Combination of rifaximin and lactulose improves clinical efficacy and mortality in patients with hepatic encephalopathy

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Background: Rifaximin and lactulose are common effective agents for hepatic encephalopathy (HE). Whether a combination of rifaximin and lactulose improves the efficacy and mortality in patients with HE compared with lactulose alone needs to be analyzed.

Methods: A systematic search was performed in electronic databases and other sources for possible studies focusing on combination therapy of rifaximin and lactulose for HE between January 2000 and February 2018. A meta-analysis was performed by the method recommended by the Cochrane Collaboration, and estimated effect size was presented as risk difference (RD), 95% CI, and the number needed to treat (NNT). Subgroup analysis, sensitivity analysis, and Trial Sequence Analysis were comprehensively performed to indicate the source of heterogeneity and risk of bias.

Results: Five randomized and five observational studies involving 2,276 patients were included. Combination therapy had a significant advantage in both clinical efficacy increase (RD 0.26, 95% CI 0.19–0.32, NNT 5) and mortality decrease (RD = –0.16, 95% CI –0.20–0.11, NNT 9) in overall analysis. In the pooled analysis of randomized studies, combination therapy showed similar results in clinical efficacy (RD 0.25, 95% CI 0.16–0.35, NNT 4) and mortality (RD = –0.22, 95% CI –0.33–0.12, NNT 5). Compared with lactulose, hospital stay was also reduced in combination therapy, and there was no significant difference in treatment-related adverse events between the two groups.

Conclusion: Combination of rifaximin and lactulose has beneficial effects on HE. Compared with lactulose alone, additional rifaximin increases clinical efficacy and decreases mortality. However, its effects on different types of HE are still uncertain.

Keywords: rifaximin, lactulose, combination therapy, HE, meta-analysis

Introduction

Hepatic encephalopathy (HE) is described as a brain dysfunction caused by liver insufficiency and/or portosystemic shunting; it manifests itself as a wide spectrum of neurological or psychiatric abnormalities ranging from subclinical alterations to coma. HE is a severe complication of acute or chronic liver failure due to cirrhosis; its prevalence and severity is closely related to the underlying liver status. The prevalence of minimal HE in cirrhosis ranged from 20% to 80%. For overt HE, the prevalence was 10%–14% in general cirrhosis, 16%–21% in decompensated cirrhosis, and 10%–50% in patients who adopted transjugular intrahepatic portosystemic, which was a major mini-invasively shunting surgery. According to an over-5-year analysis, the mortality of HE in hospital was about 15%. For cirrhosis patients without stable management or cure, 30%–40% of them suffered repeatedly recurrent HE in their survival periods.
Besides intensive care and underlying liver disease treatment, 90% of HE patients can be treated through managing precipitating factors, including gastrointestinal bleeding, excessive protein intake, infection, hypokalemic alkalosis, constipation, hypoxia, or the use of sedatives and tranquilizers. About 60%–80% of HE patients showed an elevated serum ammonia, and thus therapy was recommended, including non-absorbable disaccharides (NAD), rifaximin, L-ornithine-L-aspartate (LOLA), and branched chain amino acids (BCAA), which mostly aim to reduce the level of serum ammonia. Among them, lactulose is the first choice and the most widely adopted NAD, while rifaximin emerged as an effective non-absorbed oral antimicrobial agent in recent years. However, the two drugs presented similar clinical effects, which were demonstrated by large-scale meta-analysis studies.

Because rifaximin and lactulose had different pharmacological mechanisms for HE, it is important to understand whether a combination of them would further increase the clinical efficacy compared to lactulose alone. Investigated by a series of small-scale studies, the issue was not fully evaluated and has yet to provide a confirmed conclusion. Therefore, we performed a comprehensive systematic review and meta-analysis of published clinical studies aiming to determine the comparative efficacy and safety of combined rifaximin and lactulose with lactulose alone for current HE treatment.

Methods

Search strategy

Current meta-analysis was reported in accordance with Preferred Reporting Items for Systematic Reviews and Meta-Analyses Guideline. Comprehensive searches including electronic databases, clinical register centers, scholar search engine and manual search were completed using modified search strategy. Search terms were “hepatic encephalopathy, rifaximin, lactulose, plus, and combination”. Online search was carried out by searching the websites of PubMed, EMBASE, the Cochrane Library, CNKI, Wanfang, ClinicalTrials.gov, Google scholar, and Baidu scholar between January 2000 and February 2018. Manual search was done by screening the references and citations of the similar studies. Publication language was limited to English and Chinese.

Study inclusion and data extraction

Both randomized and observational controlled studies investigating specific topics were considered in our analysis. Relevant meta-analyses and systematic reviews were also searched. Participants were patients suffering from covert (the West Haven criteria, < grade 2) and overt HE (the West Haven criteria, ≥ grade 2) due to liver cirrhosis. Combination therapy of rifaximin and lactulose was compared with lactulose alone in the treatment of HE, and the specific dose and usage of rifaximin were not restricted; also, there were no restrictions of the control type (bland control or placebo). Outcome measurements were clinical efficacy, mortality, and treatment-related adverse events. Two reviewers independently assessed the eligibility of all potential citations obtained from initial search, and a third reviewer checked the included studies.

Data extraction was also completed by two independent reviewers. Baseline study information, patients’ characteristic, comparison, type of HE, etiology and severity of HE, treatment duration, and follow-up period for evaluation of clinical outcomes were extracted from each study; the number of events of interest in studies were extracted for further statistical analysis purpose. Discrepancies regarding the extraction of data were resolved by the third reviewer.

Outcome measures

Primary outcome measures included clinical efficacy and mortality. Clinical efficacy was defined as improvement in the HE clinical syndrome with improved neurological status or a significant decrease in the HE index after treatment. Secondary outcome measure was treatment-related adverse events such as severe diarrhea, episodes of intense abdominal pain, and other gastrointestinal system reactions.

Assessment of risk of bias

Current analysis included both randomized and observational studies, and risk of bias located in each study was independently assessed by two reviewers by using different scales. Modified JADAD scale was used for randomized controlled trials, which contained items assessing sequence generation for the randomization, allocation of treatment concealment, blinding of participant and outcome measures, follow-up, and drop out. Trials with a score of ≤3 (total 7 scores) were considered as having high risk of bias and >3 were considered as having low risk of bias. Newcastle-Ottawa Scale (NOS) was used for observational cohort studies, which contained items assessing risk in exposed cohort representativeness, non-exposed cohort selection, ascertainment of exposure, comparability of cohort, interested outcome assessment, and follow-up periods. Studies achieving a score ≥4 (total 9 scores) were considered to be low risk of bias. Specific checklists for randomized controlled trials and observational
Combined rifaximin and lactulose for hepatic encephalopathy

Statistical analysis
Review Manager (version 5.3, Cochrane Center, Copenhagen, Denmark) was used to perform this meta-analysis. In the software, $I^2$ and corresponding $P$-value were used to investigate the significance of statistical heterogeneity. Moderate, considerable, and substantial heterogeneity were set based on the value of $I^2$ range from 30% to 60%, 60% to 75%, and >75%. Clinical heterogeneity was handled by subgroup analysis according to study design (randomized studies or not). According to $I^2$ value, random-effects model or fixed-effects model was selected to analyze the data extracted from published reports. For each outcome measure under random-effects model, the result of fixed-effect model was reported only if the difference between the two models existed. Current meta-analysis used risk difference (RD) indicating the effect size of categorical variables and mean difference (MD) indicating the effect size of continuous variables together with their 95% CI. Both of them represented an average MD between the groups, and $P$-value $<0.05$ was considered to be statistically significant.

For primary outcome measures, the number needed to treat (NNT) was also calculated, which represented the number of patients needed to treat to achieve a different clinical outcome. Based on the data of absolute risk reduction (ARR), experimental event rate (EER), and control event rate (CER) in low risk of bias studies, NNT was obtained as $NNT = 1/ARR$ and $ARR = CER$. The 95% CI of NNT was calculated based on the lower limit (LL) and upper limit (UL) of 95% CI of ARR: $NNT_{LL} = 1/ARR_{UL}$ and $NNT_{UL} = 1/ARR_{LL}$. Statistical significance was identified by the range of 95% CI that it did not cross the value of 1. We further confirmed the reliability of the results in Trial Sequential Analyses (TSA) by using TSA software (Copenhagen Trial Unit, Copenhagen, Denmark). The sequential analyses were based on random-effects model, with $\alpha=5\%$ and a test power of 80%, and model-based heterogeneity (diversity). The analysis results were judged as highly reliable when the z-curve crossed the trial monitoring curve, which indicated that cumulative result of included trials was relative enough to achieve a certain result even if the required sample size was not reached.

Results
Study information
During the search process, 360 citations including 64 duplicates were obtained. After screening the titles and abstracts, 23 studies were assessed for eligibility by reading full-texts (Figure 1). Finally, seven studies with full text and two cohort studies of Scottish Intercollegiate Guidelines Network were also used.

Figure 1 Process of including studies evaluating combination therapy for hepatic encephalopathy.
Table 1 Baseline characteristics of the included studies

<table>
<thead>
<tr>
<th>Studies</th>
<th>Design</th>
<th>Type</th>
<th>Case (n)</th>
<th>Age (years)</th>
<th>Sex (n, M:F)</th>
<th>Type of HE</th>
<th>Etiology</th>
<th>Severity</th>
<th>Comparison</th>
<th>Treatment duration (days)</th>
<th>Follow-up</th>
</tr>
</thead>
<tbody>
<tr>
<td>Gao 2012&lt;sup&gt;3&lt;/sup&gt;</td>
<td>RCT</td>
<td>Full-text</td>
<td>31/31</td>
<td>58/61</td>
<td>22.9/19.12</td>
<td>HE</td>
<td>Cirrhosis</td>
<td>Not reported</td>
<td>RA 1,100 mg + LA 60 mL/LC</td>
<td>10–15</td>
<td>10–15</td>
</tr>
<tr>
<td>Neff et al 2013&lt;sup&gt;3&lt;/sup&gt;</td>
<td>OCT</td>
<td>Abstract</td>
<td>40/352</td>
<td></td>
<td></td>
<td>HE</td>
<td>Cirrhosis</td>
<td>MELD: 13.1/10.2</td>
<td>RA 1,100 mg + LC/LC</td>
<td>Not reported</td>
<td>Not reported</td>
</tr>
<tr>
<td>Sharma et al 2013&lt;sup&gt;5&lt;/sup&gt;</td>
<td>RCT</td>
<td>Full-text</td>
<td>63/57</td>
<td>40.4±8.5/37.5±10.5</td>
<td>47:16/42:15</td>
<td>Overt HE</td>
<td>Alcohol: 40/32; HBV: 22/10; HCV: 7/3; Other: 19/10</td>
<td>CPT: 9.9±2.9/9.2±2.5; MELD: 24.9±6.6/23.8±5.18</td>
<td>HE grade (2/3/4): 22/40/58</td>
<td>≤10</td>
<td>During hospital stay</td>
</tr>
<tr>
<td>Gill et al 2014&lt;sup&gt;4&lt;/sup&gt;</td>
<td>RCT</td>
<td>Abstract</td>
<td>100/100</td>
<td>40</td>
<td>70/30/70.30</td>
<td>Overt HE</td>
<td>Cirrhosis</td>
<td>HE grade (2/3/4): 60/70/70</td>
<td>RA 1,100 mg + LA 60–180 mL/placebo + LA</td>
<td>10</td>
<td>During hospital stay</td>
</tr>
<tr>
<td>Miu et al 2014&lt;sup&gt;4&lt;/sup&gt;</td>
<td>OCT</td>
<td>Full-text</td>
<td>80/80</td>
<td>41±8.9</td>
<td>89:71</td>
<td>Overt HE</td>
<td>Alcohol: 3; HBV: 7; HCV: 139; Other: 11</td>
<td>CPT (A/B/C): 1/49/110; HE grade (1/2/3/4): 3/29/51/77</td>
<td>RA 1,100 mg + LA 90 mL/LA</td>
<td>≤7</td>
<td>7 days</td>
</tr>
<tr>
<td>Courson et al 2015&lt;sup&gt;18&lt;/sup&gt;</td>
<td>RCT</td>
<td>Full-text</td>
<td>62/87</td>
<td>58±11/59±12</td>
<td>22.40/47.40</td>
<td>Acute HE</td>
<td>Alcohol: 52/27; HBV: 4/1; HCV: 16/6; Other: 35/37</td>
<td>MELD: 18/21</td>
<td>RA 1,100 mg + LA/LA</td>
<td>≤10</td>
<td>180 days</td>
</tr>
<tr>
<td>Kang et al 2017&lt;sup&gt;19&lt;/sup&gt;</td>
<td>OCT</td>
<td>Full-text</td>
<td>145/276</td>
<td>58.6±11.5/60.2±12.0</td>
<td>92.53/167:109</td>
<td>HE</td>
<td>Alcohol: 55/90; Viral: 64/126; Others: 26/60</td>
<td>CPT (A/B/C): 15/174/232</td>
<td>Same as Ref 15</td>
<td>Not reported</td>
<td>540 days</td>
</tr>
<tr>
<td>Kang et al 2017&lt;sup&gt;19&lt;/sup&gt;</td>
<td>OCT</td>
<td>Full-text</td>
<td>173/448</td>
<td>63.28±9.8/64.23±9.9</td>
<td>143:30/35:197</td>
<td></td>
<td>Alcohol: 134/353; Viral: 24/59; Others: 15/36</td>
<td>CPT (A/B/C): 14/170/337</td>
<td>Same as Ref 15</td>
<td>Not reported</td>
<td></td>
</tr>
<tr>
<td>Hasan et al 2018&lt;sup&gt;11&lt;/sup&gt;</td>
<td>RCT</td>
<td>Full-text</td>
<td>45/46</td>
<td>44.7±10.6/44.9±10.1</td>
<td>36.9/38.8</td>
<td>Overt HE</td>
<td>Alcohol: 42/39; Others: 3/7</td>
<td>Not reported</td>
<td>RA 1,200 mg + LA 60–120 mL/placebo + LA</td>
<td>≤10</td>
<td>10 days</td>
</tr>
</tbody>
</table>

Abbreviations: CTP, Child–Turcotte–Pugh score; HBV, hepatitis B virus; HCV, hepatitis C virus; HE, hepatic encephalopathy; LA, lactulose (the amount of lactulose was 667 mg/mL); MELD, model for end-stage liver disease; OCT, observational controlled trials; RA, rifaximin; RCT, randomized controlled trial.
studies with abstracts were included, among them one study divided patients into two groups (with HCC or not), which were regarded as two separate trials. A total of 771 patients were assigned to combination group (rifaximin and lactulose) and 1,505 patients were assigned to control group (lactulose alone). Demographic characteristics are presented in Table 1. Six studies included overt HE patients (the West Haven criteria, ≥ grade 2), and three studies included new onset and recurrent HE patients without detailed grade of the West Haven criteria. The etiology of cirrhosis included alcohol, hepatic virus infection, and others. The severity of HE was judged by the abovementioned HE grade. The Child Pugh score and model for end-stage liver disease score judged the severity of underlying liver disease. Among them, HE grade >1 was diagnosed as overt HE, which indicated that clinical findings such as lethargy or apathy, disorientation for time, and obvious personality change were reproducible. The dose of rifaximin was fixed in the studies that 1,100 mg therapy was adopted in five studies, and 1,200 mg therapy was adopted in four studies. The volume of lactulose varied in each study, ranging from 60 mL to 180 mL (the amount was 667 mg in 1 mL). The treatment duration was mainly less than or equal to 10 days, and only one study reported a maximum treatment duration of 15 days. The follow-up period for analysis of clinical efficacy and mortality was similar to treatment duration, as the outcomes were mostly measured during hospital stay, except for two studies that had a follow-up period >180 days.

**Risk of bias**

Five randomized studies were assessed by JADAD scale, three of them had a score >3 and three of them designed a placebo control. Four observational studies were assessed by NOS scale, and three of them had a score >4. Both of them were assessed by SIGN scale, where two studies were judged as high quality, five studies were judged as acceptable, and only one study was judged as low quality. The assessment result of risk of bias is shown in Table 2.

**Clinical efficacy**

Six studies evaluated the effects of the combination of rifaximin and lactulose on HE (Figure 2). The meta-analysis result showed that combination therapy significantly increased clinical efficacy compared with lactulose alone in HE patients (RD 0.19, 95% CI 0.09–0.29, P=0.0002). The heterogeneity value of I² was 59%, and one study was identified to be responsible for this moderate heterogeneity through inverted funnel plot. After excluding it, the value of I² was reduced to 35%; the sensitivity analysis result was consistent with that of before (RD 0.26, 95% CI 0.19–0.32, P=0.00001).

Randomized studies were regarded to be better than non-randomized studies in the study design; accordingly, we analyzed the result based on only randomized studies (Figure 3). In the new pooled analysis, three studies containing 342 HE patients also demonstrated a significant increase in clinical efficacy of combination therapy (RD 0.25, 95% CI 0.16–0.35, P<0.00001). The NNT was 5 (95% CI 3.45–11.11) in primary analysis, while it was 4 in sensitivity analysis (95% CI 3.13–5.26) and randomized study analysis (95% CI 2.86–6.25). In TSA analysis, we set α to 5%, test power to 80%, control group incidence to 54%, relative risk reduction to −26%, and heterogeneity correction to 35%. The required sample size was 615 participants. The monitoring boundary was crossed in 2013, which confirmed the reliability of current meta-analysis result with enough required sample size (Figure 4). Inverted funnel plot indicated a low risk of bias (Figure 5).

**Mortality**

Seven studies reported the data of mortality (Figure 6). In the pooled analysis, combination therapy significantly reduced the mortality in HE patients compared with lactulose alone (RD −0.11, 95% CI −0.19 to −0.03, P=0.009). Considerable heterogeneity was found (I²=70%). Two studies were identified to be different with others through an inverted funnel plot. Exclusion of the studies decreased the value of I² to 41%, and sensitivity analysis result showed a similar trend to before (RD −0.16, 95% CI −0.20 to −0.11, P<0.00001).

In the new pooled analysis based on randomized studies, two studies containing 280 patients demonstrated a significant

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Table 2 Risk of bias assessment of the included studies

<table>
<thead>
<tr>
<th>Studies</th>
<th>SIGN</th>
<th>JADAD</th>
<th>NOS</th>
</tr>
</thead>
<tbody>
<tr>
<td>Gao 2012</td>
<td>Acceptable</td>
<td>3</td>
<td>–</td>
</tr>
<tr>
<td>Neff et al 2013</td>
<td>Low quality</td>
<td>–</td>
<td>4</td>
</tr>
<tr>
<td>Sharma et al 2013</td>
<td>High quality</td>
<td>6</td>
<td>–</td>
</tr>
<tr>
<td>Gill et al 2014</td>
<td>Acceptable</td>
<td>3</td>
<td>–</td>
</tr>
<tr>
<td>Miu et al 2014</td>
<td>Acceptable</td>
<td>–</td>
<td>5</td>
</tr>
<tr>
<td>Courson et al 2015</td>
<td>Acceptable</td>
<td>4</td>
<td>–</td>
</tr>
<tr>
<td>Kang et al 2017</td>
<td>High quality</td>
<td>–</td>
<td>9</td>
</tr>
<tr>
<td>Ahire and Sonawale 2017</td>
<td>High quality</td>
<td>–</td>
<td>8</td>
</tr>
<tr>
<td>Hasan et al 2018</td>
<td>High quality</td>
<td>6</td>
<td>–</td>
</tr>
</tbody>
</table>

**Abbreviations:** JADAD, JADAD scale; NOS, Newcastle-Ottawa Scale; SIGN, Scottish Intercollegiate Guidelines Network.
decrease of mortality in combination therapy (RD –0.22, 95% CI –0.33 to –0.12, P<0.0001, I²=0%; Figure 7).¹⁵,¹⁶

The NNT was 9 (95% CI 5.26–33.33) in primary analysis, 6 (95% CI 5.00–9.09) in the sensitivity analysis, and 5 (95% CI 3.03–8.33) in randomized study analysis. In TSA analysis, we set α to 5%, power to 80%, control group incidence to 67%, relative risk reduction to 16%, and heterogeneity correction to 41%. The required sample size was 1,248 participants. The monitoring boundary was crossed in 2014, which confirmed the reliability of current meta-analysis result with enough sample size (Figure 8). Inverted funnel plot indicated low risk of bias (Figure 9).

Based on available data of one study including 120 patients (HE grade 2/3/4 was 22/40/58) with a mortality of 35.83%;¹⁵ deaths were caused by sepsis (7/17, P=0.01), hepatorenal syndrome (4/7, P>0.05), and gastrointestinal bleeding (4/4, P>0.05) in each group.

Hospital stay
Three studies reported the data of hospital stay. Two of them were pooled analyzed, and meta-analysis results showed that combination therapy significantly reduced the duration of hospital stay (MD –2.89, 95% CI –3.52 to –2.25, P<0.00001).¹⁵,¹⁶ The other study only reported the data of median hospital stay between the groups (6/8, P=0.09),¹⁸ which found no significant difference.

Adverse events
Four studies compared the treatment-related adverse events (Figure 10). In the pooled analysis, there was no significant difference in combination therapy and lactulose alone (RD –0.06, 95% CI –0.24 to 0.13, P=0.56). One study was found to be responsible for the substantial heterogeneity (I²=90%); after excluding it,¹⁴ the I² value was obviously reduced (I²=0%). The sensitivity result was similar to before (RD 0.0, 95% CI –0.03 to 0.02, P=0.63).

Discussion
HE is one of the most common causes of death and would cause many precipitating factors in cirrhosis patients.²² According to the recommendations in the 2014 practice guideline, controlling precipitating factors is of paramount importance in the management of HE. Besides precipitating factor management, lactulose is the first recommended agent. Rifaximin, oral BCAA, intravenous LOLA, neomycin, and metronidazole can also be used as alternative or additional agents.¹ However, high-level evidence such as meta-analysis or systematic review is still lacking.
The current meta-analysis was the first to quantitatively evaluate the effective effects of combination therapy of rifaximin and lactulose vs lactulose alone for the management of HE. We included five randomized and five observational controlled studies, with a sample size of 2,276 HE patients, and demonstrated that combination therapy significantly improved the clinical efficacy, mortality, and hospital stay without increasing treatment-related adverse events in HE.

As stated within the practice guideline, current definition, diagnosis, classification, or the treatment of HE are not universally accepted, and the complex pathogenesis of HE is difficult to clarify. Currently, clinical observations always exhibit a high average level of ammonia in high grade HE patients; thus, the ammonia hypothesis is a widely accepted premise that leads to frequent assessment of ammonia concentrations. Conventional pharmacologic treatment of HE consisted of NAD since 1966, which mainly included lactulose and lactitol. A meta-analysis including 38 randomized trials published in 2016 had sufficiently confirmed the effectiveness of the routine use of NAD to reduce the production and absorption of gut-derived neurotoxin ammonia in clinical practice, as well as the potential roles of catharsis, ammonia metabolism, and gut microbiome adjustment.

For a specific HE patient, the serum level of ammonia sometimes did not correlate with the severity of clinical symptoms.
Therapies targeting the infection and inflammation became a promising agent. Inflammation deriving from the gut because of bacterial overgrowth was related to a non-response rate as high as 22%. Accordingly, improved therapy based on lactulose is warranted since no certain method was introduced to identify a non-responder of lactulose therapy. Confirming by sensitivity analysis, subgroup analysis, Trial Sequential Analysis, and publication bias analysis, our meta-analysis shows that a combination of rifaximin to lactulose can significantly improve both clinical efficacy of HE manifestations (NNT =4) and mortality (NNT =5) mainly for overt HE patients. Based on limited data in one randomized study, the cause of deaths was significantly reduced only in aspect of sepsis, which indicated that combination therapy may improve clinical outcome by reducing the incidence of severe and systemic infection and inflammation. In past years, multiple factors were included in the explanation of HE progress; among them systemic inflammation and infection burden were originated and induced by gut because of bacterial overgrowth and gut-derived endotoxin seemed to be the most attractive explanation. Therapies targeting the infection and inflammation derived from the gut are effective, and a new development of non-absorbed rifaximin with patient tolerance became a promising agent. The therapeutic actions of rifaximin were reported to be twofold in reducing the number of ammonia-producing enteric bacteria in the gastrointestinal tract, and would initially inhibit the occurrence of systemic inflammation because of enteric bacteria translocation-induced endotoxin. Therefore, improved clinical outcomes in HE patients after applying additional rifaximin may be achieved by interrupting the pathway of gut-derived local and systematic infection and inflammations, as well as by inhibiting the potential of involvement and injury in organs including liver, lung, brain, and the body’s immune system. All of these would contribute to a significant reduction in length of hospital stay. Lactulose is reported to be associated with increased mild adverse events in the gastrointestinal system, such as diarrhea and abdominal pain. Although it may lead to some temporary discomfort in HE patients, most do not need clinical interventions. Compared with lactulose, rifaximin can be better tolerated with fewer incidences of adverse events. It was reported that compared with other absorbed and systemic antibiotics, non-absorbed rifaximin induced lower risk of bacteria resistance; its plasma concentration was negligible, which indicated a very low risk of bacteria selection outside the gut. In the current study, a combination of the two agents did not increase any risk of adverse events related to the treatment.

<table>
<thead>
<tr>
<th>Study or subgroup</th>
<th>Combination therapy events</th>
<th>Lactulose events</th>
<th>Weight (%)</th>
<th>Risk difference M–H, random, 95% CI</th>
<th>Risk difference M–H, random, 95% CI</th>
</tr>
</thead>
<tbody>
<tr>
<td>Hasan 2018</td>
<td>13</td>
<td>10</td>
<td>9.9</td>
<td>0.07 (–0.11, 0.25)</td>
<td></td>
</tr>
<tr>
<td>Sharm 2018</td>
<td>15</td>
<td>28</td>
<td>10.5</td>
<td>–0.25 (–0.42, –0.09)</td>
<td></td>
</tr>
<tr>
<td>Gill 2014</td>
<td>20</td>
<td>100</td>
<td>13.1</td>
<td>–0.20 (–0.32, –0.08)</td>
<td></td>
</tr>
<tr>
<td>Haq 2014</td>
<td>17</td>
<td>80</td>
<td>12.1</td>
<td>–0.20 (–0.34, –0.06)</td>
<td></td>
</tr>
<tr>
<td>Courson 2015</td>
<td>20</td>
<td>19</td>
<td>11.8</td>
<td>0.10 (–0.04, 0.25)</td>
<td></td>
</tr>
<tr>
<td>Kang 2017-a</td>
<td>53</td>
<td>157</td>
<td>14.8</td>
<td>–0.20 (–0.30, –0.11)</td>
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<tr>
<td>Kang 2017-b</td>
<td>130</td>
<td>448</td>
<td>16.4</td>
<td>–0.08 (–0.15, –0.00)</td>
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</tr>
<tr>
<td>Ahire 2017</td>
<td>2</td>
<td>4</td>
<td>11.3</td>
<td>–0.08 (–0.23, 0.07)</td>
<td></td>
</tr>
<tr>
<td>Total (95% CI)</td>
<td>700</td>
<td>1,122</td>
<td>100</td>
<td>–0.11 (–0.19, –0.03)</td>
<td></td>
</tr>
</tbody>
</table>

Figure 6: Meta-analysis result of mortality in overall analysis between combination therapy and lactulose alone.

Meanwhile, clinical analysis reported that lactulose therapy was related to a non-response rate as high as 22%. Accordingly, improved therapy based on lactulose is warranted since no certain method was introduced to identify a non-responder of lactulose therapy. Confirming by sensitivity analysis, subgroup analysis, Trial Sequential Analysis, and publication bias analysis, our meta-analysis shows that a combination of rifaximin to lactulose can significantly improve both clinical efficacy of HE manifestations (NNT =4) and mortality (NNT =5) mainly for overt HE patients. Based on limited data in one randomized study, the cause of deaths was significantly reduced only in aspect of sepsis, which indicated that combination therapy may improve clinical outcome by reducing the incidence of severe and systemic infection and inflammation. In past years, multiple factors were included in the explanation of HE progress; among them systemic inflammation and infection burden were originated and induced by gut because of bacterial overgrowth and gut-derived endotoxin seemed to be the most attractive explanation. Therapies targeting the infection and inflammation derived from the gut are effective, and a new development of non-absorbed rifaximin with patient tolerance became a promising agent. The therapeutic actions of rifaximin were reported to be twofold in reducing the number of ammonia-producing enteric bacteria in the gastrointestinal tract, and would initially inhibit the occurrence of systemic inflammation because of enteric bacteria translocation-induced endotoxin. Therefore, improved clinical outcomes in HE patients after applying additional rifaximin may be achieved by interrupting the pathway of gut-derived local and systematic infection and inflammations, as well as by inhibiting the potential of involvement and injury in organs including liver, lung, brain, and the body’s immune system. All of these would contribute to a significant reduction in length of hospital stay. Lactulose is reported to be associated with increased mild adverse events in the gastrointestinal system, such as diarrhea and abdominal pain. Although it may lead to some temporary discomfort in HE patients, most do not need clinical interventions. Compared with lactulose, rifaximin can be better tolerated with fewer incidences of adverse events. It was reported that compared with other absorbed and systemic antibiotics, non-absorbed rifaximin induced lower risk of bacteria resistance; its plasma concentration was negligible, which indicated a very low risk of bacteria selection outside the gut. In the current study, a combination of the two agents did not increase any risk of adverse events related to the treatment.

<table>
<thead>
<tr>
<th>Study or subgroup</th>
<th>Combination therapy events</th>
<th>Lactulose events</th>
<th>Weight (%)</th>
<th>Risk difference M–H, random, 95% CI</th>
<th>Risk difference M–H, random, 95% CI</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sharm 2013-RCT</td>
<td>15</td>
<td>28</td>
<td>35.5</td>
<td>–0.25 (–0.42, –0.09)</td>
<td></td>
</tr>
<tr>
<td>Gill 2014-RCT</td>
<td>20</td>
<td>40</td>
<td>64.5</td>
<td>–0.20 (–0.32, –0.08)</td>
<td></td>
</tr>
<tr>
<td>Total (95% CI)</td>
<td>163</td>
<td>157</td>
<td>100</td>
<td>–0.22 (–0.32, –0.12)</td>
<td></td>
</tr>
</tbody>
</table>

Figure 7: Meta-analysis result of mortality in randomized controlled trial between combination therapy and lactulose alone.
Limitations existing in the meta-analysis mainly included study quality, outcome measures, and targeted population. Randomized studies are regarded as higher level evidence than non-randomized studies. The current meta-analysis included both randomized and non-randomized studies to enable a large enough sample size. For the sensitivity analysis, excluding non-randomized studies did not alter the results despite a small difference in the value of effect size. For outcome measures, the current meta-analysis can only analyze data extracted from original reports. Clinical efficacy was judged by different centers; this difference across the studies may induce moderate heterogeneity although it is comparable between groups within each study. Development of a valid and standard test scale of HE is necessary in future research. Currently, HE treatment is mainly targeting at overt HE, because of lower diagnosis rate in covert HE. Combination therapy achieves better clinical outcomes than lactulose alone. However, whether rifaximin and lactulose therapy should be immediately recommended as the first-line treatment for overt HE or an additive therapy of rifaximin for only lactulose-non-response overt HE still needs to be determined. Kang et al performed a cost-analysis study, and showed that the 1-year incremental cost was as high as $85,560 to increase the survival rate in one patient. In another study, Courson et al supposed that although the cost of rifaximin was high, it reduced the overall cost in
patients receiving combination therapy than lactulose alone.\textsuperscript{18} Further cost-effective analyses are needed. Combination therapy of rifaximin and lactulose has beneficial effects on HE. Compared with lactulose alone, combination therapy increases clinical efficacy and decreases mortality in HE patients. However, its effects on different types of HE are still uncertain.

**Conclusion**

Combination therapy of rifaximin and lactulose has beneficial effects on HE. Compared with lactulose alone, combination therapy increases clinical efficacy and decreases mortality in HE patients. However, its effects on different types of HE is still uncertain.

**Disclosure**

The authors report no conflicts of interest in this work.

**References**


