assessed within 24 h; 1.1.2 outcome assessed 24 h–6 months; A); haemodynamic response rate (1.2.1 outcome assessed within 24 h; 1.2.2 outcome assessed 24 h–6 months; B); post-treatment MAP (1.3.1 outcome assessed within 24 h; 1.3.2 outcome assessed 24 h–6 months; C).

Post-treatment MAP at 24 h to 6 months: Four trials with an unclear (2) or a high (2) risk of bias reported the short-term post-treatment MAP. No significant heterogeneity was found. The fixed-effects model found
Carvedilol versus other drugs

Carvedilol versus nadolol

One trial with 121 participants reported on carvedilol versus nadolol plus ISMN.26 The trial only focused on clinical outcomes (mortality, upper gastrointestinal bleeding, etc). Treatment effects were assessed with a follow-up of ∼30 months (table 1 and online supplementary appendixes 2 and 3). There were no statistically significant differences in all-cause mortality (HR 1.07, 95% CI 0.45 to 2.55; RR 0.87, 95% CI 0.48 to 1.57), bleeding-related mortality (RR 0.66, 95% CI 0.11 to 3.79) or upper gastrointestinal bleeding (HR 1.28, 95% CI 0.76 to 2.17; RR 0.98, 95% CI 0.74 to 1.31). However, the carvedilol group had fewer adverse events (5/61 vs 23/61, RR 0.21, 95% CI 0.09 to 0.53).

Carvedilol versus propranolol

This review showed that carvedilol is more effective than propranolol in decreasing HVPG acutely and over short-term follow-up. In addition, the short-term haemodynamic response rates were greater in the carvedilol group. Although no long-term outcomes were reported, one might infer that carvedilol would be effective...
beyond 6 months because the number of haemodynamic responders in the acute setting was almost identical to the outcomes at 6 months. One trial reported mortality and upper gastrointestinal bleeding. This trial was designed to evaluate haemodynamic effects and it is difficult to compare carvedilol with propranolol on mortality with a follow-up within 90 days.10 Recently, a non-randomised study including 104 participants with a follow-up of 2 years had assessed the efficacy of carvedilol for propranolol non-responders.11 It was reported that a significant proportion of propranolol non-responders could achieve haemodynamic responses to carvedilol treatment. In addition, the variceal bleeding rate, hepatic decompensation rate and mortality rate were significantly decreased in the haemodynamic response group. This study indicated that carvedilol might be better than propranolol in decreasing the HVPG and improving the survival of patients with cirrhosis.

Systemic hypotension is the main cause of drug discontinuance among patients using carvedilol. Although some participants developed severe systemic hypotension and withdrew from the trials,25 27 our study showed no significant differences between groups in post-treatment MAP for over acute-term and short-term follow-ups. The haemodynamic effect of carvedilol is dose dependent; an increase in the carvedilol dose from 6.25–12.5 to 25–50 mg/day significantly decreased MAP and HR further without an additional effect on HVPG.25 41 Thus, it is advised that carvedilol be started at a low dose (6.25 mg/day); if tolerated, the dose could be increased stepwise up to 12.5 mg/day. It must be noticed that carvedilol or NSBBs can increase the mortality of patients with decompensated cirrhosis (cirrhosis with refractory ascites or spontaneous bacterial peritonitis).12 13 These drugs can aggravate the disordered systemic circulation and induce acute kidney injury or other life-threatening complications under these circumstances.42 43

Carvedilol versus EVL

EVL is recommended for the primary and secondary prevention of variceal bleeding. It was reported that band ligation was better than NSBBs in preventing variceal bleeding, but it could not improve overall survival.44 In this study, we found no significant differences in overall mortality, bleeding-related mortality or upper gastrointestinal bleeding between groups for the primary and secondary prophylaxis. Carvedilol seems to have the same efficacy in prevention of variceal bleeding as EVL. However, only three studies were analysed, and the quality of evidence is low. More studies are needed to make firm conclusions.

Complications from carvedilol are often mild and subside after dose reduction or drug discontinuation. In contrast, the complications of EVL often require hospitalisation and can be lethal. It may be appropriate to restrict EVL to carvedilol non-responders or to patients who have contraindications to carvedilol. Furthermore, EVL requires frequent follow-up endoscopies because recurrence of varices requiring retreatment occurs in more than 50% of cases during the first year,45 significantly increasing the burden of patients physically and economically.
REFERENCES


