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Editorial Board Member of World Journal of Hepatology, Salvatore Sutti, MSc, PhD, Assistant Professor, Department of Health Sciences, University of East Piedmont, Novara 28100, Italy. salvatore.sutti@med.uniupo.it

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META-ANALYSIS

Effects and safety of natriuretic peptides as treatment of cirrhotic ascites: A systematic review and meta-analysis

Rasmus Hvidbjerg Gantzel, Mikkel Breinholt Kjær, Peter Jepsen, Niels Kristian Aagaard, Hugh Watson, Lise Lotte Gluud, Henning Grønbæk

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BACKGROUND
Natriuretic peptides are involved in the cascade of pathophysiological events occurring in liver cirrhosis, counterbalancing vasoconstriction and anti-natriuretic factors. The effects of natriuretic peptides as treatment of cirrhotic ascites have been investigated only in small studies, and definitive results are lacking.

AIM
To examine the effects and safety of natriuretic peptides in cirrhosis patients with ascites.

METHODS
We searched MEDLINE, Web of Science, Scopus, Cochrane Library and Embase for all available studies applying intravenous administration of any natriuretic peptide to patients suffering from cirrhotic ascites. Inclusion was not limited by treatment duration or dose, or by follow-up duration. Both randomised controlled trials and non-randomised studies were eligible for inclusion. The primary outcome was change in renal sodium excretion. Secondary outcomes included safety measures and changes in renal water excretion, plasma aldosterone concentration, and plasma renin activity.

RESULTS
Twenty-two studies were included. Atrial natriuretic peptide (ANP) was the only intensively studied treatment. Sodium excretion increased in response to
continuous ANP infusion and was more pronounced when infusion rates of > 30 ng/kg/min were administered compared with ≤ 30 ng/kg/min (P < 0.01). Moreover, natriuresis was significantly higher in study subgroups with mild/moderate ascites compared with moderate/severe and refractory ascites (P < 0.01). ANP infusions increased renal water excretion, although without reaching a statistically significant dose-response gradient. Plasma aldosterone concentration and plasma renin activity were significantly lower at baseline in study subgroups achieving a negative sodium balance in response to an ANP administration compared with treatment non-responders (P < 0.01). Blood pressure decreases occurred less frequently when ANP doses ≤ 30 ng/kg/min were applied. The quality of evidence for a natriuretic response to ANP was low, mainly due to small sample sizes and considerable between-study heterogeneity. Data were sparse for the other natriuretic peptides; B-type natriuretic peptide and urodilatin.

CONCLUSION
Intravenous ANP infusions increase sodium excretion in patients with cirrhotic ascites. Continuous infusion rates > 30 ng/kg/min are the most effective. However, safety increases with infusion rates ≤ 30 ng/kg/min.

Key Words: Atrial natriuretic peptide; B-type natriuretic peptide; Urodilatin; Cirrhosis; Ascites; Refractory ascites

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Core Tip: Pharmacotherapies for cirrhotic ascites have remained largely unchanged for the past four decades. 5%-10% of cirrhosis patients with ascites become refractory to available treatments, and the majority of these patients require frequent large-volume paracenteses. This justifies a continued search for new and improved treatments for ascites. This is the first systematic review and meta-analysis to investigate the effects and safety of natriuretic peptides as treatment of cirrhotic ascites. We demonstrate a significant natriuretic effect of intravenously administered atrial natriuretic peptide and summarise the safety findings.

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INTRODUCTION
The cirrhosis-derived health-care burden is growing globally[1]. Patients suffering from liver cirrhosis may develop portal hypertension with substantial neuro-hormonal impacts. Portal hypertension directly induces vasodilation of the splanchnic vascular system with blood pooling in the splanchnic bed. This reduces the central blood volume, leading to arterial hypotension. When these pathophysiological regulatory events become manifest, vasoconstrictor and anti-natriuretic factors are activated, ultimately resulting in retention of sodium and water[2,3]. Ascites formation is the most common manifestation of decompensation due to cirrhosis with portal hypertension. After the appearance of ascites, patients experience a poor prognosis, frequent hospital contacts, and an impaired quality of life [3,4]. In general, dietary sodium restriction, anti-aldosterone agents, and loop-diuretics are effective treatments to manage ascites[4,5]. However, 5%-10% progress to refractory ascites[6]. Invasive treatments, such as liver transplantation, transjugular intrahepatic portosystemic shunt, or implantation of an automated low-fluid ascites pump (Alfapump®), are restricted to a minor proportion of patients with refractory ascites, while the majority require frequent hospitalizations for large-volume paracentesis[3,4]. New pharmacotherapies to counteract the mechanisms involved in ascites development, as well as to treat established ascites and reduce the need for paracenteses in refractory ascites, are highly warranted.

Atrial natriuretic peptide (ANP) is the most extensively studied member of the natriuretic peptide family, which also comprises B-type natriuretic peptide (BNP) and two active forms of C-type natriuretic peptide (CNP), as well as the renal paracrine peptide urodilatin[7,8]. Urodilatin is identical to ANP except for an N-terminal extension of 4-amino acids, which makes it less vulnerable to degradation by the renal neutral endopeptidase[8,9]. Therefore, the renal effects of urodilatin are stronger than those
The majority of patients with cirrhotic ascites have marked activation of vasoconstrictor systems\cite{11-14}. Moreover, plasma concentrations of ANP and BNP are elevated in patients with cirrhosis and ascites\cite{15,16}, consistent with a physiological attempt to counterbalance vasoconstriction and antinatriuretic factors. However, in advanced cirrhosis, endogenous natriuretic peptides cannot override the pathophysiological events leading to ascites formation\cite{11}.

Earlier studies, that have investigated natriuretic peptides as treatment of cirrhotic ascites, have been too small to provide a definitive result. Thus, a review of the literature is justified. This systematic review and meta-analysis evaluates the benefits and harms of intravenous administration of the natriuretic peptides, ANP, BNP, or urodilatin in patients with liver cirrhosis and ascites. We aim to perform a meta-analysis of available data, especially regarding the effects on natriuresis, to determine the therapeutic potential of natriuretic peptides in cirrhotic ascites. Moreover, we intend to identify patient subpopulations for which treatment with natriuretic peptides may be particularly beneficial.

**MATERIALS AND METHODS**

This systematic review was performed in accordance with the PRISMA guidelines and the Cochrane Handbook\cite{17,18}. The preregistered review protocol (CRD42020195619) is available in PROSPERO.

We included trials evaluating mono-therapy of ANP, BNP, urodilatin, or any synthetic equivalent of these in patients with cirrhosis and ascites. Randomised controlled trials (RCTs) and non-randomised studies, e.g., trials investigating the pharmacodynamics of natriuretic peptides in cirrhosis patients with ascites and dose-finding trials, were considered for inclusion.

Trials were included regardless of dose and duration of the intervention or follow-up. Treatment administration had to be intravenous. As sodium retention is a main characteristic of decompensated cirrhosis, and existing pharmacological treatments aim to increase renal sodium excretion, the primary outcome in this systematic review was the maximum difference in natriuresis from baseline. To further evaluate renal and systemic treatment effects, secondary outcomes comprised the maximum difference in diuresis, plasma osmolality, urine osmolality, plasma creatinine concentration, plasma aldosterone concentration, and plasma renin activity from baseline. Change in bodyweight and waist circumference were included as objective surrogate measures of ascites burden. Non-serious and serious adverse events (AEs) were assessed focusing on the expected risk of blood pressure decreases and hypotension.

Trials were eligible for inclusion regardless of publication status, publication year, or language. The databases MEDLINE, Web of Science, Scopus, Cochrane Library and Embase were searched for eligible publications on the 13th of August 2020 (Supplementary material, page 1). References were managed in the reference sorting tool Covidence® (Covidence systematic review software, Veritas Health Innovation, Melbourne, Australia).

**Data management**

Study screening and data extraction to a spreadsheet with predefined variables were performed independently by two reviewers (RHG and MBK). Discrepancies were resolved through discussion.
before analyses. To characterise study populations, we collected background details including age, gender, cirrhosis aetiology, ascites severity, dietary sodium restriction, and diuretic withdrawal prior to treatment. We extracted data concerning the intervention including the name of the experimental intervention (generic and international non-proprietary name), comparator intervention, duration and dose of treatment, and whether a bolus or continuous infusion was applied. Finally, we collected information on study design, size of the trial, and follow-up duration.

Outcomes measured at baseline, during the intervention and at the end of follow-up were extracted from the text, tables or figures using a measuring tool from Adobe Acrobat 2017 (Adobe Inc., California, United States). The baseline outcome value was selected at the time point closest to the intervention exposure. For data concerning primary and secondary outcomes, the peak response or concentration measured during the intervention was used for trials administering a continuous intervention, while the peak response or concentration measured after the intervention was used for bolus infusion trials. Analyses were made using means and standard deviations (assumptions regarding units and unit conversions are described in Supplementary material, page 2).

**Risk of bias, quality assessments, and evidence**

Risk of bias assessments were performed at study level using the Cochrane risk of bias tool version 2 (RoB2)[19] for RCTs. An overall risk of bias judgment; low, moderate, or high was independently conducted by RHG and MBK. Non-randomised studies were assessed using the Newcastle-Ottawa-Scale (NOS)[20] (Supplementary material, pages 2 and 3) with a score from 0 (lowest) to 9 (highest). The score was transformed to generate a RoB2-comparable entity; NOS-scores 8-9 were classified as “low”, 5-7 as “moderate”, and ≤ 4 as “high”. The Grading of Recommendations Assessment and Evaluation was used to judge the quality of evidence for the study questions[21].

**Data synthesis and statistical analyses**

The effects of natriuretic peptides were summarised as overall peak mean and compared with overall baseline mean. This was performed for all trials and for study subgroups stratified by type of natriuretic peptide and continuous vs bolus infusions. When non-randomised studies outnumbered RCTs, baseline sodium excretion and peak sodium excretion were meta-analysed separately. We were unable to gather data allowing meta-analyses of safety data. Safety outcomes were therefore reviewed without statistical analyses.

The statistical software Stata, version 16.1, (StataCorp Lp, TX, United States) was used for data analyses. We applied a random-effects model with a DerSimonian-Laird tau² estimator for all meta-analyses. I² was used as a consistency measure. In situations with high levels of heterogeneity, meta-regressions were performed to elucidate study moderators. Moreover, the reasons for heterogeneity were addressed through subgroup analyses and leave-one-out analyses.

Small study effects were assessed using the Egger regression-based test. Funnel plots were included for graphical illustration of publication bias, while Trim-and-fill analysis were performed to estimate and account for this.

**RESULTS**

The literature search identified a total of 589 unique references. Ultimately, 22 publications describing 22 studies were included for data extraction (Figure 1)[17] covering 34 subgroups eligible for review and analyses (Table 1). ANP was studied in 19 studies (215 patients)[11-15,22-35], BNP in one study (seven patients)[16], and urodilatin in two studies (21 patients)[36,37].

The 22 included studies were conducted in eight countries and published between 1988 and 2007. Eighteen studies were non-randomised without a comparator, while four were cross-over RCTs using either placebo (3) or albumin (1) as a comparator. As the majority of included trials were non-randomised studies, we were unable to perform paired comparisons to compute effect sizes in the subsequent data analysis, except for urodilatin which was assessed in two RCTs. Fifteen studies investigated a continuous drug infusion, six studies applied a bolus injection, and one study assessed both a continuous infusion and a bolus injection (Table 1).

We classified ascites as mild, moderate, severe, or refractory. In a few situations, necessary assumptions were made regarding treatment dose and ascites severity to permit comparability (Table 1 and Supplementary material, pages 3 and 4).

**Background characteristics**

Two hundred and forty-three patients with cirrhosis and ascites were treated with intravenous natriuretic peptide (Table 1). The overall ratio of male and female participants was 3:1. Mean age ranged between 44.5 and 64.3 years (median 56.2 years). Ascites severity was reported in 21 of the included studies, covering 239 participants. Eighty-four (35%) had mild ascites, 81 (34%) had moderate ascites, 22 (9%) had severe/tense ascites, and 52 (22%) had refractory ascites. Only one study did not withdraw loop-diuretics and anti-aldosterone agents prior to the intervention[23]. The time from discontinuation...
Table 1 Main characteristics of the 22 included studies [34 subgroups; receiving bolus injection (n = 8) or continuous infusion (n = 26)]. Atrial natriuretic peptide was tested in 19 studies (215 patients), B-type natriuretic peptide in one study (7 patients), and urodilatin in two studies (21 patients).

<table>
<thead>
<tr>
<th>Ref.</th>
<th>Country</th>
<th>Study type</th>
<th>Patient n</th>
<th>Drug</th>
<th>Comparator</th>
<th>Adm</th>
<th>Dose (ng/kg/min or ng/kg)</th>
<th>Duration (min)</th>
<th>Post-infusion follow-up (min)</th>
<th>Age (mean)</th>
<th>Male (%)</th>
<th>Ascites severity: Mild/moderate/severe/refractory</th>
<th>S- or P-sodium (mmol/L) mean ± SE</th>
<th>Quality assessments (risk of bias)</th>
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<td>138.3 ± 0.7</td>
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<td>Bolus</td>
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<td>90</td>
<td>57.1</td>
<td>80</td>
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<td>133.3 ± 2.0</td>
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of diuretics until intervention in the remaining 21 studies ranged between one day and 14 d (median 7 d). Finally, a sodium restrictive diet was applied before the intervention in 18 studies ranging from 15 to 150 mmol/d (median 20 mmol/d). The duration of continuous infusions ranged from 30 to 180 min (median 120 min) with post-infusion follow-up period ranging from 0 to 180 min (median 60 min). Post-bolus follow-up ranged from 60 to 900 min (median 90 min). For studies administering a continuous infusion, the time point for peak natriuresis in general coincided with the treatment duration. For studies using a bolus injection, a peak response after 30 min was most commonly observed.

We found a substantial inter-study variation in definitions of responder and non-responder subgroups. As the majority of included trials introduced dietary sodium restriction prior to intervention, we redefined treatment responsiveness as the achievement of a negative sodium balance during or after the intervention.
One study investigated the natriuretic effect of continuous BNP infusion. No significant change in natriuresis was observed, nor any effect on diuresis and blood pressure\[16\]. Therefore, due to the lack of effect on natriuresis, BNP did not receive further attention in our analyses.

**Natriuresis**
Continuous infusion of ANP increased sodium excretion from 5.6 μmol/min [95% confidence interval (CI): 3.7-7.4] pre-intervention (baseline) to 27.0 μmol/min (95%CI: 41.5-103.5) at peak response. Interestingly, the peak response was significantly higher in study subgroups receiving an infusion rate of > 30 ng/kg/min compared with ≤ 30 ng/kg/min (P < 0.01) (Figure 2). The between-study heterogeneity was considerable (peak $I^2 = 88\%$, baseline $I^2 = 89\%$). Leave-one-out analyses did not recognise a single study responsible for the large $I^2$. To search for interactions to explain the heterogeneity we performed subgroup analyses exploring the effects of infusion duration, ascites severity, and dietary sodium restriction. No statistically significant differences were observed between subgroups exposed to short-term or long-term infusions (threshold at 60 min) (P = 0.07), nor between subgroups with a marked or moderate restriction of dietary sodium intake (threshold at 20 mmol/d) (P = 0.11). The baseline and peak sodium excretion were higher in subgroups with mild/moderate ascites compared with moderate/severe and refractory ascites (P < 0.01). Meta-regression found that the effect size was moderated by treatment dose and ascites severity, although adjusting for these parameters had only limited influence on the overall $I^2$. Effect size was not affected by quality assessment score, mean age, dietary sodium intake, and treatment duration. Study subgroups exposed to a bolus ANP injection were heterogeneous (peak $I^2 = 94\%$, baseline $I^2 = 93\%$) affecting the interpretability of the results. However, overall sodium excretion increased marginally (Supplementary Figure 1). Leave-one-out analyses did not identify any single study responsible for the heterogeneity. As only seven studies (covering eight study subgroups) applied a bolus injection, subgroup analyses and meta-regression were not justified. However, we performed a sensitivity analysis excluding the single study that did not draw diuretics prior to ANP injection. This manoeuvre slightly lowered the baseline and peak natriuresis.

Two cross-over RCTs investigated the natriuretic effect of urodilatin in cirrhosis patients with ascites\[36,37\], originating from the same study site. The patients in the earliest and larger trial had mild/moderate ascites and received 60 min infusion with 20 ng/kg/min, while the patients in the latest trial had refractory ascites and received 90 min of infusion with 20 ng/kg/min. Both studies reported a significant increase in sodium excretion as a response to the treatment compared with placebo, although the response was most pronounced in the larger trial investigating cirrhosis patients with mild/moderate ascites (Figure 3). The results suffered from substantial heterogeneity (P = 64%). A summary of the findings with quality of evidence are listed in Table 2. For natriuresis, the primary outcome of this systematic review, the quality of evidence was assessed as low for ANP and very low for urodilatin. These conclusions were primarily based on few RCTs being available, resulting in the dominance of non-randomised studies contributing to considerable heterogeneity, inconsistency, and imprecision due to small sample sizes.

**Diuresis**
Water excretion increased from baseline 1.3 mL/min (95%CI: 1.0-1.6) to 2.2 mL/min (95%CI: 1.7-2.7) in study subgroups receiving a continuous ANP infusion. The peak response tended to be higher in study subgroups receiving an infusion rate of > 30 ng/kg/min compared with ≤ 30 ng/kg/min (P = 0.08) (Figure 4). In our subgroup analysis, we found higher baseline and peak water excretion in study subgroups with a marked or moderate restriction of dietary sodium intake (P < 0.01). Dose, ascites severity, and quality assessment score explained between-study heterogeneity to some extent and especially in the low-dose group. When we adjusted for these factors overall heterogeneity remained substantial (P = 77%), but was reduced to moderate in the low-dose group (P = 54%). We were unable to identify parameters explaining heterogeneity in the high-dose group. Effect size was unaffected by mean age and treatment duration. Leave-one-out analyses also failed to identify any single study to explain the heterogeneity for this outcome.

Study subgroups exposed to bolus ANP injections had a tendency to increase diuresis with an estimated effect size similar to continuous infusion (Supplementary Figure 2). However, there was also considerable heterogeneity in this analysis (peak $I^2 = 94\%$, baseline $I^2 = 90\%$).

Urodilatin significantly increased urine production in the first and larger RCT\[36\] compared with placebo. However, in the most recent RCT with patients suffering from refractory ascites, water excretion was largely unaltered by urodilatin compared with placebo\[37\] (Supplementary Figure 3).

**Bodyweight, waist circumference, osmolality and creatinine**
Data were sparse on bodyweight and non-existent on waist circumference, thus these outcomes did not receive further evaluation. Two studies measured urine osmolality\[14,24\]. Laffi et al\[24\] reported a reduction in urine osmolality at the end of ANP-infusion and at the end of the recovery period compared with baseline for a subgroup of four participants. An increase was observed for five participants in another subgroup, while urine osmolality was unaffected in a third subgroup of six participants. Ferrier et al\[14\] reported no significant change in urine osmolality; however, a tendency...
Table 2 Effect of natriuretic peptides as treatment of cirrhotic ascites grouped by drug

<table>
<thead>
<tr>
<th>Outcomes</th>
<th>Anticipated absolute effects (95%CI)</th>
<th>No of participants (study subgroups)</th>
<th>Publication bias</th>
<th>Quality of evidence (GRADE)</th>
<th>Comments</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Baseline</td>
<td>Peak</td>
<td>Egger test</td>
<td>Trim-and-fill</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Baseline</td>
<td>Peak</td>
<td></td>
</tr>
<tr>
<td>Atrial natriuretic peptide</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Natriuresis (µmol/min)</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Continuous infusion</td>
<td>5.6 (3.7-7.4)</td>
<td>27.0 (19.4-34.7)</td>
<td>152 (23)</td>
<td></td>
<td>Low2,3</td>
</tr>
<tr>
<td>Low dose (≤ 30 ng/kg/min)</td>
<td>3.6 (2.0-5.3)</td>
<td>15.6 (8.7-22.4)</td>
<td>89 (13)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>High dose (&gt; 30 ng/kg/min)</td>
<td>24.2 (13.8-34.5)</td>
<td>72.5 (41.5-103.5)</td>
<td>63 (10)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Bolus injection</td>
<td>10.3 (4.3-16.2)</td>
<td>18.4 (8.8-28.0)</td>
<td>63 (8)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Diuresis (mL/min)</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Continuous infusion</td>
<td>1.3 (1.0-1.6)</td>
<td>1.8 (1.2-2.3)</td>
<td>62 (8)</td>
<td></td>
<td>Very low2,4</td>
</tr>
<tr>
<td>Low dose (≤ 30 ng/kg/min)</td>
<td>1.0 (0.7-1.3)</td>
<td>15.6 (8.7-22.4)</td>
<td>89 (13)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>High dose (&gt; 30 ng/kg/min)</td>
<td>1.8 (1.2-2.4)</td>
<td>2.8 (1.8-3.8)</td>
<td>63 (10)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Bolus injection</td>
<td>1.1 (0.8-1.5)</td>
<td>2.3 (1.3-3.2)</td>
<td>63 (8)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>P-Aldosterone (pmol/L)</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Overall</td>
<td>1182 (879-1484)</td>
<td>739 (553-925)</td>
<td>148 (21)</td>
<td></td>
<td>Very low2,4</td>
</tr>
<tr>
<td>Continuous infusion</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Low dose (≤ 30 ng/kg/min)</td>
<td>1895 (929-2861)</td>
<td>1110 (670-1553)</td>
<td>56 (8)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>High dose (&gt; 30 ng/kg/min)</td>
<td>753 (491-1015)</td>
<td>607 (412-801)</td>
<td>50 (8)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Bolus injection</td>
<td>1095 (449-1741)</td>
<td>435 (38-812)</td>
<td>42 (5)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>P-Renin activity (ng/mL/h)</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Overall</td>
<td>5.0 (3.7-6.2)</td>
<td>5.6 (4.2-6.9)</td>
<td>137 (20)</td>
<td></td>
<td>Very low2,4</td>
</tr>
<tr>
<td>Continuous infusion</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Low dose (≤ 30 ng/kg/min)</td>
<td>5.5 (2.8-8.2)</td>
<td>4.0 (2.2-5.8)</td>
<td>50 (7)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>High dose (&gt; 30 ng/kg/min)</td>
<td>2.6 (1.8-3.4)</td>
<td>6.5 (3.7-9.2)</td>
<td>54 (9)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Bolus injection</td>
<td>10.5 (3.8-17.2)</td>
<td>11.0 (2.9-19.1)</td>
<td>33 (4)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Bodyweight</td>
<td>NA</td>
<td>NA</td>
<td>NA</td>
<td>NA</td>
<td>Data too sparse for synthesis and quality assessment</td>
</tr>
<tr>
<td>Waist circumference</td>
<td>NA</td>
<td>NA</td>
<td>NA</td>
<td>NA</td>
<td>No available data</td>
</tr>
<tr>
<td></td>
<td>NA</td>
<td>NA</td>
<td>NA</td>
<td>NA</td>
<td>NA</td>
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<td>------</td>
<td>------</td>
<td>------</td>
</tr>
<tr>
<td><strong>P-Osmolality</strong></td>
<td>NA</td>
<td>NA</td>
<td>NA</td>
<td>NA</td>
<td>NA</td>
</tr>
<tr>
<td><strong>U-Osmolality</strong></td>
<td>NA</td>
<td>NA</td>
<td>NA</td>
<td>NA</td>
<td>NA</td>
</tr>
<tr>
<td><strong>P-Creatinine</strong></td>
<td>NA</td>
<td>NA</td>
<td>NA</td>
<td>NA</td>
<td>NA</td>
</tr>
<tr>
<td><strong>AEs</strong></td>
<td>Risk of any AEs: 20 AEs/70 participants = 29%</td>
<td>NA</td>
<td>NA</td>
<td>NA</td>
<td>NA</td>
</tr>
<tr>
<td><strong>BP reduction</strong></td>
<td>Subgroups reporting BP drops</td>
<td>NA</td>
<td>NA</td>
<td>NA</td>
<td>NA</td>
</tr>
<tr>
<td><strong>Continuous infusion</strong></td>
<td>53%</td>
<td>125 (19)</td>
<td>NA</td>
<td>NA</td>
<td>NA</td>
</tr>
<tr>
<td><strong>Bolus injection</strong></td>
<td>100%</td>
<td>63 (8)</td>
<td>NA</td>
<td>NA</td>
<td>NA</td>
</tr>
<tr>
<td><strong>B-type natriuretic peptide</strong></td>
<td>NA</td>
<td>NA</td>
<td>NA</td>
<td>NA</td>
<td>NA</td>
</tr>
<tr>
<td><strong>Urodilatin</strong></td>
<td>Mean difference (Urodilatin vs Placebo): 75 (−33–183)</td>
<td>21 (2)</td>
<td>Mean difference; NA</td>
<td>Very low²,⁴</td>
<td></td>
</tr>
<tr>
<td><strong>Natriuresis (µmol/min)</strong></td>
<td>Mean difference (Urodilatin vs Placebo): 2.5 (−0.9–5.8)</td>
<td>21 (2)</td>
<td>Mean difference; NA</td>
<td>Very low²,⁴</td>
<td></td>
</tr>
<tr>
<td><strong>Diuresis (mL/min)</strong></td>
<td>Risk of any AEs: 6 AEs/21 participants = 29%</td>
<td>21 (2)</td>
<td>NA</td>
<td>NA</td>
<td>NA</td>
</tr>
<tr>
<td><strong>BP reduction</strong></td>
<td>No BP drops observed for subgroups</td>
<td>21 (2)</td>
<td>NA</td>
<td>NA</td>
<td>NA</td>
</tr>
</tbody>
</table>

1For natriuresis and diuresis trim-and-fill analyses were performed separately for continuous and bolus applications, for all other outcomes exclusively on overall data. Trim-and-fill analyses were restricted to outcomes of atrial natriuretic peptide studies.

2Since the vast majority of studies were non-randomised studies, rating started at low quality of evidence. For urodilatin, two cross-over randomised controlled trials were available justifying rating initiation from high quality of evidence. No limitations due to risk of bias, since the majority of included studies obtained a low risk of bias score and the remaining studies a moderate risk of bias (Table 1). Overall results suffered from substantial or considerable heterogeneity with large I². However, the importance of inconsistency for decision-making is uncertain, since the majority of studies were early phase non-randomised clinical trials. Nevertheless, quality of evidence was rated 1 down for the presence of serious inconsistency. Study populations were considered similar without indirectness in interventions.

3Rated 1 up because of a dose-response gradient. Rated 1 up because of large effect defined as a more than two-fold increase in the outcome measure.

4Outcomes for atrial natriuretic peptide: Rated 1 down for imprecision due to small sample sizes. Outcomes for urodilatin: Rated 2 down for imprecision due to small sample sizes and the possibility of no effect.

5We were unable to estimate quality of evidence for the occurrence of adverse events (AEs), although our results indicated that one atrial natriuretic peptide or urodilatin induced AE may be expected to occur in approximately every fourth patient exposed to the drugs.

6We were unable to estimate quality of evidence for the occurrence of blood pressure drops, although our results clearly indicated a frequent occurrence of the event when exposed to atrial natriuretic peptide. For the two urodilatin studies, blood pressure reductions were not observed.

7Only one trial with seven participants evaluated the effect of B-type natriuretic peptide and two studies with 21 participants in total evaluated the effect of urodilatin. A justified quality of evidence assessment requires more data.

8For urodilatin, data were too sparse for plasma aldosterone concentration and renin activity to justify quality of evidence grading. Furthermore, no data were available for the outcomes bodyweight, waist circumference, plasma and urine osmolality, and plasma creatinine.

Patients or population: Patients with liver cirrhosis and ascites.

Setting: Hospital

Intervention: Intravenous bolus injections or continuous infusions of natriuretic peptides (atrial natriuretic peptide, B-type natriuretic peptide, or urodilatin).

Duration of continuous infusions: 30 to 180 min; Follow-up period bolus injections: 60 to 900 min; Follow-up period continuous infusions: 0 to 180 min; Comparison: Baseline levels/concentrations.

ANP: Atrial natriuretic peptide; BNP: B-type natriuretic peptide; AEs: Adverse events; BP: Blood pressure; NA: Not applicable.
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Figure 2 Effect of intravenous continuous atrial natriuretic peptide infusion on sodium excretion (μmol/min). Analyses were performed on overall data and separately for studies applying low dose infusion (≤ 30 ng/kg/min) and high dose infusion (> 30 ng/kg/min). ANP: Atrial natriuretic peptide; CI: Confidence interval.

towards a reduction was observed in the recovery period compared with baseline. None of the included studies published data on changes in plasma osmolality. Plasma creatinine was unaffected in the seven subjects investigated by Ferrier et al[14].

**Plasma aldosterone concentration and renin activity**

Repeated measurements of plasma aldosterone concentration were performed in 13 studies covering 21 study subgroups exposed to intravenous ANP, while another two studies covering three study subgroups restricted aldosterone concentration measurements to baseline. Plasma renin activity was
measured before and after initiation of the intervention with ANP in 20 study subgroups originating from 12 studies. Two additional studies with three study subgroups measured plasma renin activity exclusively at baseline. Hence, for evaluation of ANP effects on these two parameters we defined peak response as the value that deviated the most from baseline.

We obtained an impression of decreased aldosterone concentration when exposed to ANP, which was similar for low- and high-dose continuous infusions as well as bolus injections (Supplementary Figure 4). For plasma renin activity, no overall significant change was observed, although study subgroups exposed to high-dose continuous ANP infusion tended to increase their plasma renin activity (Supplementary Figure 5). For both outcomes, heterogeneity was considerable and meta-regression revealed minor modulation by administration approach (bolus or continuous) and ascites severity. Of specific interest, subgroup analyses revealed that baseline plasma aldosterone concentrations and renin activity were significantly lower in study subgroups that achieved a negative sodium balance on intervention, responders, compared with non-responder study subgroups ($P < 0.01$) (Figures 5 and 6).

Only the largest study with urodilatin reported characteristics on plasma renin activity and aldosterone concentration. Plasma renin activity during the intervention decreased insignificantly from baseline levels, while plasma aldosterone concentration significantly decreased during the intervention compared with baseline.

Safety
Specific AEs (Supplementary Table 1) were reported in five of the included trials[16,24,34,36,37]. Three studies stated that AEs were absent[12,25,26], while no information on AEs was provided by 14 studies [11,13-15,22,23,27-33,35]. The most frequently reported AEs were facial flushing and hypotensive episodes. The latter were reported on an individual level in studies without any effect on blood pressure at study subgroup level.

In general, the included studies presented thorough illustrations or descriptions of the blood pressure course before, during and after the intervention, with only three studies[28-30] not mentioning this variable. Since the studies were of small size, we considered both statistical significance and trends as occurrence of blood pressure reductions. Blood pressure was unaffected in the three trials investigating urodilatin and BNP[16,36,37] (Supplementary Table 2).

Eighteen study subgroups exposed to ANP, including all eight subgroups that received a bolus injection, showed reductions in either mean arterial blood pressure or mean systolic blood pressure during the intervention while the blood pressure remained stable in nine study subgroups. Change in blood pressure was unreported in four study subgroups. Full recovery in the post-infusion follow-up period was reported in six study subgroups, all exposed to a bolus infusion. Partial recovery occurred in five study subgroups. No recovery was observed in one study subgroup subsequent to a continuous infusion. The remaining six study subgroups lacked follow-up data (Supplementary Table 2). Regarding
Figure 4 Effect of intravenous continuous atrial natriuretic peptide infusion on water excretion (mL/min). Analyses were performed on overall data and separately for study subgroups receiving low dose infusion (≤ 30 ng/kg/min) and high dose infusion (> 30 ng/kg/min). ANP: Atrial natriuretic peptide; CI: Confidence interval.

the effect on blood pressure, our results clearly indicated continuous infusions of ≤ 30 ng/kg/min as well-tolerated, while higher doses almost guaranteed blood pressure reductions with a reduced likelihood of recovery in the follow-up period (Supplementary Table 2).

**Publication bias**
For all outcomes, Egger tests indicated statistically significant publication bias (Table 2). For peak natriuresis induced by ANP, funnel plots showed large asymmetry (Supplementary Figure 6). This may
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Figure 5 Baseline plasma aldosterone concentration in selected study subgroups. Non-responders failed to obtain negative sodium balance following intervention with atrial natriuretic peptide (ANP) and were characterised by significantly higher baseline plasma aldosterone concentration compared with ANP responders ($P < 0.01$). Baseline aldosterone levels were similar for subgroups exposed to bolus injections and continuous infusions, and extremely elevated in subgroups with refractory ascites compared with non-refractory ascites. Dotted line represents overall baseline mean plasma aldosterone concentration. ANP: Atrial natriuretic peptide; CI: Confidence interval.

DISCUSSION

We present here the first systematic review and meta-analysis of the benefits and harms of natriuretic peptides in patients with cirrhosis and ascites. In an attempt to investigate alternative treatments for cirrhotic ascites, this systematic review demonstrated a clinically relevant natriuretic effect in addition to diuretic and hormonal effects of natriuretic peptides. However, the levels of evidence are judged as low or very low, mainly due to considerable between-study heterogeneity, inconsistency, and imprecision due to small sample sizes. The most robust data exists for natriuresis induced by intravenous ANP administration, and especially when given by continuous infusion. Moreover, high-dose continuous infusions induce stronger sodium excretion than low-dose infusions, and subgroups with mild/moderate ascites have the most pronounced responses. Based on results from one single study, BNP is probably ineffective in this setting, although the patients included in the trial had marked sodium retention and activation of vasoconstrictor systems, and thus may have limited response to any natriuretic therapy[16]. Results from two studies investigating continuous urodilatin infusion suggest a clinically relevant natriuretic effect.
Baseline plasma renin activity - ANP

<table>
<thead>
<tr>
<th>Study subgroups</th>
<th>No. of study subgroups</th>
<th>Effect size with 95% CI</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Negative sodium balance:</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Non-responders</td>
<td>7</td>
<td>12.1 (8.3, 15.9)</td>
</tr>
<tr>
<td>Responders</td>
<td>13</td>
<td>2.5 (1.7, 3.3)</td>
</tr>
<tr>
<td>Test of group differences: $Q_0(1) = 22.91, P = 0.00$</td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Bolus/continuous:</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Studies applying a bolus injection</td>
<td>4</td>
<td>10.5 (3.8, 17.2)</td>
</tr>
<tr>
<td>Studies applying a continuous infusion</td>
<td>16</td>
<td>4.1 (3.0, 5.3)</td>
</tr>
<tr>
<td>Test of group differences: $Q_0(1) = 3.35, P = 0.07$</td>
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<td></td>
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<tr>
<td><strong>Ascites severity:</strong></td>
<td></td>
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</tr>
<tr>
<td>Mild to moderate</td>
<td>7</td>
<td>3.5 (1.8, 5.2)</td>
</tr>
<tr>
<td>Moderate to severe</td>
<td>9</td>
<td>4.2 (2.6, 5.9)</td>
</tr>
<tr>
<td>Refractory</td>
<td>3</td>
<td>12.7 (-14.2, 68.2)</td>
</tr>
<tr>
<td>Not reported</td>
<td>1</td>
<td>27.0 (-14.2, 68.2)</td>
</tr>
<tr>
<td>Test of group differences: $Q_0(3) = 7.76, P = 0.05$</td>
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</tr>
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</table>

Random-effects DerSimonian-Laird model

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Figure 6 Baseline plasma renin activity in selected study subgroups. Non-responders failed to obtain negative sodium balance following intervention with atrial natriuretic peptide (ANP) and were characterised by significantly higher baseline plasma renin activity compared with ANP responders ($P < 0.01$). Baseline renin activity tended to be lower in subgroups exposed to continuous infusion than for subgroups exposed to bolus injection, while subgroups with refractory ascites had markedly elevated renin activity compared with non-refractory ascites. Dotted line represents overall baseline mean plasma renin activity. ANP: Atrial natriuretic peptide; CI: Confidence interval.

Since the effects of ANP were discovered in the early 1980’s[38], investigations have been conducted to determine the therapeutic potential of natriuretic peptides in a broad range of diseases[39,40]. Through binding to specific natriuretic peptide receptors, ANP increases renal sodium and water excretion, increases glomerular filtration rate, inhibits aldosterone secretion, suppresses the release of arginine vasopressin, and reduces vascular resistance by smooth muscle relaxation[41-45]. These effects on fluid, electrolyte, and neuro-hormonal homeostasis reduce vasoconstriction and the intravascular volume, which may be beneficial for patients with fluid-accumulating disorders, including liver cirrhosis which is frequently complicated by ascites and peripheral oedema. Mild and moderate degrees of ascites are effectively treated with salt restriction and diuretics, but when ascites becomes refractory even to high doses of diuretics, treatment options are restricted to invasive procedures. Advances in diagnostic and therapeutic measures for refractory ascites remain unmet needs. Although a range of drugs, intended to alter essential events in the pathophysiological cascade, have undergone clinical investigations to relieve refractory ascites, none are currently approved.

Fyhrquist et al[40] were the first to report on the natriuretic and diuretic effects induced by intravenous ANP in refractory cirrhotic ascites and in the following decades further investigations with natriuretic peptides administered to cirrhotic patients were conducted. However, the trials sought primarily to elucidate the pharmacodynamics of natriuretic peptides in patients with cirrhosis and ascites, and some used ANP responsiveness to select eligible patients for other experiments e.g., combined treatment of ANP and norepinephrine to stabilize blood pressure[33], compare ANP responsiveness with responsiveness to head-out water immersion[25], compare ANP responsiveness with sympathetic nerve activity[13], test the inhibitory potential of Angiotensin-II on ANP responsiveness[28], and compare ANP responsiveness before and after establishment of a peritoneovenous shunt[29].
Thus, the general limitations of the trials are the small number of participants, different study designs, short-term interventions, varying treatment doses, and differences in ascites severity of participants. These limitations manifest as considerable heterogeneity in the current meta-analysis and reduce the level of evidence obtained for all outcomes. Eight studies investigating a bolus ANP injection were also included in this work. Although natriuresis may be induced with this setup, due to a short half-life bolus injection as therapy has limited clinical potential. As continuous intravenous infusions are more effective and applicable, this may be the preferred method of administration for future trials.

We included the outcomes bodyweight and waist circumference in our protocol since they may function as surrogate markers of ascites burden. Unfortunately, only a few trials weighed their participants and none measured girth; thus, data on these parameters are too sparse for analysis. A maximum ascites reabsorption capacity of less than one litre per day\cite{46} probably explains why these parameters were neglected for the majority of studies investigating short-term infusions of natriuretic peptides. However, due to the absence of these data the clinical relevance of natriuresis is still to be demonstrated. Serum sodium and potassium concentrations were not included as outcomes in the present study, although these electrolytes may be affected by natriuretic peptides. Baseline sodium concentration was measured in 12 study subgroups (Table 1). In only one study a follow-up measurement after ANP infusion, which was unchanged from baseline, was reported. The baseline serum sodium concentration generally reflects the disease severity, with the lowest concentrations in study subgroups characterised by refractory ascites, and thus, may be a potential predictor of ANP responsiveness. Exclusively at baseline, serum potassium concentration was measured in six study subgroups, which showed similar results. A RCT investigating longer-term urodilatin infusion is currently recruiting patients, and includes bodyweight, waist circumference, plasma sodium and plasma potassium as outcomes\cite{47}.

A notable finding of this meta-analysis is the observation that non-responders unable to achieve a negative sodium balance when treated with a natriuretic peptide were characterised by high baseline plasma aldosterone concentrations and renin activity. Plasma levels of these parameters indicate the severity of renin-angiotensin-aldosterone (RAAS)-activation, which directly parallels the degree of cirrhosis decompensation with portal hypertension. Therefore, we hypothesise that a threshold exists where RAAS-activation is too severe for ANP to counteract. Future trials may benefit from inclusion of these hormones to validate their ability to identify treatment responders and non-responders, and further investigate if a threshold predicting responsiveness can be estimated.

AEs and especially effects on blood pressure were assessed in this systematic review. The results indicated the incidence of any AE in every fourth patient receiving treatment with either ANP or urodilatin, and the AEs described were predominantly subjective measures, which may easily be explained by the well-known physiological effects of natriuretic peptides. Blood pressure is sensitive to intravenous administration of natriuretic peptides, which is well-documented in other diseases, e.g., decompensated heart failure\cite{48}. Our results clearly indicate that bolus ANP injections guarantee reductions in blood pressure, although with a high chance of rapid recovery. For continuous ANP infusions, doses ≤ 30 ng/kg/min are less likely to affect blood pressure compared with higher doses. Future trials are needed before a definite conclusion on the tolerability of natriuretic peptides in cirrhotic patients with ascites can be made.

The present study is limited to the inclusion of cirrhosis patients with ascites treated with either ANP, BNP, or urodilatin. Thus, we have not included results from studies investigating natriuretic peptide effects in cirrhosis patients without ascites. Furthermore, we have excluded studies focusing on interventions intended for blockade of natriuretic peptide degradation. One trial has tested the effects of sinorphan, an enkephalinase inhibitor, in patients with cirrhosis and ascites. Enkephalinase is a neutral membrane endopeptidase with a broad presence in plasma and tissues including the kidneys\cite{49}. By blocking this enzyme, plasma levels of ANP and the second messenger cyclic guanosine monophosphate increase. Sixteen cirrhosis patients with persistent ascites received treatment with sinorphan in a double-blind, placebo-controlled, cross-over trial. Plasma concentrations of ANP increased significantly after drug administration and sodium excretion doubled (mean increase 17.2 µmol/min) in the 2-h post-administration period\cite{49}. Interestingly, mean arterial blood pressure was unaffected. Therefore, blockade of ANP degradation may be of relevance for future trials, potentially with limited effects on blood pressure.

Another limitation of the present study is the age of the included studies. All but one of the studies were performed and published more than 20 years ago. The distribution of cirrhosis aetiologies has changed during the past two decades, particularly due to an increased frequency of cirrhosis due to metabolic disease. However, we focus here on cirrhosis decompensation with ascites, which occurs late in the disease course, allowing translation of the results to the present.

**CONCLUSION**

Ultimately, this systematic review and meta-analysis provides new perspectives into future medico-pharmaceutical research in patients with advanced cirrhosis and elucidates the course for future
research in the field. A clear natriuretic and diuretic effect of intravenous ANP is evident from our results, and we suggest that future trials would benefit from a RCT design with a comparator group, longer-term infusion, and clinical follow-up. The preferred dose has to be determined on the basis of larger natriuretic and diuretic effects with doses > 30 ng/kg/min, but better tolerability of doses ≤ 30 ng/kg/min. Although the natriuretic and diuretic effects are modest in patients with severe or refractory ascites, new treatments are most needed for this indication. This argues for a particular focus on these patients in future trials. To increase the treatment duration and maintain increased natriuresis, alternative methods of administration must be considered e.g., continuous subcutaneous infusion. Finally, clinicians may find a prolongation of the interval between paracentesis as a sufficiently meaningful treatment effect and not necessarily a reversion from refractory ascites to treatment responsive ascites.

ARTICLE HIGHLIGHTS

Research background
Ascites formation is the most frequent event of cirrhosis decompensation with no approved non-invasive treatment for patients with refractory ascites. However, natriuretic peptides may induce beneficial effects in patients with cirrhosis and ascites, by counterbalancing vasoconstriction and anti-natriuretic factors.

Research motivation
Only small studies with a broad pallet of designs, have investigated the renal and systemic effects and safety of natriuretic peptides in patients with cirrhosis and ascites. We were motivated to compile results regarding natriuretic peptides as treatment for ascites to advance pharmaceutical research concerning this disease.

Research objectives
We aimed to systematically review the effects and safety of natriuretic peptides in patients with cirrhosis and ascites. We collected results regarding changes in renal sodium and water excretion, plasma aldosterone concentration, and plasma renin activity, and to perform meta-analyses providing overall estimates of the effects of natriuretic peptide infusions. Safety assessments, in particular blood pressure decreases, were included and descriptively summarised.

Research methods
We adhered to the PRISMA guidelines and the Cochrane Handbook. A review protocol was preregistered at PROSPERO. The databases MEDLINE, Web of Science, Embase, Scopus, and Cochrane Library were searched and references reporting the renal and systemic effects of mono-therapy with atrial natriuretic peptide (ANP), B-type natriuretic peptide, urodilatin, or any synthetic equivalent in patients with cirrhosis and ascites were considered for inclusion. Treatment administration had to be intravenous, but trials were included regardless of dose, follow-up duration, and whether a continuous infusion or bolus injection was administered. The study screening and data extraction were performed independently by two reviewers. A random-effects model was applied for the meta-analysis. Reasons for heterogeneity were explored through subgroup and leave-one-out analyses.

Research results
Twenty-two studies were included and short-term ANP infusion was the only intensively studied treatment. Renal sodium and water excretion increased in response to continuous ANP infusion and was most pronounced when infusion rates > 30 ng/kg/min were applied. Furthermore, sodium excretion was higher in study subgroups with mild/moderate ascites compared with refractory ascites. The baseline plasma aldosterone concentration and renin activity were significantly lower in subgroups achieving a negative sodium balance compared with treatment non-responders, and may be relevant response predictors to include in future trials. Blood pressure decreases occurred less frequently with doses ≤ 30 ng/kg/min.

Research conclusions
Intravenous infusions of natriuretic peptides induce meaningful sodium and water excretion in patients with cirrhotic ascites. Continuous ANP infusions > 30 ng/kg/min induce the most pronounced renal effects, but with a higher risk of blood pressure decreases than doses ≤ 30 ng/kg/min.

Research perspectives
Future larger-scale clinical trials are justified to determine the therapeutic potential of natriuretic peptides in patients with cirrhotic ascites.
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FOOTNOTES

Author contributions: Gluud LL and Grønbæk H contributed equally to the work. All authors contributed to the design of the study. Gantzel RH and Kjær MB acquired the study data; Gantzel RH and Gluud LL performed the statistical analyses; all authors contributed to the interpretation of data; Gantzel RH drafted the manuscript; and all authors critically reviewed and approved the final manuscript.

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PRISMA 2009 Checklist statement: This review was performed in accordance with the PRISMA guidelines.

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