Efficacy and tolerability of renzapride in irritable bowel syndrome: a meta-analysis of randomized, controlled clinical trials including 2528 patients

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Abstract

Introduction: By targeting different subtypes of 5-hydroxytryptamine (5HT) receptors in the gastrointestinal (GI) tract, several drugs have been introduced for the management of irritable bowel syndrome (IBS). Renzapride is a full agonist for 5HT4 receptor and an antagonist to 5HT2b and 5HT3 receptors which is thought a promising therapeutic agent for constipation predominant IBS (C-IBS) patients due to its accelerating effect on the GI tract. In this meta-analysis, our aim was to evaluate the efficacy and tolerability of renzapride in the management of IBS.

Material and methods: A search was done from 1992 to February 2013 for placebo-controlled trials that investigated the efficacy of renzapride in IBS. Renzapride is a full agonist for 5HT4 receptor and an antagonist to 5HT2b and 5HT3 receptors which is thought a promising therapeutic agent for constipation predominant IBS (C-IBS) patients due to its accelerating effect on the GI tract. In this meta-analysis, our aim was to evaluate the efficacy and tolerability of renzapride in the management of IBS.

Results: Relative risk (RR) for clinical efficacy in IBS patients treated for 5 weeks or less comparing renzapride to placebo was 1.07 (95% CI = 0.89–1.29, p = 0.38). This value for IBS patients treated for more than 5 weeks was 1.04 (95% CI = 0.78–1.23, p = 0.77). The RR for clinical efficacy in IBS patients treated with renzapride (4 mg) for 5 weeks or less and more than 5 weeks in comparison to placebo was 1.2 (95% CI = 0.97–1.48, p = 0.1) and 1.16 (95% CI = 0.98–1.37, p = 0.08), respectively, which were statistically non-significant but clinically important. The analysis of tolerability demonstrated that amongst different reported adverse effects, renzapride caused diarrhea more than placebo (RR = 1.61 with a 95% CI = 1.16–2.24, p = 0.004). The RR for withdrawals from renzapride compared to placebo was 1.58 (95% CI = 1.26–2.07, p = 0.0007).

Conclusions: Renzapride is not superior to placebo in relieving IBS symptoms and causes significant incidences of diarrhea and drop-outs due to adverse effects in treated patients vs. placebo. Thus, this medicine might be a cost burden to patients without providing good effectiveness.

Key words: renzapride, 5-hydroxytryptamine, irritable bowel syndrome, clinical trial, meta-analysis, systematic review.

Introduction

Irritable bowel syndrome (IBS) is a highly prevalent chronic functional disorder of the gastrointestinal (GI) tract which has different prospects
in patients. Besides predominant symptoms such as abdominal pain and bloating, altered bowel habits classify the syndrome as diarrhea, constipation or alternative predominant types. Several pathophysiological mechanisms have been identified in IBS such as genetics, visceral hypersensitivity, GI motility dysfunction, inflammation and altered bowel microbial flora, and imbalance in secretion of 5-hydroxytryptamine (5HT) [1, 2]. Forasmuch as the exact cause of this disorder is not fully understood, management of IBS is limited to relieving the main symptoms in patients. Fiber, antispasmodics [3], tricyclic antidepressants [4], serotonin reuptake inhibitors [5], antibiotics [6], probiotics [7] and several herbal preparations [8], besides cognitive behavioral therapy [9], are currently administered alone or in combination by physicians. Other types of medicines which have been introduced are 5HT receptor modulators. The 5HT secreted from enterochromaffin cells regulates the GI tract motility. By targeting different subtypes of receptors (5HT1, 5HT3, 5HT4), several drugs have been introduced such as alosetron, cilansetron (5HT3 antagonist) and tegaserod (5HT4 partial agonist) [10, 11]. Prucalopride, mosapride (5HT4 agonist) and ramosetron (5HT3 antagonist) are novel substances under evaluation in phase III of a clinical trial; we have recently reviewed their efficacy, safety and pharmacokinetic issues [12].

Renzapride (a novel benzamide substitute) is a full agonist for 5HT4 receptor and an antagonist to 5HT2b and 5HT3 receptors. By stimulating 5HT4 and 5HT2b receptors, it accelerates the GI tract transit and motility [13]. It is thought a promising therapeutic agent for constipation predominant IBS (C-IBS) patients. Forasmuch as pharmacokinetic and safety studies demonstrated its well tolerability with less adverse effects [14, 15], several clinical trials have been performed to evaluate its potential efficacy in IBS patients. Several clinical trials have confirmed that renzapride does not cause cardiac arrhythmias in clinical dosages, unlike cisapride [14, 15]. Renzapride is excreted renal and it is not metabolized by cytochrome P450 enzymes. Thus no drug interactions via affecting cytochrome P450 enzymes have been reported [13, 15]. Due to its demonstrated prokinetic property, most trials have shown that renzapride increases colonic transit and reduces transit time and pain in IBS patients which is beneficial especially in those with constipation [16]. In addition, in dose finding studies, a dose-dependent relation has been observed for its effects [15]. In a meta-analysis performed in 2009, renzapride showed no benefit in IBS patients [17].

In the present meta-analysis, the updated results of clinical trials on efficacy and tolerability of renzapride in IBS patients have been evaluated.

Material and methods

Renzapride (C16H22ClN3O2) molecular formula and structure was presented in Figure 1.

Data sources

We searched PubMed, Google Scholar, Web of Science, Scopus, and Cochrane Central Register of Controlled trials for placebo-controlled trials that investigated the efficacy of renzapride in IBS management. Data were collected from 1992 to February 2013. The search terms were “renzapride”, “5-hydroxytryptamine”, “irritable bowel syndrome”, “constipation”, “functional bowel disease” and “irritable colon”. All published studies as well as abstracts presented at the meetings were evaluated without any language limitations. The reference list from retrieved articles was reviewed for additional applicable studies.

Study selection

Two reviewers independently examined the title and abstract of each article to eliminate duplicates, reviews, case studies, pilot studies and uncontrolled trials. All studies investigating effects of renzapride in IBS patients compared to placebo were considered. Trials were disqualified if their outcomes did not have any relation to clinical improvement. We included studies that used Rome criteria for IBS diagnosis. The reviewers independently extracted data on patient’s characteristics, sample size, dosage, trial duration, and outcome measures. There were no disagreements between reviewers.

Figure 1. Molecular formula and structure of renzapride
Assessment of trial quality

Quality of studies was rated by use of the Jadad score, which is based on adherence of studies to randomization, blinding, and dropouts (withdrawals) [18]. The quality scale ranges from 0 to 5 points with a low quality report of score 2 or less and a high quality report of score at least 3.

Statistical analysis

Data from selected studies were extracted in the form of 2 × 2 tables by study characteristics. Included studies were weighted and pooled. Data were analyzed using StatsDirect software version 2.7.9. Relative risk (RR) and 95% confidence intervals (95% CI) were calculated using Mantel-Haenszel, Rothman-Boice (for fixed effects) or Der Simonian-Laird (for random effects) methods. The Cochran Q test was used to test heterogeneity and p < 0.05 was considered significant. In case of heterogeneity or few included studies, the random effects model was used. Funnel plot was used as a publication bias indicator. Clinical importance was evaluated by the Edwards-Nunnally method.

Results

The electronic searches yielded 752 items: 16 from PubMed, 601 from Google Scholar, 106 from Scopus, 23 from Web of Science and 6 from the Cochrane Central Register of Controlled Trials. Of these, six were scrutinized in full text, and four were considered eligible, had a well-defined global response outcome and were included in this analysis (Figure 2). Two of the studies had a quality score of 4 [19, 20] and the two other studies had a score of 3 [21, 22] (Table I). These four trials included 2528 patients randomized to receive either renzapride or placebo. Of the total, 2421 (95.77%) were women and 107 (4.23%) were men. In three of the trials C-IBS patients (meeting the Rome criteria) were involved [19, 21, 22] and in one trial non C-, non D-IBS patients were involved [20]. Patients’ characteristics, type, and dosage of renzapride and placebo, duration of treatment, and outcomes (clinical improvement and the relief of abdominal pain and discomfort) for each study are shown in Tables II and III. Different adverse events of renzapride compared to placebo in IBS patients are summarized in Table IV.

Effectiveness

Clinical efficacy of renzapride in comparison to placebo in irritable bowel syndrome patients for 5 weeks or less than 5 weeks therapy

The summary of RR for clinical efficacy in IBS patients treated for 5 weeks or less in 3 included trials comparing renzapride to placebo [20–22] was 1.07 with 95% CI = 0.89 to 1.29 (p = 0.38, Figure 3 A). The Cochran Q test for heterogeneity indicated that the studies are not heterogeneous (p = 0.51, Figure 3 B) and could be combined but because of few included studies the random effects for individual and summary of RR was applied. Regression of normalized effect vs. precision for all included studies for clinical efficacy in IBS patients treated for 5 weeks or less with

| Study             | Randomization | Blinding | Withdrawals and dropouts | Total score |
|-------------------|---------------|----------|---------------------------|-------------|
| George et al. 2008 | 1             | 1        | 1                         | 3           |
| Lembo et al. 2010  | 2             | 1        | 1                         | 4           |
| Camilleri et al. 2004 | 1          | 1        | 1                         | 3           |
| Spiller et al. 2008 | 2             | 1        | 1                         | 4           |
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Clinical efficacy of renzapride in comparison to placebo in irritable bowel syndrome patients for more than 5 weeks therapy

The summary of RR for clinical efficacy in IBS patients treated for more than 5 weeks in 3 included trials comparing renzapride to placebo [19, 20, 22] was 1.04 with 95% CI = 0.78 to 1.39 (p = 0.77, Figure 4 A). The Cochrane Q test for heterogeneity indicated that the studies are heterogeneous (p = 0.004, Figure 4 B) and could not be combined; thus the random effects for individual and summary of RR was applied. Regression of normalized effect vs. precision for all included studies for clinical efficacy in IBS patients treated for more than 5 weeks with renzapride vs. placebo could not be calculated because of too few strata.

Clinical efficacy of renzapride 4 mg daily in comparison to placebo in irritable bowel syndrome patients for 5 weeks or less than 5 weeks therapy

The summary of RR for clinical efficacy in IBS patients treated for 5 weeks or less in 3 included trials comparing renzapride to placebo [20–22] was 1.2 with 95% CI = 0.97 to 1.48 (p = 0.1, Figure 5 A), a statistically non-significant result but clinically important. The Cochrane Q test for heterogeneity indicated that the studies are not heterogeneous (p = 0.37, Figure 5 B) and could be combined but because of few included studies the random effects for individual and summary of RR was applied. Regression of normalized effect vs. precision for all included studies for clinical efficacy in IBS patients treated for 5 weeks or less with renzapride 4 mg daily vs. placebo could not be calculated because of too few strata.

Clinical efficacy of renzapride 4 mg daily in comparison to placebo in irritable bowel syndrome patients for more than 5 weeks therapy

The summary of RR for clinical efficacy in IBS patients treated for more than 5 weeks in 3 included trials comparing renzapride to placebo [19, 20, 22] was 1.16 with 95% CI = 0.98 to 1.37 (p = 0.08, Figure 6 A), a statistically non-significant result but clinically important. The Cochrane Q test for heterogeneity indicated that the studies are heterogeneous (p = 0.2, Figure 6 B) and could be combined but because of few included trials the random effects for individual and summary of RR was applied. Regression of normalized effect vs. precision for all included studies for clinical efficacy in IBS patients

### Table II. Characteristics of studies included in the meta-analysis

| Trial | Treatment duration [weeks] | Time of reporting [weeks] | Patients | Outcomes |
|-------|---------------------------|--------------------------|----------|----------|
| George et al. 2008 | 12 | 4 | 510 C-IBS (Rome II) | ↑ Responders rate (relief of pain/discomfort) dose dependently; improved bowel movement frequency and stool consistency; more effective in females |
| Lembo et al. 2010 | 12 | 48 (for long-term safety study) | 1798 C-IBS (Rome II) | Improved stool consistency and frequency; bloating, abdominal distention and pain; ↑ quality of life score; ischemic colitis in long-term study |
| Camilleri et al. 2004 | 2 | 0 | 48 C-IBS (Rome II) | ↑ Colon transit time; improved bowel function and satisfactory symptom relief; more effective in females |
| Spiller et al. 2008 | 4 | 4.8 | 4.8 C-IBS (Rome II) | No sig IBS symptoms improvement; relief of pain/distress most effective in females |

b.d. – twice daily, C-IBS – constipation predominant IBS, D-IBS – diarrhea predominant IBS, F – female, IBS – irritable bowel syndrome, M – male, ↑ – increase
treated for more than 5 weeks with renzapride 4 mg daily vs. placebo could not be calculated because of too few strata.

**Tolerability**

Different adverse events of renzapride compared to placebo in irritable bowel syndrome patients

The summary RR for diarrhea in 4 trials [19–22] was 1.61 with a 95% CI of 1.16–2.24 and a significant RR \( p = 0.004 \). The Cochrane Q test for heterogeneity indicated that the studies are homogeneous \( p = 0.28 \) and could be combined; thus the fixed effects for individual and summary of RR was applied.

The summary RR for headache in 4 trials [19–22] was 1.21 with a 95% CI of 0.93–1.56 and a non-significant RR \( p = 0.16 \). The Cochrane Q test for heterogeneity indicated that the studies are not heterogeneous \( p = 0.39 \) and could be combined; thus the fixed effects for individual and summary of RR was applied.

**Table III.** Clinical improvement (IBS relief) in included studies

| Study             | Time of reporting (duration) |
|-------------------|-------------------------------|
|                   | ≤ 5 weeks | > 5 weeks |
|                   | Renzapride | Placebo | Renzapride | Placebo |
| George et al. 2008 | 188/384 | 58/125 | 190/384 | 71/125 |
| Lembo et al. 2010 | – | – | 378/1198 | 146/600 |
| Camilleri et al. 2004 | 15/36 | 2/11 | – | – |
| Spiller et al. 2008 | 57/126 | 18/42 | 68/126 | 23/42 |

**Table IV.** Different adverse events of renzapride compared to placebo in IBS patients

| Adverse events      | RR combined meta-analysis | 95% CI | Value of \( p \) for RR | Value of \( p \) for heterogeneity | Publication bias |
|---------------------|---------------------------|--------|--------------------------|-------------------------------|-----------------|
| Diarrhea            | 1.61                      | 1.16–2.24 | 0.004                     | 0.28                          | No              |
| Headache            | 1.21                      | 0.93–1.56 | 0.16                      | 0.39                          | No              |
| Abdominal pain      | 1.37                      | 0.95–1.98 | 0.09                      | 0.51                          | NA              |
| Constipation (aggravated) | 0.98                  | 0.55–1.74 | 0.94                      | 0.98                          | NA              |
| Nausea              | 1.27                      | 0.76–2.11 | 0.36                      | 0.27                          | NA              |
| Dyspepsia (gas)     | 0.94                      | 0.45–1.96 | 0.88                      | 0.96                          | NA              |
| Vomiting            | 1.2                       | 0.74–1.95 | 0.47                      | 0.75                          | No              |

NA – not applicable
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Figure 4. A – Individual and pooled relative risk for the outcome of “clinical efficacy treated for more than 5 weeks” in the studies considering renzapride compared to placebo therapy in IBS patients. B – Heterogeneity indicators for the outcome of “clinical efficacy treated for more than 5 weeks” in the studies considering renzapride compared to placebo therapy in IBS patients.

Figure 5. A – Individual and pooled relative risk for the outcome of “clinical efficacy treated for 5 weeks or less” in the studies considering renzapride 4 mg daily compared to placebo therapy in IBS patients. B – Heterogeneity indicators for the outcome of “clinical efficacy treated for 5 weeks or less” in the studies considering renzapride 4 mg daily compared to placebo therapy in IBS patients.

Figure 6. A – Individual and pooled relative risk for the outcome of “clinical efficacy treated for more than 5 weeks” in the studies considering renzapride 4 mg daily compared to placebo therapy in IBS patients. B – Heterogeneity indicators for the outcome of “clinical efficacy treated for more than 5 weeks” in the studies considering renzapride 4 mg daily compared to placebo therapy in IBS patients.
The summary RR for abdominal pain in 3 trials [19, 20, 22] was 1.37 with a 95% CI of 0.95–1.98 and a non-significant RR ($p = 0.09$). The Cochrane Q test for heterogeneity indicated that the studies are homogeneous ($p = 0.51$) and could be combined but because of few included studies the random effects for individual and summary of RR was applied.

The summary RR for constipation (aggravated) in 2 trials [20, 22] was 0.98 with a 95% CI of 0.55–1.74 and a non-significant RR ($p = 0.94$). The Cochrane Q test for heterogeneity indicated that the studies are homogeneous ($p = 0.98$) and could be combined but because of few included studies the random effects for individual and summary of RR was applied.

The summary RR for nausea in three trials [19, 21, 22] was 1.27 with a 95% CI of 0.76–2.11 and a non-significant RR ($p = 0.36$). The Cochrane Q test for heterogeneity indicated that the studies are homogeneous ($p = 0.27$) and could be combined but because of few included studies the random effects for individual and summary of RR was applied.

The summary RR for dyspepsia (gas) in two trials [21, 22] was 0.94 with a 95% CI of 0.45–1.96 and a non-significant RR ($p = 0.88$). The Cochrane Q test for heterogeneity indicated that the studies are homogeneous ($p = 0.96$) and could be combined but because of few included studies the random effects for individual and summary of RR was applied.

The summary RR for vomiting in 4 trials [19–22] was 1.2 with a 95% CI of 0.74–1.95 and a non-significant RR ($p = 0.47$). The Cochrane Q test for heterogeneity indicated that the studies are homogeneous ($p = 0.75$) and could be combined; thus the fixed effects for individual and summary of RR was applied (Table IV).

Withdrawal due to adverse effect, non-compliance and lack of efficacy with renzapride compared to placebo therapy in irritable bowel syndrome patients

The summary of RR for withdrawal in IBS patients in 3 included trials comparing renzapride to placebo [19, 20, 22] was 1.58 with 95% CI = 1.26 to 2.07 ($p = 0.0007$, Figure 7 A). The Cochrane Q test for heterogeneity indicated that the studies are not heterogeneous ($p = 0.73$, Figure 7 B) and could be combined but because of few included studies, the random effects for individual and summary of RR was applied. Regression of normalized effect vs. precision for all included studies for withdrawal in IBS patients receiving renzapride vs. placebo therapy could not be calculated because of too few strata.

**Discussion**

This meta-analysis included a total of 2528 C-IBS and non C-, non D-IBS patients according to the mentioned inclusion criteria from randomized placebo-controlled clinical trials. In all studies, the criteria for diagnosis of IBS patients was based on Rome criteria. Regarding the statistical analysis, the present results confirm that renzapride has no significant advantage over placebo in relieving symptoms in IBS patients. Regarding the prevalence and high burden of IBS, developing novel therapeutic agents for this syndrome is of great value. To reach a convincing conclusion on effectiveness of each new suggested drug, assessing clinical trials in which drug was compared to placebo could be clarifying. We analyzed data for efficacy twice based on duration of treatment. The results demonstrated that increasing the duration of treatment (more than 5 weeks) does not influence the efficacy of renzapride compared to placebo.
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Several systematic reviews have assessed the efficacy of 5-HT receptor modulators in IBS patients. Ford et al. performed a systematic review on the efficacy of known 5-HT3 antagonists and 5-HT4 agonists in IBS in 2009 [17]. They conducted their meta-analysis by reviewing placebo-controlled clinical trials up to the year 2008. However, our meta-analysis included a 12-week double-blind, randomized, placebo-controlled clinical trial by Lembo et al. [19] in 2010, in which a total of 1798 female IBS patients were included for both efficacy and safety analysis. As in the present meta-analysis, Ford et al. confirmed that renzapride and cisapride were not more effective than placebo in IBS patients. Another group of researchers also evaluated the efficacy of mixed 5HT3 antagonists/5HT4 agonists (cisapride and renzapride) in IBS patients [22] and found that renzapride in doses of 1 mg and 2 mg was not effective in relieving IBS symptoms, which further supports the present meta-analysis. They demonstrated that a dose of 4 mg was significantly more effective than placebo. For clarification, we analyzed the clinical efficacy of renzapride 4 mg separately in comparison to placebo for 5 weeks or less, and more than 5 weeks treatment durations. Although differences were not statistically significant, the results were clinically important and significant for both treatment durations. Therefore, these results could be considered for renzapride 4 mg while more trials are necessary to conclude the effectiveness of this novel drug more precisely. Several adverse effects have been reported for renzapride that were mainly in the GI tract. By analyzing the reported adverse effects, we found that there was no statistically significant difference between renzapride and placebo, except in the occurrence of diarrhea, which was higher in patients receiving renzapride.

In addition, renzapride caused more withdrawals due to adverse effects and/or low efficacy in patients. In the same study, no significant difference was observed between renzapride and placebo for occurrence of adverse effects in patients [23].

One of the limitations of this meta-analysis is the differences in the characteristics of involved patients (age, sex, life style and compliance), which is inevitable and results from deficits of included trials. In addition, included trials had different duration of treatment and endpoints. Treatment durations ranged from 2 weeks [21] to 12 weeks [19, 22]. Therefore, to avoid heterogeneity, we divided our data into two groups according to treatment duration and time of reporting the results (5 weeks or less, and more than 5 weeks), although there were few data in each group. In these 4 assessed clinical trials, 3 doses of renzapride were used in patients (1, 2, and 4 mg daily). We did not observe any significant difference between renzapride and placebo, except for 4 mg, which in comparison to placebo had a clinically significant difference. To have more reliable results on the efficacy of renzapride and to find a possible dose-dependent effect, we analyzed data based on renzapride dose and compared the 4 mg group to placebo separately. But, similar to duration of treatment, there were few patients in each group. In one of the trials the experimental population members were female [19] and in other three studies the authors reported better efficacy of renzapride in females than males [20–22]. Therefore to have a bigger population of patients with more homogeneity in sex and age to reach a more powerful conclusion, further clinical trials are essential. Although the results of this meta-analysis do not confirm the significant effectiveness of renzapride in IBS patients, regarding the acceptable quality score of included trials and lack of other suitable choices, it can still be recommended in C-IBS patients due to its accelerating effects on GI motility.

In conclusion, renzapride is not only superior to placebo in relieving IBS symptoms (abdominal pain and discomfort) but also causes significant incidence of diarrhea and drop-outs due to adverse effects in patients. This means that this drug might be a cost burden to patients without providing good efficacy or advantages.

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